

Journal of Experimental Botany, Vol. 74, No. 17 pp. 5273–5293, 2023 https://doi.org/10.1093/jxb/erad207 Advance Access Publication 8 June 2023



RESEARCH PAPER

Contrasting sap flow characteristics between pioneer and late-successional tree species in secondary tropical montane forests of Eastern Himalaya, India

Manish Kumar^{1,2,3,*,}, Gladwin Joseph^{1,4,}, Yangchenla Bhutia^{1,2,5,} and Jagdish Krishnaswamy^{1,2,6,}

- ¹ Ashoka Trust for Research in Ecology and the Environment (ATREE), Bangalore 560064, Karnataka, India
- ² Manipal Academy of Higher Education (MAHE), Manipal 576104, Karnataka, India
- ³ School of Geography, Earth and Environmental Sciences, University of Birmingham, Birmingham B15 2TT, UK
- ⁴ Conservation Biology Institute, Corvallis, Oregon 97333, USA
- ⁵ Sikkim State Council of Science & Technology, Gangtok 737102, Sikkim, India
- ⁶ School of Environment and Sustainability, Indian Institute for Human Settlements, Bangalore 560080, Karnataka, India

Received 12 August 2022; Editorial decision 24 May 2023; Accepted 30 May 2023

Editor: Menachem Moshelion, The Hebrew University of Jerusalem, Israel

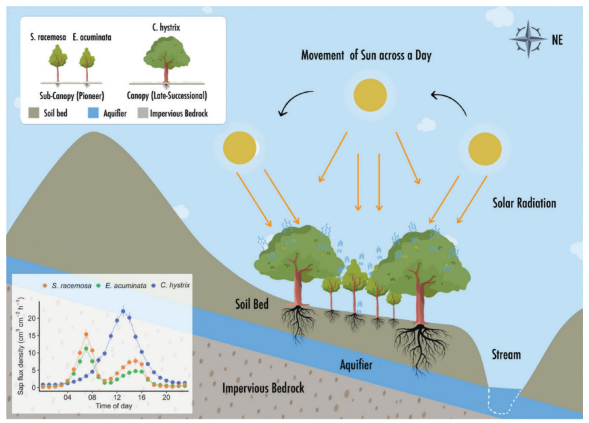
Abstract

The interactive role of life-history traits and environmental factors on plant water relations is crucial for understanding the responses of species to climate change, but it remains poorly understood in secondary tropical montane forests (TMFs). In this study, we examined differences in sap flow between the pioneer species Symplocos racemosa and Eurya acuminata, and the late-successional species Castanopsis hystrix that co-occur in a biodiverse Eastern Himalayan secondary broadleaved TMF. The fast-growing pioneers had sap flux densities that were 1.6-2.1 times higher than the late-successional species, and exhibited characteristics of long-lived pioneer species. Significant radial and azimuthal variability in sap flow (V) between species was observed and could be attributed to the life-history trait and the access of the canopy to sunlight. Nocturnal V was 13.8% of the daily total and was attributable to stem recharge during the evening period (18.00-23.00 h) and to endogenous stomatal controls during the pre-dawn period (00.00-05.00 h). The shallow-rooted pioneer species both exhibited midday depression in V that was attributable to photosensitivity and diel moisture stress responses. In contrast, the deep-rooted late-successional species showed unaffected transpiration across the dry season, indicating their access to groundwater. Thus, our results suggest that secondary broadleaved TMFs, with a dominance of shallow-rooted pioneers, are more prone to the negative impacts of drier and warmer winters than primary forests, which are dominated by deep-rooted species. Our study provides an empirical understanding of how life-history traits coupled with microclimate can modulate plant water use in the widely distributed secondary TMFs in Eastern Himalaya, and highlights their vulnerability to warmer winters and reduced winter precipitation due to climate change.

Abbreviations: ACF, autocorrelation coefficient; corARMA, second-order autoregressive moving average; CV, coefficient of variability; GLS, generalized least squares; KGE, Kling–Gupta efficiency; MAP, mean annual precipitation; masl, meters above sea level; MLRM, multiple linear regression models; SPAC, soil–plant–atmosphere continuum; TDP, thermal dissipation probes; TMF, tropical montane forests; VPD, vapour pressure deficit.

^{*} Correspondence: m.kumar.2@bham.ac.uk or manish.kumar@atree.org

Graphical Abstract



The access to soil moisture and sunlight dictates diurnal sap flux density (J in cm³ cm⁻² h⁻¹) patterns among co-occurring pioneer and emergent species in a secondary tropical montane forest in Eastern Himalaya.

Keywords: Ecohydrology, Himalaya, late-successional, pioneer species, plant water relations, sap flow, secondary forests, transpiration.

Introduction

Plant water relations play a crucial role in modulating carbon and water fluxes in recovering secondary tropical forests. In most terrestrial systems, the availability of energy and water are the primary drivers of water use by vegetation, mainly through transpiration (Heimann and Reichstein, 2008; Asbjornsen et al., 2011). Transpiration is the movement of water from the soil to the atmosphere via the stem based on a gradient of total water potential along the soil-plant-atmosphere-continuum (SPAC), and is an active area of research in ecohydrology (Asbjornsen et al., 2011; Manzoni et al., 2013; Mencuccini et al., 2019). Transpiration is driven by environmental variables such as vapour pressure deficit (VPD), light, and soil moisture availability, and by ecophysiological traits controlling plant water use such as wood anatomy, rooting depth, stomatal density, and sapwood density (Asbjornsen et al., 2011; Manzoni et al., 2013). These ecophysiological traits have evolved through competition and adaptation between co-occurring species belonging to different functional groups, such as pioneer or late-successional, and they contribute to the interspecific variability in sap flow (O'Brien et al., 2004; Hernandez-Santana et al., 2015). Research typically divides interspecific variation along a fast-slow/growth-survival axes, where pioneer species tend to adopt a 'live fast and die young' approach to maximize growth (with high plant water use), while climax or late-successional species adopt a low-risk, slow-growth strategy (with conservative water use) with high survival rates (Rüger et al., 2020; Lai et al., 2021).

In terms of intraspecific variations, these are manifested as radial and azimuthal (between cardinal directions) variability in sap flow and are influenced by growth stage, plant size, stem anatomy (ring-porous versus diffuse-porous species), and individual responses to the microclimate (Fiora and Cescatti, 2006; Berdanier et al., 2016; Berry et al., 2018). Individual plant- and species-level studies are therefore crucial for understanding

the responses of co-occurring species belonging to similar or different functional groups to changing environmental conditions, especially in biodiverse tropical montane forests (TMFs; Forrester, 2015; Bretfeld et al., 2018; Basnett and Devy, 2021). Such studies are also crucial to our understanding of the responses of secondary forest communities to the predicted intensifications of hydrological cycles with climate change, especially to extreme precipitation events, seasonal droughts, and reductions in winter precipitation (Donohue et al., 2007; Zeppel et al., 2008, 2011; Kulkarni et al., 2013; Bretfeld et al., 2018). The impact of atmospheric CO₂ fertilization on plant water use and carbon assimilation rates, and hence on carbon sequestration potential, is another important area of research in TMFs (Bugmann and Bigler, 2011; Eckert et al., 2021).

Yet, ecohydrological studies describing the linkages between micro-environment and ecophysiological traits associated with transpiration and their underlying mechanisms are relatively sparse in TMFs. Most studies have been restricted to the tropical Andes and South-East Asia, and very little research has come from the Himalaya region (Bruijnzeel et al., 2011; Poyatos et al., 2021); such studies that have been carried out have mainly focused on interspecific patterns in leaf water potential with environmental gradients, with most coming from the drier and relatively higher latitudes of Western Himalaya (Zobel et al., 2001; Poudyal et al., 2004; Singh et al., 2006). Direct measurements of whole-tree transpiration are rare, with Ghimire et al. (2014) being the first to compare the variability in transpiration rates between needle-leaved Chir pine (Pinus roxburghii) and broadleaved oak forests in Central Himalaya using sap-flow probes. Other studies have also highlighted the crucial role of fluctuations in seasonal soil moisture on phenology and growth cycles in broadleaved trees in the Himalaya (Singh et al., 2000; Tewari et al., 2016; Chand et al., 2017). The variability in ecophysiological responses of Himalayan trees to seasonal moisture deficit has also been examined across differing elevations, species ranges, and canopy structures (Zobel et al., 2001; Singh et al., 2006; Tewari et al., 2018). As a result of the long history of human forest use, the majority of broadleaved TMFs in the Himalaya are secondary forests, and they remain understudied (Ramakrishnan and Kushwaha, 2001; Kanade and John, 2018). Secondary TMFs differ significantly from their primary counterparts in that species of different successional stages such pioneer and late-successional co-exist, together with mixed stand structures (sub-canopy and emergent trees) (Ramakrishnan and Kushwaha, 2001; Nogueira et al., 2004). In the Himalayan state of Sikkim in northeast India, secondary TMFs constitute a third of the total forest cover and form a significant carbon sink, and are a source of the springs and streams that provide principal water resources (Wohl et al., 2012; Brown and Lugo, 1990; Daniel et al., 2021; Kumar et al., 2023). The majority of these forests are under protected area management, and understanding their plant water relations is critical to the water and ecological security of the Eastern Himalaya (Kanade and John, 2018). The region

is threatened by climate change, with projections indicating a high rate of warming (+5 °C), a significant increase in precipitation (+40%), and a high loss of biodiversity (33%) by the end of the this century (Kulkarni et al., 2013; Krishnan et al., 2019; Dahal et al., 2021). The forests in parts of the Himalaya have already shown trends of browning in response to warminginduced moisture stress in recent decades (Krishnaswamy et al., 2014), making a better understanding of plant water relations a research priority.

To the best of our knowledge, plant water relations and transpiration have not been studied in the wet secondary TMFs of Eastern Himalaya using direct whole-tree methods such as sap flow. Further, the variability in water-use strategies among different functional groups in response to seasonally moist and dry conditions and its impact on plant productivity and the carbon sequestration potential also remains to be explored in these human-influenced biodiverse forest communities (Bretfeld et al., 2018). The current study aims to address this knowledge gap using a response-based approach to understanding the variability in plant water use among co-occurring pioneer and late-successional species in relation to micro-meteorological and environmental factors under a SPAC framework (Mencuccini et al., 2019; Kannenberg et al., 2022). Specifically, this study addresses the following two questions within the context of an East Himalayan broadleaved wet evergreen TMF. (i) How do the patterns of sap flow differ between co-occurring pioneer and late-successional species in a secondary forest? (ii) What are the environmental and ecophysiological drivers of variability in sap flow? We hypothesized that plant water use by the pioneer species would be high and would likely be highly sensitive to environmental extremes, whereas late-successional species would have relatively stable water use patterns. We also predicted that, unlike other TMFs, the ecosystem productivity was more likely to be limited by energy availability than water in the wet East Himalayan eco-climate.

Materials and methods

Site description

The forest stand used as the experimental site represents the East Himalayan broadleaved evergreen wet montane forest classification (Sudhakar et al., 2008; Kanade and John, 2018; Bhutia et al., 2019). The site was 2150 m above sea level (masl) with a mean annual precipitation (MAP) of 4650 (±120) mm, making it one of the highest elevation and wettest tropical montane forest (TMF) site to date where direct sap flow-based plant water flow measurements have been carried out (Bruijnzeel et al., 2011; Poyatos et al., 2021). Such TMFs are distinguished from temperate broadleaved evergreen forests (high latitude and low elevation) in terms of being relatively closer to the tropics (20-35°N) and having high elevation (800-2800 masl), a wet/humid climate (MAP 3000-5000 mm), moderate energy conditions (12 °C mean annual temperature), and high biodiversity (Ohsawa, 1993; Tang and Ohsawa, 1999; Kanade and John, 2018). The field study was conducted in a secondary forest stand (27.35°N, 88.56°E) in the Fambong-Lho Wildlife Sanctuary (FWS), Sikkim, India, with annual daily mean temperatures ranging from -2 °C to 24 °C. The hydrologic year at the site is divided into three

distinct seasons, as follows. Summer, (March to May), characterized by warm days, cloudy afternoons, significant pre-monsoon precipitation, and high evapotranspiration; monsoon (June to October), with concentrated precipitation, high humidity, and low evapotranspiration; and winter (November to February) with sunny days, cold nights, occasional snowfall, and moderate evapotranspiration (Kumar *et al.*, 2021). The forest is sloped at 10–35°, predominantly facing northeast, and the soil is a well-drained sandy-loam with moderate depth (40–100 cm).

Vegetation data were derived from five vegetation surveys each of 100 × 10 m, which recorded 321 trees of 16 species belonging to 12 families (Bhutia et al., 2019). The composition marked the stand as an early-successional secondary forest with a short-statured canopy dominated by pioneer species such as Symplocos and Eurya species. Relatively older, taller, and larger-girthed individuals of the Fagaceae family, such as Castanopsis and Quercus species, stood out as emergent trees (Sudhakar et al., 2008; Bhutia et al., 2019; Gurung and Chettri, 2019). Tree density provided a better estimate than basal area for understanding the population structure of pioneer species because of the higher number of smallsized individuals, whereas the late-successional species were larger-sized and fewer in number. The selection of species for sap-flow measurements was based on their representation of the successional stages (pioneer and late-successional), and on selection of dominant species in each of the successional stages according to the tree density and basal area distribution of the stand. Symplocos racemosa and Eurya acuminata were selected as representing the fast-growing pioneer species community, whilst Castanopsis hystrix was selected as the most dominant species belonging to the climax or late-successional stage (Supplementary Fig. S1). Information on rooting depth, phenology, and other ecophysiological characteristics of the three species collated from available literature and personal observations is presented in Table 1. Symplocos racemosa was observed to be flowering and fruiting in June-July during the monsoon season, E. acuminata flowered in December-January at the peak of the dry season and fruited during February-March, whilst C. hystrix flowered in February and fruited in March-April. None of the three species was observed to have any concentrated leaf fall.

Sap flux measurements

Sapwood thickness was measured from wood cores extracted using an increment borer from each selected tree at the start of the experiment. The cores from *C. hystrix* showed distinct heartwood formation and only the outer xylem (4–5 cm) was observed to be functional. No heartwood was detected in *S. racemosa* and *E. acuminata*, and the entire xylem was assumed to be functional, a characteristic of diffuse-porous species (Berdanier *et al.*, 2016). After installing the probes, wood cores were collected from each tree concurrently (14 December 2013), wrapped in plastic cling film immediately after extraction to avoid moisture loss, and then subsequently processed in the laboratory. Each core was cut into 1-cm long sections to estimate wood density and moisture following the method of Chave (2006). Sap flux measurements were carried out from November 2013 to May 2014 for five trees each of *S. racemosa* and *E. acuminata*, and three trees of *C. hystrix*

using the Granier's thermal dissipation probe (TDP) method (Table 2) (Granier, 1987; Lu et al., 2004). A total sample size of 13 trees and 3-5 replicate trees is consistent with the average number that have been used in previous sap flow studies, and the installation design was optimized to cover maximum xylem variability (Asbjornsen et al., 2011; Guyot et al., 2015). Davis et al. (2012) suggested modifications to the lab-built TDP design template to measure sap flow at the top 1 cm of the probe and validated the probes against the commercial Dynamax TDP30 sap flow probe. Our study used lab-built probes based on this modified design template (for further details see Phillips et al. 1996; James et al. 2002; Davis et al. 2012). TDPs the most widely used technique for measuring sap flow and they perform well in cold and lowflow conditions (Lu et al., 2004; Chan and Bowling, 2017; Wang et al., 2023). They are also known to suffer from linear systemic bias and thus are suitable for relative comparisons between species (Peters et al., 2018; Flo et al., 2019). Radial probes of 1–5 cm length were installed at 1 cm incremental depths between 1–5 cm, respectively, from the cambium in one tree for each of the three species (henceforth referred to as 'radial trees'), in a spiral design. The remaining trees for each species (henceforth referred to as 'replicate trees') were fitted with two 2 cm length probes that were installed in the north and south aspects to observe azimuthal variability (Shinohara et al., 2013; Komatsu et al., 2016). A design limitation of using multiple sap-flow probes of different depths in a spiral is the potentially confounding interactions between radial and circumferential variability, which can result in significant biases in the calculation of whole-tree water use, especially in trees with nonuniform xylem growth patterns (James et al., 2002).

The probe signals (in mV) were converted to temperature differences (ΔT , °C) between the heater and reference sensors in a TDP probe at 30 min intervals and stored in a multi-channel data logger (DL2e, Delta-T, UK). The zero-sap flow assumption, at which the temperature difference between the probes is maximum ($\Delta T_{\rm max}$), is critical for Granier's method and a significant source of error in TDP-based sap-flow estimation. When comparing different approaches for determining $\Delta T_{\rm max}$, Rabbel *et al.* (2016) showed that daily $\Delta T_{\rm max}$ has lower variability than studies using moving-window or regression-based approaches, and hence they recommended daily $\Delta T_{\rm max}$ for research in humid environments without significant water limitations. Consequently, we computed sap flux density (J, cm³ cm⁻² h⁻¹) following Granier's empirical equation below, where $\Delta T_{\rm max}$ was determined daily for each probe:

$$J = 119 \times 10^{-6} \times 3600 \times \left(\frac{(\Delta T_{\text{max}} - \Delta T)}{\Delta T}\right)^{1.231}$$
 (1)

The first 2 weeks of data were ignored to avoid errors due to installation wounds (Wiedemann *et al.*, 2016). After quality checks, 114 unique days of sap-flow data between 7 December 2013 and 15 May 2014 were used for the final analysis, and this included 33 d where data were present for all three species (Supplementary Fig. S2; Supplementary Table S1B).

Table 1. Life-forms and ecophysiological characteristics of the three species

Species	Family	Form	Functional group	Canopy position	Rooting depth	Wood type
S. racemosa	Symplocaceae	Small-medium trees	Pioneer	In-canopy	Shallow	Diffuse-porous
E. acuminata	Pentaphylacaceae	Shrubs/small trees	Pioneer	In-canopy	Shallow	Diffuse- porous
C. hystrix	Fagaceae	Large trees	Late-successional	Emergent	Deep	Semi ring- porous

References: Ohsawa et al., 1986; Suzuki et al., 1991; Sundriyal and Sharma, 1996, 2003; Chettri et al., 2002; Sharma et al., 2011; Li et al., 2013.

Table 2. Basic morphological details of the trees measured in the experiment, the number of radial probes installed per species, and number of unique days of sap flow data per species.

Species	No. of trees	DBH (m)	Basal area per tree (m²)	Sapwood area per tree (m²)	No. of radial probes per tree	No. of unique data days
S. racemosa	5	0.21 ± 0.06	0.036 ± 0.02	0.032 ± 0.02	5	49
E. acuminata	5	0.19 ± 0.04	0.028 ± 0.01	0.026 ± 0.01	4	87
C. hystrix	3	0.34 ± 0.2	0.11 ± 0.12	0.037 ± 0.02	5	99

DBH, diameter at breast height. Means (±SD) are shown.

Environmental measurements

Soil water potential was recorded at 10 cm incremental depths from the surface at 10 min resolution using granular matrix-based (watermark) sensors (Virtual Electronics, Roorkee, India) and converted to volumetric water content using site-specific van Genuchten water retention curve parameters. The curve parameters were developed using the percentage of soil organic matter and organic carbon, particle size distribution (sand, silt, and clay), bulk density, and porosity as input parameters in the Rosetta software (Schaap et al., 2001). The hourly total soil moisture (S, in mm) for the topsoil (0-30 cm depth) was computed using the trapezoidal method and smoothed using a three-step moving-average window to fill stray gaps resulting from missing values (Nachabe et al., 2005). In-canopy air temperature and relative humidity were recorded at 10-min resolution (iButton Hygrochrons, Maxim Int., USA). Air temperature (°C), relative humidity $(R_h, \text{ in } \%)$, wind speed $(u, \text{ in m s}^{-1})$, and incoming short-wave radiation $(R_{sol}, \text{ in kW m}^{-2}, \text{ henceforth referred to as solar radiation})$ were recorded with 10 min resolution using an automatic weather station (AWS;Vantage Pro2, Davis Instruments, USA). Vapour pressure deficit (VPD, kPa) was calculated from the hourly air temperature and relative humidity recorded by the Hygrochrons and the AWS and averaged. Precipitation (P, in mm h⁻¹) was recorded using a tipping-bucket rain gauge (Spectrum Technologies, India) fitted with an odyssey data logger (Dataflow systems Ltd., New Zealand). Data processing, analysis, and visualization were done in the R software (https://www.r-project.org).

Data analysis

The data analysis consisted of three main steps: scaling from the probes to whole-tree sap flow, assessing sap flow variability within and between species, and linking the environmental drivers of the observed patterns in sap flow. The scaling procedure followed established literature, and relative biases in whole-tree sap flow were estimated. The intraspecies differences in sap flow were investigated within an individual tree and between individuals of the same species along radial (three metrics) and azimuthal (one metric) axes. The interspecific variability was explored between the two co-occurring functional groups, pioneer and late-successional species. Species-wise multiple linear regression models were used to understand the drivers of day and night sap-flow patterns. SPAC interactions were explored using lag regression analysis while accounting for the relative time lags across the different seasons. The observed temporal lag was combined with a generalized least-squares (GLS) regression time-series model to quantify the relative influence of key environmental drivers on sap flow in the three species, including solar radiation and VPD.

Scaling from sap-flux density to whole-tree sap flow.

A combination of the zero-averaged technique and weighted mean method was used to estimate the whole-tree sap flow in trees with radial probes ($V_{\rm rad}$) in kg h⁻¹) (for details, see Hatton et al. 1990; Pausch et al. 2000). Another estimate of whole-tree sap flow, $V_{\rm rad.2cm}$, was calculated in the radial-probe trees using sap flux density at 2 cm depth only (J_{2cm}) and the total sapwood area (A_{total}) , assuming homogenous radial flow for comparison against V_{rad} :

$$V_{\rm rad.2cm} = J_{\rm 2cm} \times A_{\rm total} \times 10^{-3}$$
 (2)

 $V_{
m rad,2cm}$ showed significant overestimation (15%) in comparison with sap flow estimated using the radial profile $V_{\rm rad}$. Previous studies have used simple linear regression models (LRMs) between integrated whole-tree sap flow (V_{rad}) and sap flow in the outer xylem for trees with only shallow probes ($V_{\text{rad.2cm}}$) (Paudel et al., 2013; Berdanier et al., 2016). However, given the diel variability in their relationships, we fitted linear regression models without intercepts for each hour of the day for each species (Supplementary Table S2). The slopes (m) of the LRMs for each hour were multiplied by the mean sap-flux density between the north- (J_{north}) and south-facing (J_{south}) probes and by the total sapwood area (A_{total}) to estimate whole-tree sap flow in the replicate trees using a linear model (LM) method (V_{rep,LM}, in kg h⁻¹) (Paudel et al., 2013; Berdanier et al., 2016), as follows:

$$V_{\text{rep.LM}} = \left(\frac{J_{\text{north}} + J_{\text{south}}}{2}\right) \times A_{\text{total}} \times m \times 10^{-3}$$
(3)

For the subsequent analyses, $V_{\rm rad}$ and $V_{\rm rep,LM}$ are combined under the term 'whole-tree sap flow' and represented by $V (\text{kg h}^{-1})$.

Biases in the estimation of whole-tree sap flow.

The relative biases in whole-tree sap flow were estimated in replicate trees by systematically ignoring radial and azimuthal variability and compared with whole-tree sap flow as estimated by incorporating radial variability ($V_{\text{rep.LM}}$, Equation 3) (see Shinohara et al., 2013). First, whole-tree sap flow was estimated for replicate trees using the conventional method (CM) without incorporating the radial variability ($V_{\text{rep.CM}}$, in kg h⁻¹) was estimated by multiplying the A_{total} by the average J determined from the two diametrically opposite 2 cm probes:

$$V_{\text{rep.CM}} = \left(\frac{J_{\text{north}} + J_{\text{south}}}{2}\right) \times A_{\text{total}} \times 10^{-3}$$
 (4)

Second, whole-tree sap flow was estimated for replicate trees based on the north-facing side only ($V_{\rm rep.N}$, in kg ${\rm h^{-1}}$) using the sap flux density for the north-facing probes (J_{north}) and A_{total} while ignoring azimuthal variability

$$V_{\text{rep.N}} = J_{\text{north}} \times A_{\text{total}} \times 10^{-3}$$
 (5)

Intraspecific variability in sap flow.

Within a species, the sap-flow variabilities were visualized and quantified as the variabilities in xylem conductivity along the radial and azimuthal profiles. The radial sap flow variability was observed in one tree per species and quantified using three metrics (Delzon *et al.*, 2004; Fiora and Cescatti, 2006), as follows. (i) Daily maximum sap flux density (J_{max}) was plotted against the depth from the cambium to indicate regions of high sap flux in the sapwood. (ii) The percentage contribution of sapwood annuli at each depth to daily V was estimated as a product of the sapwood area of the annulus and the corresponding J values. (iii) Radial variability was quantified by means of hourly correction factors (C_h), which were estimated for sunny days for trees with radial probes as the sum of hourly ratios of J at different depths (J_i) with reference to $J_{2\text{cm}}$, normalized by the sapwood area of each annulus (A_i) over A_{total} (Delzon *et al.*, 2004), as follows:

$$C_{\rm h} = \sum_{i=1}^{n} \left(\frac{J_i}{J_{\rm 2cm}} \right) \times \left(\frac{A_i}{A_{\rm total}} \right) \tag{6}$$

The azimuthal variability in sap flow was estimated as hourly ratios between $J_{\rm north}$ and $J_{\rm south}$ ($R_{\rm NS}$) and the results were averaged across the replicate trees for each species (Shinohara *et al.*, 2013). The coefficients of variability (CV) were estimated for individual trees and averaged for each species for the key sap flow parameters of sap flux density (J), whole-tree sap flow (V), nocturnal sap flow ($V_{\rm Night}$, see below), and the sap flux ratios ($R_{\rm NS}$) (Wang *et al.*, 2018).

Interspecific variability in sap flow.

Diel patterns in the sap flow of the three species were assessed for variability across seasons. Our assessment of interspecific variability was focused on the relative differences in sap flow along the radial and azimuthal profiles among the three species. Species-wise multiple linear regression models (MLRMs) were fitted for the day (06.00-18.00 h) and night (18.00-06.00 h) periods to assess the relative roles of the environmental variables, R_{sol} , VPD, and S in driving indices of radial (C_h) and azimuthal (R_{NS}) variability (Moore et al., 2011; Barbeta et al., 2012). Values of C_h above and below 1 suggest under- and overestimation, respectively, by the 2 cm probe compared to the rest of the xylem. Similarly, $R_{\rm NS}$ values above 1 suggest dominant flow on the north-facing part of the trunk, while values below 1 suggest dominant flow on the south side. Sap flow between 18.00–06.00 h was considered nocturnal ($V_{\rm night}$), and its percentage contribution to daily V was estimated. $V_{\rm night}$ was further divided into evening (18.00-00.00 h, V_{evening}) and pre-dawn (00.00-06.00 h, $V_{\text{pre-dawn}}$) components (Forster, 2014). Species-wise MLRMs using V_{night} as the response variable and VPD, S, and wind speed (u) as predictor variables were fitted to quantify the environmental drivers of nocturnal sap flow (Barbeta et al., 2012). In the MLRMs, VPD, u, and S were normalized by their respective daily means to facilitate the interpretation of the results.

Lag analysis between sap flow and SPAC variables.

Exploratory linear regression models yielded poor results for interactions between sap flow and environmental drivers, which were temporally lagged due to phase differences (Bond et al., 2002). Hence, lag correlation analysis was used to compute seasonal changes in the time lag between the environmental drivers and whole-tree sap flow (V) (Moore et al., 2011). The time-series for S was de-trended to extract the diel signals (S_{diu}) using the stlplus package in R. The hourly lag was computed using the ccf function in the stats package in R for each day between R_s versus V, VPD versus V, and $S_{\rm diu}$ versus V for each of the three species. The days with significant auto-correlation coefficients (ACF $\geq |0.4|$) were plotted for the three species. Positive ACF values signify that a high value of the driver variable is followed by a high value of the response variable after the corresponding lag hours. Conversely, negative ACF values signify that a high value of the driver variable is followed by a low value of the response variable. Similarly, positive lag hours indicate that the driver variable leads the response variable and that negative lag hours indicate that the driver variable lags behind the response variable.

GLS linear regression models for whole-tree sap flow.

The relative influences of environmental (predictor) variables on the whole-tree sap flow (response) variable were quantified using a combination of a temporal lag and generalized least-squares (GLS) regression models using the gls function in the nlme package in R with a suitable correlational structure. The GLS regression method estimates the maximum likelihood of the regression coefficients using generalized least-squares and is suitable for analysing time-series data with autocorrelational structures (Pinheiro and Bates, 2006; Krishnaswamy et al., 2012). The GLS model used R_s , VPD, and S as predictor variables, and they were normalized by subtracting the daily mean from the time-series and checked for collinearity. The second-order autoregressive moving average (corARMA) was found to be the best-fit correlational structure. The GLS model with corARMA structure was run separately for each tree to capture the interactions at both intra- and interspecific levels. Contiguous subsets longer than 5 d without missing values were tested for the time lag between the driver and response variables using the cof function in R. If any significant lag (ACF $\geq |0.4|$) was observed, the driver variable was lagged by a suitable range of hours to develop a multiple-lagged time-series starting from zero lag time, and a linear regression model was fitted with the response variable. The procedure was repeated for each of the driver variables, and the lagged time-series with the lowest lag hours ($P \le 0.05$) was chosen to develop a reconstituted dataset. The GLS model with corARMA structure was compared against a similar GLS model without any autocorrelation structure using ANOVA. The GLS model with the lowest Akaike information criterion and the highest likelihood ratio and significance was chosen for interpretations. The model performance was assessed using a linear regression model and the Kling-Gupta Efficiency (KGE) score for the observed and predicted values of V (Gupta and Kling, 2011).

Results

Intraspecific variability in sap flux density and sap flow

Radial variability in sap flux density and sap flow.

Radial patterns in J_{max} varied considerably across the three species. The pioneer species S. racemosa and E. acuminata both exhibited concave curves for I_{max} , with comparable peaks at the outer- and innermost xylem (37.4 \pm 15 cm³ cm⁻² h⁻¹ and $41.3 \pm 23 \text{ cm}^3 \text{ cm}^{-2} \text{ h}^{-1}$, respectively; Fig. 1A). Correspondingly, the percentage contribution of each depth to daily V declined uniformly from the outermost xylem inwards for these species (Fig. 1B). Conversely, the late-successional species C. hystrix showed highest J_{max} (23.1 \pm 14 cm³ cm⁻² h⁻¹) at the inner xylem (4 cm depth from the cambium) and its contribution to daily V was disproportionately higher (40%) than the rest of the depths. Following the above patterns, the highest wood moisture and lowest wood density were also observed at 1 cm depth in S. racemosa and E. acuminata, and at 4 cm depth in C. hystrix (Supplementary Fig. S3). Overall, mean hourly J increased with wood moisture and declined with wood density for a given sapwood annulus ring, except for E. acuminata (Supplementary Fig. S4). The relative contributions of sapwood

annuli to whole-tree sap flow were similar for both day and night periods and across the species. The innermost probe (5 cm depth) in C. hystrix stopped showing sap flux movement after January, indicating loss of conductivity in the annuli as the inner sapwood transitioned into non-functioning heartwood (Brodersen et al., 2019).

The three species displayed contrasting diel patterns in the variability of the radial correction factor, C_h (Supplementary Fig. S5A). In S. racemosa and E. acuminata, the 2 cm probe underestimated sap flow in relation to the rest of the xylem $(C_b>1)$ during the evening-night period (17.00–04.00 h) and overestimated it (C_h <1) during the day (04.00–1700 h). In contrast, in C. hystrix the 2-cm probe overestimated sap flow during the night-morning (23.00-09.00 h) and overestimated it during the noon-evening period (10.00–22.00 h). Closer examination indicated that the variability in C_h was driven by the higher sap-flux density at the outer xylem (1 cm depth) in S. racemosa and E. acuminata and at the inner xylem (4 cm depth) in C. hystrix compared to the rest of the xylem. The results of MLRMs suggested that daytime radial variability was strongly predicted by R_{sol} (negative slope) and S (positive slope) in S. racemosa (r^2 =0.43, P<0.001), whereas none of the predictors was significant for either E. acuminata ($r^2=0.01$, P<0.6) or C. hystrix (r^2 =0.01, P<0.15) (Supplementary Table S3). On the other hand, VPD was a strong negative predictor of night-time C_h in E. acuminata (r^2 =0.4, P<0.001), but not in S. racemosa $(r^2=0.02, P<0.2)$ or C. hystrix $(r^2=0.001, P<0.6)$.

Both S. racemosa and E. acuminata exhibited consistently high azimuthal sap flux ratios (R_{NS}) throughout the day (Supplementary Fig. S5B). Interestingly, in C. hystrix the northern probes dominated in the morning, with large standard deviations that suggested significant intraspecific variability. The southern probes showed peaks from the afternoon onwards in this species. The MLRMs predicting R_{NS} performed

poorly across the species ($r^2 \sim 0.03 - 0.18$; Supplementary Table S4). However, $R_{\rm sol}$ (positive slope) was a significant predictor of $R_{\rm NS}$ at moderate values of VPD and S across the three species (P<0.09). Night-time azimuthal variability was best predicted by VPD (P<0.06) and S (P<0.08) in S. racemosa, and by VPD in C. hystrix (P<0.004). S was a significant predictor of nighttime R_{NS} in E. acuminata (P < 0.04).

Intraspecific variability in sap flux density and sap flow.

The CV between the three species was very high for all sapflow parameters (J, 255%; V, 246%; V_{Night} , 147%; and R_{NS} , 236%). The CV in I was significantly higher among individuals of S. racemosa (282 \pm 32%) and E. acuminata (267 \pm 30%) than C. hystrix (179 \pm 12%). Similarly, the CV in V was significantly higher between individuals of S. racemosa (279 \pm 49%) and E. acuminata (273 \pm 22%) than C. hystrix (170 \pm 5%). The intraspecific variabilities in nocturnal sap flow in the pioneer species S. racemosa (155 \pm 32%) and E. acuminata (140 \pm 34%) were almost double that of the late-successional C. hystrix (78 \pm 3%). Conversely, the CV in azimuthal variability in C. hystrix (273 \pm 28%) was over twice that of S. racemosa $(114 \pm 56\%)$ and E. acuminata $(96 \pm 60\%)$.

Interspecific variability in sap flow and environmental drivers

Comparisons of sap-flux density and whole-tree sap flow.

The pioneer species S. racemosa and E. acuminata exhibited double the rates of sap-flux density (J) compared to the latesuccessional species C. hystrix (Table 3). The whole-tree sap flow (V) was linearly related to the total sapwood area ($r^2 > 0.71$, P < 0.001). The larger girth in C. hystrix was offset by a significant proportion of non-conducting heartwood (Table 2), whereas S. racemosa and E. acuminata showed a significantly

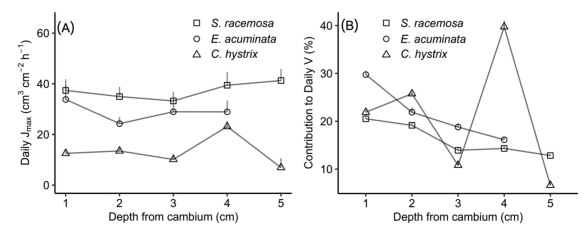


Fig. 1. Radial variability in sap flow in the pioneer species Symplocos racemosa and Eurya acuminata and the late-successional species Castanopsis hystrix measured in situ in an Eastern Himalayan forest. (A) Mean daily maximum sap-flux density (J_{max}, cm³ cm⁻² h⁻¹) and (B) percentage contribution of the sap flow at different depths to daily whole-tree sap flow (V, kg h⁻¹). Data are means (+SE), 56 d of measurements including overlapping sample days for Symplocos racemosa (Tree 1, n=16), Eurya acuminata (Tree 1, n=12) and Castanopsis hystrix (Tree 3, n=47).

Table 3. Estimates of sap flux density (*J*), whole-tree sap flow (*V*), and percentage biases in the estimation of *V* due to ignoring radial and azimuthal variability.

Species	J (cm³ cm ⁻² h ⁻¹)	<i>V</i> (kg h ⁻¹)	Radial bias (%)	Azimuthal bias (%)
S. racemosa	28 (2)	2.4 (1.2)	8 (4)	-5 (12)
E. acuminata	21 (10)	1.52 (0.8)	1 (3)	5 (12)
C. hystrix	13 (5)	2.39 (1.7)	8 (7)	-21 (41)

Data are means (±SD); 114 unique day replicates, tree-wise breakup of number of days is given in Supplementary Table S1B.

higher proportion of conducting sapwood, contributing to high V values. Radial bias, due to assumptions of homogenous flow along the xylem, led to overestimations of V in the three species. The failure to account for azimuthal variability (azimuthal bias) caused a significant underestimation of V in S. racemosa and C. hystrix but an overestimation in E. acuminata.

Diel patterns in sap flow in relation to environmental drivers.

The peak sap flowin S. racemosa and E. acuminata occurred relatively early in the day (07.00–08.00 h), whereas in C. hystrix it peaked around 12.00 h (Fig. 2; Supplementary Table S1A). Both S. racemosa and E. acuminata showed low sap flow at high values of $R_{\rm sol}$ and VPD (Supplementary Fig. S6). The heightened sensitivity to the environmental extremes manifested in bimodal peaks in diel sap flow with significant midday depression of flow across the trees for S. racemosa and E. acuminata. This depression coincided with peaks in VPD and $R_{\rm sol}$ and a trough in S. In contrast, C. hystrix maintained relatively high Vat high values of $R_{\rm sol}$ and VPD with unimodal peaks, although minor signs of a midday depression in sap flow were seen in two of the three trees.

Nocturnal sap flow.

Nocturnal sap flow (V_{night}) as a fraction of daily V was highest in E. acuminata (17.2 \pm 9%), followed by C. hystrix (13.5 \pm 5%) and S. racemosa (11.3 \pm 7%) (Table 4). A significant proportion of V_{night} occurred during the pre-dawn period in S. racemosa $(6.2 \pm 7\% \text{ of daily } V)$, whereas evening flux dominated in C. hystrix (9.7 \pm 5%). The proportion during the evening period (9.5 \pm 8%) was marginally higher than during pre-dawn $(7.7 \pm 7\%)$ in E. acuminata. The environmental conditions during the night fluctuated between high wind and high VPD to relatively quieter periods with low VPD. Across the species, high values of V_{night} were associated with low VPD (<0.1 kPa), low wind velocity (<0.5 m s⁻¹), and moderately saturated soil moisture conditions (Supplementary Fig. S7). The MLR models predicting V_{night} concurred with the observations that VPD, soil moisture, and wind acted as limiting variables to nocturnal sap flux (Supplementary Table S5). Soil moisture (negative slope) was a significant predictor of V_{night} in E. acuminata $(r^2=0.12, P<0.05)$ and VPD (negative slope) was a significant predictor for C. hystrix (r^2 =0.13, P<0.009). None of the predictors were significant for S. racemosa ($r^2=0.02$, P>0.5). Both evening and pre-dawn sap flow were independent of tree size in the three species.

Seasonal changes in diel patterns of environment, sap flow, and transpiration.

Changes in diel patterns of the SPAC variables as the growing season progressed from winter to summer are shown in Fig. 3. The winter months of December-February marked the dry season with sunny but cold days, sub-zero night temperatures, low VPD, and declining soil moisture. In March, rains replenished the soil moisture reserves and promoted leaf flush. The summer months of March-May saw increased day length, abundant moisture, warmer temperatures, and higher VPD. Consequently, V also increased from winter to summer, but with considerable variation across the three species (Fig. 3D-F). Both S. racemosa and E. acuminata exhibited peak sap flows in December and March, whereas C. hystrix had peaks in March and April. The intraspecific variability in sap flow also followed similar patterns. Radial variability in sap flux (differences in J_{max} between inner and outer xylem) increased in the dry season (December) for S. racemosa, whereas C. hystrix exhibited the highest variability in March and April (Supplementary Fig. S8). The fractional contribution of the outer xylem to daily V increased from December to March for *E. acuminata* and *C. hystrix*, whereas there were only marginal differences observed in S. racemosa (Supplementary Fig. S9). Sap flux ratios (R_{NS}) , the metric for azimuthal variability, also increased from winter to summer for S. racemosa, whereas in C. hystrix it was highest in January (Supplementary Fig. S10). The peak sap flow values increased as the growing season progressed, except for S. racemosa, and the daily timing of peak Vremained consistent across the species (Fig. 3D–F). The bimodal peaks in diel sap flow were consistent for S. racemosa and E. acuminata across the seasons, with higher rates in the morning than in the afternoon. As noted above, during the dry winter, a mild midday depression in sap flow was observed in two out of three trees of C. hystrix, resulting in a shift to unimodal peaks being observed from March onwards. Weak declining trends in $V_{
m night}$ were observed from the dry to the wet season in S. racemosa and C. hystrix Supplementary Fig. S11. The diel peaks in soil moisture (S_{din}) shifted from the afternoon to the evening and had increasing amplitude from winter to summer (Fig. 3C).

Environmental controls on transpiration

Lag analysis between transpiration and SPAC variables.

V consistently lagged behind solar radiation (R_{sol}) in all three species by an average of 1–2 h with positive correlations in and summer, with the lowest variability in C. hystrix (Fig. 4A,

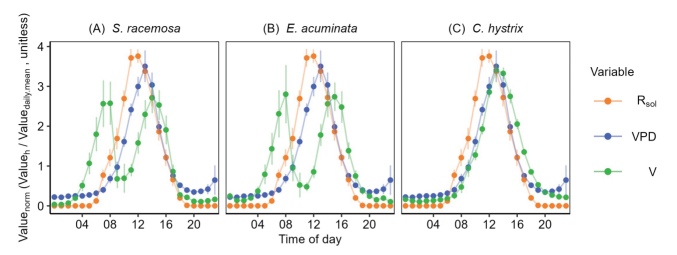


Fig. 2. Diel patterns of sap flow (V, kg h⁻¹) in relation to incoming short-wave radiation (R_{sol}, kW m⁻²) and vapour pressure deficit (VPD, kPa) in the pioneer species Symplocos racemosa and Eurya acuminata and the late-successional species Castanopsis hystrix measured in situ in an Eastern Himalayan forest. The data are normalized by their respective daily means, and are means (±SE), for the 28 concurrent data days which coincided for all species from total days for each species: Symplocos racemosa (Tree 1-5, 100 d), Eurya acuminata (Tree 1-5, 80 d) and Castanopsis hystrix (Tree 1-3, 65 d), a tree-wise breakup of n is given in Supplementary Table S1A.

Table 4. Percentage contributions of day, whole-night, evening, and pre-dawn sap flows to total daily sap flow in the three species.

Species	Day (06.00-17.00 h)	Night (18.00-05.00 h)	Evening (18.00-23.00 h)	Pre-dawn (00.00-05.00 h)
S. racemosa	88.7 (7)	11.3 (7)	5.1 (5)	6.2 (7)
E. acuminata	82.8 (9)	17.2 (9)	9.5 (8)	7.7 (7)
C. hystrix	86.5 (5)	13.5 (5)	9.7 (5)	3.8 (4)

Data are means (SD); 114 unique day replicates, tree-wise breakup of number of days is given in Supplementary Table S1B.

D, R_{sol} versus V). However, in winter, V preceded R_{sol} in S. racemosa and E. acuminata by an average of 1-2 h with positive correlations. In S. racemosa and E. acuminata, VPD lagged V in both seasons with positive correlations. In contrast, VPD led V in both seasons in C. hystrix (Fig. 4B, E, VPD versus V). On closer inspection, the days when V preceded R_{sol} and VPD with strong positive correlations were marked by significant pre-dawn flux, and the frequency of such days was significantly higher in S. racemosa and during the dry winter months of January and February (Fig. 2). Interactions between V and S_{diu} were more complicated (Fig. 4C, F, V versus $S_{\rm diu}$). On most days in winter (75% of total days), across the three species $V \log S_{\text{diu}}$ with an average lag of 2.2–7.5 h and positive correlations, while on the remaining days there were negative correlations and V lagged behind S_{diu} by an average lag of 2.2-4.1 h. However, the patterns were reversed from March onwards with the advent of rains, and on most days in summer (77% of total), S_{diu} led V by an average of 1.1-1.6 h with negative correlations in all three species, while on the remaining days S_{diu} lagged behind V by an average of 0.7–4 h and there were positive correlations. We also observed that the lag between transpiration and soil moisture was also positively correlated to tree size, with the larger-sized trees (>0.15 m DBH) experienced a smaller lag between transpiration and

soil moisture in the winter in S. racemosa and E. acuminata, and in the summer in C. hystrix, but the results were not significant.

GLS linear regression model for whole-tree sap flow.

The sap-flow modelling exercise was carried out for contiguous data periods (without missing values; Supplementary Fig. S2E-G) across winter and summer for the instrumented trees of S. racemosa (five periods, 8 ± 2 d), E. acuminata (four periods, 10 ± 5 d), and C. hystrix (10 periods, 10 ± 4 d) (Table 5). The GLS model with corARMA structure had significantly lower Akaike information criterion than the GLS model without any correlational structure. In summer, VPD (positive slope) emerged as a significant predictor for the three species of whole-tree sap flow (V) at average values of $R_{\rm sol}$ and soil moisture (S). In winter, R_{sol} was a co-driver of V for E. acuminata, and C. hystrix with positive coefficients, while S was a significant predictor of V in one of the trees of C. hystrix with a positive coefficient. Interestingly, at least one replicate tree from each species showed a negative $R_{\rm sol}$ coefficients in either the winter and/or summer, albeit with low significance (P>0.1), indicating the occasionally limiting role played by R_{sol} . S was not a significant predictor of V; however, it is important to note that soil moisture always remained well above the permanent

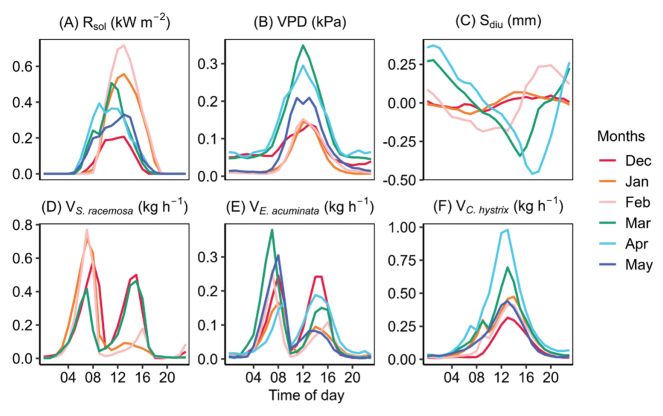


Fig. 3. Seasonal changes in diel patterns of environmental variables and whole-tree sap flow in the pioneer species *Symplocos racemosa* and *Eurya acuminata* and the late-successional species *Castanopsis hystrix* measured *in situ* in an Eastern Himalayan forest. (A) Incoming short-wave radiation (R_{sol}), (B) vapour pressure deficit (VPD), and (C) the diel component of soil moisture (S_{diu}). (D–F) The whole-tree sap-flow rates (V) of (D) *S. racemosa*, (E) *E. acuminata*, and (F) *C. hystrix*. Each line represents the monthly mean value. The sap-flow rates are based on 114 concurrent data days for the three species: *Symplocos racemosa* (Tree 1-5, 164 d), *Eurya acuminata* (Tree 1-5, 164 d) and *Castanopsis hystrix* (Tree 1-3, 206 d), a tree-wise breakup of *n* is given in Supplementary Table S1B.

wilting point of the site. The relative strength of interactions between V and the environmental drivers varied between individual trees of the same species and across the seasons.

In S. racemosa, VPD was the only significant predictor of Vin all the trees in summer (Table 5). In E. acuminata, VPD was the only significant predictor of V in one of the replicate trees in summer, while in another, VPD and R_{sol} were significant copredictors in winter. In C. hystrix, VPD and R_{sol} were significant co-predictors in six out of ten periods across both seasons with positive coefficients, except for one period in Tree 3 in summer where R_{sol} showed a negative coefficient. The GLS model performed relatively better for S. racemosa ($r^2=0.15$, P<0.001, KGE=0.12) than E. acuminata (r^2 =0.08, P<0.001, KGE=0.02) with improved concurrence between predicted and observed V. In both these species, the predicted V failed to replicate the bi-modal patterns in the observed V (Supplementary Fig. S12A, B). However, the model performance was significantly better for *C. hystrix* (r^2 =0.41, P<0.001, KGE=0.43), with the predicted V during the daytime matching well with the unimodal pattern (Supplementary Fig. 12C). However, the predicted V had sap flow in the night hours for all three species, which was absent in the observed V.

Discussion

Influence of secondary forest structure on sap flow variability in co-occurring species

The root-trunk-leaf connectivity in the xylem is a fascinating area of research under the soil-plant-atmosphere-continuum (SPAC), and it is largely dominated by the century-old cohesion-tension theory and piped model of water transport (Kim et al., 2014). The unit-pipe model postulates direct connections between the roots and lateral branches (known as sectoral flow) and is a common feature of ring-porous species. Conversely, diffuse-porous species exhibit considerable lateral connectivity (known as integrated flow), where multiple parts of the crown are supported by a cross-section of the xylem. Interestingly, pruning experiments show that, within the same individual, the outer xylem can exhibit sectoral connectivity, while the inner xylem can show an integrated flow pattern (Čermák and Nadezhdina, 2011; Dong et al., 2019). In this study, the observed Gaussian radial profile with increasing sap-flux density in the inner xylem in C. hystrix (Fig. 1), a ring-porous species, suggests sectoral connectivity, which is also typical of isolated trees with an extended

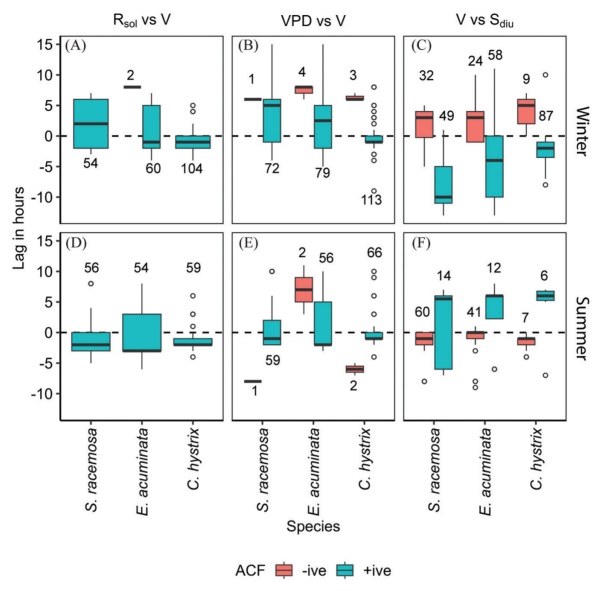


Fig. 4. Boxplots showing seasonal shifts in daily lag hours at maximum values of auto-correlation coefficients (ACFs) between environmental variables and whole-tree sap-flow rates of the pioneer species Symplocos racemosa and Eurya acuminata and the late-successional species Castanopsis hystrix measured in situ in an Eastern Himalayan forest. (A-C) Values recorded in winter and (D-F) values recorded in summer. (A, D) Incoming shortwave radiation (R_{sol}, kW m⁻²) versus whole-tree sap-flow rate (V, kg h⁻¹), (B, E) vapour pressure deficit (VPD, kPa) versus V, and (C, F) V versus the diel component of soil moisture (S_{diu}, mm). The colour of the boxplot represents the direction of the ACF values, as indicated, and the values adjacent to the boxes are the number of data points.

crown (Fiora and Cescatti, 2006). The Gaussian radial profile is ascribed to higher light availability to the older portions of the crown around the periphery, which are anatomically connected to the inner xylem. Conversely, in trees forming a continuous canopy, such as S. racemosa and E. acuminata (both diffuse-porous species), the activity of the lateral foliage is suppressed by shading, and only the top portion of the crown receives adequate light, inducing higher sap-flux density in the outer xylem. This was supported by the overestimation of radial correction factors ($C_h < 1$) that we observed in the outer xylem of the in-canopy pioneer species S. racemosa and E. acuminata, and underestimation $(C_h>1)$ in the outer xylem in the late-successional emergent trees of C. hystrix (Supplementary Fig. S5A). Delzon et al. (2004) observed that the radial correction factors were independent of meteorological factors and better represented the connectivity between xylem and crown-water use.

Landscape characteristics such as slope and aspect influence the diel changes in the angle and intensity of incident light, potentially inducing azimuthal variability in sap flow (Kumagai et al., 2007; Berry et al., 2016). Studies have shown greater connectivity between lateral branches (and leaves) and

fine-root development within the same aspect of a given tree (Čermák and Nadezhdina, 2011; Dong et al., 2019). Along the same lines, we observed that fully exposed crowns of *C. hystrix* exhibited shifting peaks in sap flow from north- to south-facing xylem following the north-east-south-west trajectory of sunlight, whereas the in-canopy *S. racemosa* and *E. acuminata* maintained dominant sap flow in the north-facing xylem (Supplementary Fig. 5B). Interestingly, the CV in azimuthal variability was almost double in the emergent trees of *C. hystrix* compared to the in-canopy pioneer species *S. racemosa* and *E. acuminata*. Daytime radial and azimuthal variability in the three species was attributable to the interaction between access to solar radiation and soil moisture, whereas night-time variability was negatively correlated with VPD (Supplementary Tables S2,

S3); Fiora and Cescatti, 2006; Shinohara *et al.*, 2013). The biases in estimating V due to ignoring azimuthal and radial variability were significant and comparable to existing literature (Fiora and Cescatti, 2006; Shinohara *et al.*, 2013).

Differential water-use strategies among pioneers and late-successional species

Functional groups play a major role in driving interspecific variability in transpiration (Nogueira et al., 2004; Motzer et al., 2005). Our observations of high sap-flux density (J) in the pioneer species S. racemosa and E. acuminata (Fig. 1) align well with a previous study of co-occurring tree species in a semi-arid region of China that showed high transpiration rates in pioneer

Table 5. Results from generalized least-squares linear regression models with corARMA correlational structure for predictor variables versus whole-tree sap flow.

Species	Tree	Season	No. of data days	Predictor variables	Coefficients	SE	Student's t	P
S. racemosa	Tree 2	Summer	9	(Intercept)	0.369	0.09	4.092	<0.001***
				S	0.014	0.067	0.204	0.839
				R_{sol}	-0.337	0.561	-0.601	0.548
				VPD	1.768	0.883	2.002	0.047*
	Tree 3	Winter	5	(Intercept)	1.164	0.447	2.605	0.01**
				S	-0.159	3.502	-0.045	0.964
				$R_{\rm sol}$	-0.41	1.715	-0.239	0.812
				VPD	6.832	7.653	0.893	0.374
		Summer	8	(Intercept)	0.564	0.058	9.703	<0.001***
				S	0.041	0.041	1.013	0.312
				R_{sol}	0.413	0.362	1.142	0.255
				VPD	3.481	0.554	6.283	<0.001***
	Tree 4	Summer	8	(Intercept)	0.255	0.052	4.905	<0.001***
				S	0.016	0.033	0.506	0.614
				$R_{\rm sol}$	0.014	0.223	0.061	0.952
				VPD	1.199	0.358	3.348	<0.001***
	Tree 5	Summer	8	(Intercept)	0.14	0.023	5.954	<0.001***
				S	0.009	0.016	0.586	0.559
				$R_{\rm sol}$	0.148	0.122	1.217	0.225
				VPD	0.619	0.189	3.282	<0.001***
E. acuminata	Tree 2	Summer	8	(Intercept)	0.146	0.025	5.901	<0.001***
				S	0.004	0.018	0.216	0.829
				R_{sol}	0.165	0.134	1.235	0.219
				VPD	0.643	0.204	3.16	0.002**
	Tree 3	Summer	9	(Intercept)	0.186	0.057	3.269	<0.001***
				S	0.001	0.043	0.016	0.987
				$R_{\rm sol}$	-0.052	0.36	-0.146	0.884
				VPD	0.297	0.562	0.528	0.598
	Tree 4	Winter	16	(Intercept)	0.342	0.045	7.606	<0.001***
				S	0.169	0.091	1.86	0.064
				R _{sol}	0.56	0.082	6.793	<0.001***
				VPD	3.128	0.268	11.669	<0.001***
		Summer	5	(Intercept)	0.869	0.474	1.836	0.069
				S	-0.01	0.317	-0.032	0.974
				R_{s}	-1.903	1.329	1.432	0.155
				VPD	0.901	2.984	0.302	0.763

Table 5. Continued

Species	Tree	Season	No. of data days	Predictor variables	Coefficients	SE	Student's t	P
C. hystrix	Tree 1	Winter	9	(Intercept)	0.48	0.128	3.745	<0.001***
				S	0.716	0.3	2.384	0.018*
				R _{sol}	3.18	0.347	9.165	<0.001***
				VPD	4.993	0.842	5.927	<0.001***
			16	(Intercept)	0.917	0.121	7.55	<0.001***
				S	-0.395	0.455	-0.868	0.386
				R _{sol}	1.59	0.365	4.358	<0.001***
				VPD	11.087	1.298	8.541	<0.001***
		Summer	13	(Intercept)	1.518	0.179	8.489	<0.001***
				S	0.1	0.063	1.591	0.113
				R _{sol}	2.443	0.417	5.857	<0.001***
				VPD	8.324	0.875	9.517	<0.001***
			10	(Intercept)	1.994	0.376	5.299	<0.001***
				S	0.105	0.164	0.639	0.523
				R_{sol}	0.3	0.44	0.684	0.495
				VPD	7.807	1.191	6.555	<0.001***
	Tree 2	Winter	9	(Intercept)	1.484	0.341	4.346	<0.001***
				S	0.75	1.359	0.552	0.581
				R_{sol}	2.1	1.411	1.488	0.138
				VPD	14.652	3.383	4.331	<0.001***
			16	(Intercept)	2.006	0.09	22.228	<0.001***
				S	-0.076	0.503	-0.152	0.879
				R_{sol}	5.618	0.364	15.417	<0.001***
				VPD	6.347	1.381	4.596	<0.001***
		Summer	7	(Intercept)	3.072	0.795	3.863	<0.001***
				S	0.174	0.706	0.246	0.806
				R_{sol}	-9.826	3.332	-2.949	0.004**
				VPD	21.676	6.44	3.366	0.001***
	Tree 3	Winter	5	(Intercept)	0.865	0.362	2.392	0.018*
				S	0.559	1.456	0.384	0.702
				R_{sol}	0.211	1.398	0.151	0.88
				VPD	4.181	3.218	1.299	0.196
			9	(Intercept)	0.725	0.072	10.015	<0.001***
				S	-0.065	0.18	-0.358	0.72
				R_{sol}	1.447	0.125	11.541	<0.001***
				VPD	5.374	0.408	13.171	<0.001***
		Summer	6	(Intercept)	1.248	0.312	4.007	<0.001***
				S	-0.141	0.199	-0.71	0.479
				R_{sol}	3.879	0.862	4.5	<0.001***
				VPD	0.52	1.757	0.296	0.768

Predictor variables: S, soil moisture (mm); R_{sol}, incoming short-wave radiation (kW m⁻²); VPD, vapour pressure deficit (kPa); significant values are highlighted in bold (*P<0.05, **P<0.01, ***P<0.001).

species, especially in the growing season (Lyu et al., 2020), and with a study from wet Brazilian rain forests, where Nogueira et al. (2004) reported pioneer species showing higher intrinsic water use efficiency (the ratio of net photosynthetic CO2 assimilation to stomatal conductance) and instantaneous transpiration efficiency (ratio of photosynthesis rate to transpiration) than late-successional species, while maintaining high sap-flux rates. Similarly, other studies have reported higher leaf-level hydraulic conductance and photosynthetic capacity in pioneer species than in late-successional species from South American tropical montane forests (TMFs; Sobrado, 2003; dos Santos et al., 2019) and from Central Panama (Bretfeld et al., 2018).

Like true pioneers, S. racemosa and E. acuminata are the first tree species to occupy any disturbance-related openings, but they also occur as sub-canopy associates to late-successional Fagaceae species in old-growth TMFs in the Sikkim region where our study was located, indicating long-term recruitment of pioneer species. Expanding the conventional segregation of species along the fast-slow/growth-survival axes, Rüger et al. (2020) suggested a third axis of stature-recruitment trade-offs

that distinguishes long-lived pioneers (fast-growing, longerlived, attaining tall canopy stature but low recruitment) from short-lived breeders (fast-growing, short-statured, low survival but a high number of offspring). In a meta-analysis of over 5000 species across 13 long-term monitoring plots, Kambach et al. (2022) showed that the fast-slow continuum and staturerecruitment are independent life-history strategies that shape tropical secondary forests globally. Although long-term studies are lacking from the landscape of our study, S. racemosa and E. acuminata can be characterized as long-lived pioneers (LLPs) owing to their ability to grow fast during the initial period after disturbance whilst also being able to adapt and live longer as the forest matures. LLPs are critical secondary forest species and dominate the intermediate successional stages in terms of carbon sequestration and water use (Rüger et al., 2020; Lai et al., 2021).

The pioneer species S. racemosa and E. acuminata also displayed increased radial and azimuthal variability in the summer (Supplementary Figs S8, S10) and consistent midday depression in *I* (Fig. 2), an indicator of efficient stomatal control over plant water use. Our GLS modelling results showed that at least one tree from each of the three species displayed a limiting effect of solar radiation (R_s) on V (i.e. negative coefficients; Table 5). The heightened sensitivity to environmental extremes in pioneer species can be attributed to their canopy position, which exposes them to strong levels of sunlight and VPD, causing midday stomatal closure (Huc et al., 1994; Franco and Lüttge, 2002; Chiariello et al., 2006). It is also attributable to their shallow-rootedness and sensitivity to diel and seasonal soil moisture fluctuations (Pavlis and Jeník, 2000), although the moisture observed in the topsoil (0-30 cm depth) in our study never went below the permanent wilting point. Bretfeld et al. (2018) quantified the impact of El Niño-induced drought years on sap flow in 76 species across different successional stages (8-80-year-old forests) in Central Panama and observed that the limiting role of seasonal soil moisture stress was stronger in pioneer species than in late-successional species. Similar reports from Central Himalayan species belonging to the Fagaceae, including Castanopsis indica, Quercus semicarpifolia, Q. leucotrichophora, and sub-canopy Rhododendron arboreum, have shown that seasonal soil moisture stress can induce a midday slump in leaf water potential (Zobel et al., 2001; Poudyal et al., 2004; Tewari et al., 2018). The absence of significant midday depression in deep-rooted species such as C. hystrix at the peak of the dry season and strong diel cycles in soil moisture provide evidence of vegetation accessing moisture from deeper soil layers (Tanaka et al., 2004; Allen, 2014; Kumar et al., 2022,

Our observations of 1–2 h of lag between R_{sol} and VPD and V were seasonally consistent (Fig. 4) and comparable with a previous report from a TMF in Costa Rica (Moore et al., 2018) and marginally higher than a reports from a TMF in Tibet (~1 h lag) (Wang et al., 2017). Since these are stem-based measurements, the lags could be confounded by delays between stem sap flow and canopy transpiration due to hydraulic capacitance and resistance, although such lags are usually ≤1 h (Phillips et al., 1997; Burgess and Dawson, 2008). The longer lags (>1 h) that we observed might also represent delays induced by low morning temperatures in winter and by leaf wetness in the summer due to night rains or dew (Aparecido et al., 2016; Bretfeld et al., 2018; Moore et al., 2018). Although rains in summer usually start around noon, significant rainfall also occurred in the pre-dawn period (Kumar et al., 2021). Hence, incorporating leaf wetness as a variable is recommended for future studies from wet TMFs such as those of Eastern Himalaya that experiences significant summer rains.

Plant water relations in wet and high-elevation Eastern Himalayan TMFs

High sap-flow rates in wet TMF climates.

The observed values of I in the three species in our study were significantly higher than their respective conspecifics from other TMFs. Thus, the *J* in *C. hystrix* at our site was found to be three times higher than conspecific C. tribuloides from the relatively drier Central Himalaya (MAP=1331 mm; Ghimire et al., 2014). Similarly, the values of J that we recorded for S. racemosa and E. acuminata were 6-9 times higher than conspecifics from drier environments, namely S. ramosissma in China (MAP ~2000 mm) and S. kuroki and E. japonica in Japan (MAP ~1700 mm) (Chiu et al., 2016; Zhang et al., 2018). Observations of higher sap-flux density and sap flow in pioneer species than in late-successional species have been previously reported from tropical lowland secondary forests of Costa Rica (40 masl), and high-elevation TMFs in Panama (1800 masl) and the Ecuadorian Andes (~2000 masl), and are commonly attributed to higher water movement potential in pioneer species (Motzer et al., 2005; McCulloh et al., 2011). The potential role of soil mycorrhizae and root hair densities in facilitating the high transpiration rates observed in these forests is another recommended area for future research (Carminati et al., 2017).

Nocturnal sap flow in Himalayan tree species.

The nocturnal sap flows that we observed constitute one of the first records for the pioneer species S. racemosa and E. acuminata globally and for C. hystrix from TMFs. For C. hystrix, previous reports from higher latitudes in China have been significantly lower (4% of daily *V*) than our observations (Chen et al., 2018; Wang et al., 2018). In contrast to a study by Wang et al. (2018), the observed intraspecific variability in nocturnal sap flow was less than the interspecific variation in late-successional C. hystrix However, in agreement with Kangur et al. (2021), higher intraspecific variability in nocturnal sap flow was seen in the pioneer species S. racemosa and E. acuminata, indicative of a broader range of responses to similar environmental stressors. According to the literature, the ecophysiological mechanisms driving nocturnal sap flux can be broadly categorized into refilling of the stem, facilitating night respiration, and endogenous stomatal controls. In our study, the majority of the nocturnal sap flux occurred in the evening (Table 3) and can be attributed to the capacitive refilling of the stem to capacity (Wang et al., 2018; Kamakura et al., 2021). In similar reports from broadleaved evergreen species in East Asia, nocturnal sap flux has been attributed to stem recharge and it has shown low correlations with environmental variables such as VPD (Chen et al., 2018; Wang et al., 2018; Zhang et al., 2020). Nocturnal movement of water to restore hydraulic equilibrium between roots, shoots, and leaves is a known feature of large trees (although not restricted to them), and in our study both evening and pre-dawn sap flux were independent of tree size (Zeppel et al., 2011; Siddig and Cao, 2018).

Interestingly, Chen et al. (2018) also reported significantly higher nocturnal sap flux in tree species with the capacity for corticular photosynthesis (C. hystrix being one of the common study species), indicating a role in facilitating respiration. They also highlighted the role of nocturnal sap flux in oxygen transport to the internal xylem during nocturnal respiration, a possible mechanism to alleviate hypoxia-like conditions created within internal photosynthesising xylem tissue during respiration. Marks and Lechowicz (2007) have further shown that nocturnal sap flux is closely linked to leaf nitrogen concentrations and higher rates of extension growth in fast-growing, shade-intolerant pioneer species and they attributed this to nocturnal respiration, which facilitates carbohydrate translocation and other processes associated with the growth of meristems in roots and shoots.

Lastly, the role of endogenous (circadian-driven) stomatal control in regulating nocturnal sap flux under conditions of stable VPD, low wind, and moderate soil moisture merits further investigations (Resco de Dios et al., 2013; De Dios et al., 2015). Under controlled conditions, Chu et al. (2009) demonstrated a significant positive correlation between wind speed and nocturnal sap flux up to a threshold when stomatal closure is assumed to limit transpiration. Similar to our observations, in a garden experiment involving temperate broadleaved species, Kangur et al. (2021) found significant nocturnal (pre-dawn) stomatal conductance in anisohydric pioneer species in comparison to isohydric late-successional species under hydrated conditions, and they attributed this to contrasting life-history traits dictating stomatal control. Kumar et al. (2021) observed a significant diel cycle in summer precipitation, starting from around noon and leaving a short 5-6 h window conducive to plant productivity and transpiration. Pre-dawn flux is an uncommon observation from the Himalaya and, as Barbeta et al. (2012) observed in the Mediterranean, could potentially be an adaptation to ensure hydraulic saturation in the stem, to avoid leaf-level moisture leakage, and to maximize photosynthetic CO₂ uptake at the start of the day. The prevalence of the phenomenon across both pioneer and primary tree species suggests that these traits might indeed be conserved as an adaptation and they merit further investigation (Kangur et al., 2021).

Energy versus moisture limitations on sap flow in Eastern Himalaya

Our study site experienced strong seasonal fluctuations in environmental conditions, with moisture- and energy-limited winters. The resulting seasonal variability in the interactive roles of sunlight and VPD as drivers of sap flow have been previously reported for TMFs (Moore et al., 2018). We observed that VPD was a stronger driver of sap flow than R_{sol} and S in both summer and winter, independent of functional groups (Fig. 4), which is in agreement with reports from Central Himalaya (Ghimire et al., 2014) and Central Panama (Bretfeld et al., 2018). However, during periods of sufficient soil moisture but limited energy availability in winter, sap flow was also driven by R_{sol} (Table 5), similar to observations from a TMF in Southern Andes (Motzer et al., 2010) and from the Alps (Fiora and Cescatti, 2006). As discussed above, we also found evidence of photosensitivity in the study species through the inhibitive impact of R_{sol} on sap flow, similar to reports from Brazilian TMFs (dos Santos et al., 2019). Soil moisture was both a limiting variable for pioneer species, as seen by the positive lags and negative autocorrelation coefficients, and a resource depleted by vegetation water use (Fig. 4C, F), something that particularly applies for the deep-rooted and large-sized individuals of Fagaceae (late-successional species) (Kumar et al., 2022, Preprint). Our results correspond to similar observations from Central Panama where older and large-sized late-successional species continue transpiring unperturbed even under drought conditions, while younger pioneer species reduce their water use (Bretfeld et al., 2018). In this respect, the effects of sapwood area (as a parameter of conducting capacity of the tree) and tree size (representing the age and form of the tree) on the ability of different species to withdraw water from different soil layers and possibly groundwater, becomes of particular interest. The linear increase in whole-tree sap flow with sapwood area is likely to plateau for mature individuals, even under saturated soil moisture conditions, due to limits imposed by stomatal conductance (Gao et al., 2015).

Our observations provide empirical evidence for the theory that the East Himalayan forests are energy-limited under abundant moisture conditions (Sebastian et al., 2019), although VPD remains the primary driver of sap flow independent of successional stages, and soil moisture plays a limiting role for pioneer species only (Bretfeld et al., 2018). Under such conditions, together with VPD, solar radiation could be a significant driver of non-monsoon vegetation productivity, as observed by Sebastian et al. (2019) using remotely-sensed enhanced vegetation index. The role of solar radiation is further accentuated in uneven-aged secondary forests in the Himalaya, where the positioning of the trees in the canopy determines the spatial availability of solar radiation to different parts of the crown (Fiora and Cescatti, 2006; Küppers et al., 2008; Zhang et al., 2019). The performance of our GLS regression model was affected by the inherent variability in sap flow, with complex

phenomena such as midday depression and pre-dawn sap flux dominating the diel patterns of the pioneer species at the site.

Implications of changing water availability on carbon sequestration in Himalayan TMFs

Climate change studies from Eastern Himalaya have highlighted an increase in mean annual temperature of 0.02 °C year⁻¹ with an estimate for daily mean temperature to increase by 1.8-4 °C by the end of the current century and precipitation to increase by 30-40% (Singh et al., 2011; Shrestha et al., 2012). Kulkarni et al. (2013) used a high-resolution PRECIS model to project an increase in temperature of up to 5°C and a 40% increase in precipitation by the year 2098 in Eastern Himalaya. In line with these regional trends, Dimri et al. (2018) used the Asian Precipitation-Highly Resolved Observational Data Integration Towards Evaluation of Water Resources (APHRODITE), a gauge-interpolated gridded precipitation product, to confirm increasing trends in pre-monsoonal (March-May) and monsoonal (June-September) precipitation in the Sikkim Himalaya region for 1970-2005. Sharma and Goyal (2020) used four global circulation models and an IMD 0.25° gridded precipitation dataset to project rises in maximum daily temperature of 0.4-1.1 °C and 0.5-1.7 °C and increases in mean annual precipitation of 220-380 mm and 150-760 mm under Representative Concentration Pathways (RCPs) 4.5 and RCP 8.5, respectively, for our study region for 2011–2100 (covered under SB6 basin in the paper). The projected rise in temperature is likely to increase both day and night-time VPD and lead to higher sap-flow rates, which we have already observed to be among the highest from tropical montane or lowland forests. The high sap-flow rates may be further increased by the predicted increases in summer and monsoon precipitation (Mcvicar et al., 2010). In winter, the reduced precipitation (low moisture availability) and increased warming (high VPD) may create seasonal drought conditions and induce water stress, especially among pioneer species.

Pioneer species are reported to have relatively high photosynthetic plasticity and leaf hydraulic conductivity that permit high sap-flux rates, while also being vulnerable to xylem embolism due to lower control over water use (Sobrado, 2003; Nogueira et al., 2004; dos Santos et al., 2019). Conversely, despite lower photosynthetic plasticity and lower hydraulic conductance, late-successional species are seemingly unaffected by seasonal drought conditions in winter due to better control of water use through stomatal closure and access to deeper moisture reserves (Sobrado, 2003; Tanaka et al., 2004; Allen, 2014). In addition, higher VPD is also likely to negatively affect nocturnal transpiration, which might in turn affect the ability of these tree species to utilize the available moisture efficiently. However, the net effect could be easily offset by the potential increase in daytime transpiration. The potential increase in annual net transpiration will most likely change the water balance by negatively affecting soil moisture and hydrological flows from these broadleaved TMFs in summers, and to a lesser degree in winters.

The impacts on carbon assimilation rates of projected increases in atmospheric CO₂ and associated fertilization effects together with increases in temperature and changes in precipitation will vary between species groups. The pioneer species are likely to maintain high photosynthesis rates by optimizing leaf-specific conductivity, which is the relationship between the conducting area and the total leaf area (Sobrado, 2003). Interestingly, Eckert et al. (2021) have shown that late-successional species have the highest rates of refixation of respiratory CO2 owing to relatively higher mesophyll resistance and thicker cell walls, which allows them to fix more CO2 with fewer open stomata and losing less water. Irrespective of the species groups, the CO₂ fertilization effect could induce lower moisture uptake as the same amount of carbon can be fixed with fewer stomata open or for less time (Mengis et al., 2015; Yang et al., 2016; Hashimoto et al., 2019). However, the gain in water-use efficiency with CO₂ fertilization could be offset by higher nocturnal transpiration in secondary forests (Marks and Lechowicz, 2007). Further, the CO₂ fertilization effect could be limited by the availability of soil nutrients, especially nitrogen and phosphorus, in poorly developed and highly leached mountain soils, such as at our study site (dos Santos et al., 2019; Terrer et al., 2019). Taken together, the postulated increase in productivity with increases in temperature and precipitation could lead to the greening of the TMFs in Sikkim Himalaya, at least in the short term (Sebastian et al., 2019). Similar evidence has already been reported from the wetter parts of eastern Nepal Himalaya, where climate warming and increased summer precipitation are likely to remove moisture constraints on photosynthesis in treeline conifer and broadleaved species (Pandey et al., 2020). Over longer time scales, the increase in productivity might not necessarily lead to higher carbon sequestration and net carbon gain. In a growth simulation study of 141 temperate tree species, Bugmann and Bigler (2011) showed that the higher productivity due to the CO₂ fertilization effect is associated with proportionately less biomass gain as well as reduced longevity, leading to net zero carbon gains.

Conclusions

It is still challenging to predict plant water relations in diverse secondary TMFs because of the variability among different functional groups and the diversity in the traits that regulate the relations (Huc et al., 1994; Nogueira et al., 2004; Küppers et al., 2008). In this study, using limited resources, the plant water relations in an East Himalayan wet tropical montane broadleaved forest have been estimated for the first time using methods covering the maximum variability of sap flow. Sap flow was found to be 3-9 times higher than in conspecific species in relatively drier regions of Central Himalaya and East Asia, highlighting the interactive role of precipitation and

elevation in modulating the available energy and moisture, and hence plant productivity (Bruijnzeel et al., 2011). Our use of radial and azimuthal probes has opened a whole new dimension of understanding of plant water use for Himalayan species, and their use is highly recommended to minimize the errors in estimating whole-tree sap flow (Shinohara et al., 2013; Komatsu et al., 2016). Our finding of significant nocturnal sap flow, dominated by the pre-dawn period, is the first report from the Himalaya and we postulate two potential roles that might have led to adaptation of pre-dawn sap flux in such wet, energy-limited montane systems: first, to facilitate carbon fixation during the cloud free, early morning hours by ensuring adequate leaf water balance and keeping stomata open, and second to ensure nutrient and O2 transport for early-morning developmental activity. Our observations of nocturnal sap flow hold significance for future SPAC work in terms of shifting the timing of pre-dawn leaf water potential measurements (Kavanagh et al., 2007), estimating water use efficiency across species (Chaves et al., 2016), quantifying hydrological services (Siddiq and Cao, 2018; Kangur et al., 2021), and improving regional climate change modelling exercises (De Dios et al., 2015). Other adaptations such as midday depression in shallow-rooted pioneer species highlight their sensitivity to environmental extremes (dos Santos et al., 2019).

Studies on climate change in the Eastern Himalaya have predicted an increase in summer rainfall, declining winter rains, and increasing summer and winter temperatures (Kulkarni et al., 2013; Krishnan et al., 2019). Drier and warmer winters are likely to affect the phenology and negatively affect the productivity of shallow-rooted pioneers such as S. racemosa and E. acuminata. In contrast, deep-rooted species such as C. hystrix and Himalayan oaks might remain unaffected by seasonal droughts (Kumar et al., 2022, Preprint). Thus, secondary broadleaved forests, with a dominance of shallow-rooted pioneers, are more prone to the negative impacts of drier and warmer winters than primary forests, which are dominated by deep-rooted species. In the summer, the onset of rain currently provides enough moisture to ensure peak vegetation productivity in April; however, increased summer precipitation in the future could result in higher cloud cover, negatively affecting both vegetation productivity and transpiration (Donohue et al., 2017), but it might also allow the opportunistic fast-growing pioneers to grow stronger alongside the latesuccessional species in the short term (Lyu et al., 2020). The CO₂ fertilization effect could potentially benefit both these species groups, but this could be limited by the availability of soil nutrients such as nitrogen and phosphorus, and any gains are likely to be offset by lower biomass gains and reduced longevity. The net effect of the postulated increase in productivity due to warming, CO₂ fertilization, and increased precipitation could lead to the greening of the TMFs in Sikkim Himalaya, at least in the short term (Krishnaswamy et al., 2014; Sebastian et al., 2019).

The overall effects of the changes in temperature and precipitation on the biodiversity of the region will be complex and their prediction will require ecohydrological models specific to the East Himalayan TMFs. More regional studies, including diel and seasonal variability in transpiration, will be critical for improving the accuracy of land-surface interaction models and predicting the impact of climate change on Himalayan ecohydrology (Wang and Dickinson, 2012; Miller et al., 2018). Our study provides the first empirical understanding of climatic controls on vegetation water use from a wet, high-elevation tropical broadleaved evergreen montane forest in Eastern Himalaya, one of the 30 global biodiversity hotspots and 200 ecoregions considered to be of importance (Chettri et al., 2008). Our GLS model highlighted the interacting roles of R_{sol} and VPD in modulating sap flow, and our study contributes to the currently scant literature on plant water relations from secondary TMFs, which have distinct water use behaviour in contrast to primary forests. Our study also highlights the pitfalls of generalizing water use across functional groups, forest types, and climatic conditions (Berdanier et al., 2016), whilst comparing plant water use strategies between co-occurring pioneer and late-successional species with a view to assessing the impacts of future changes in energy and soil moisture conditions (Bretfeld et al., 2018; Moore et al., 2018). Based on our findings, we recommend prioritizing the conservation and management of these secondary forests in the Eastern Himalaya for sustained ecosystem services.

Supplementary data

The following supplementary data are available at *JXB* online. Fig. S1. Tree densities of top seven species per hectare at the study site and their basal areas per hectare.

Fig. S2. Time-series plots of the raw daily data for the environmental variables and calculated sap flows of the three species during the study period.

Fig. S3. Variability in wood moisture and density with depth from the cambium in the three study species.

Fig. S4. Scatterplots of sap-flux density versus wood moisture and wood density for sample taken from the three species in December.

Fig. S5. Diel variability in sap flow across the three species as indicated by the correction factor C_h , the sap-flow ratio R_{NS} .

Fig. S6. Scatterplots of sap flow versus Incoming short-wave radiation and vapour pressure deficit for the three species.

Fig. S7. Scatterplots of nocturnal sap flow versus vapour pressure deficit, soil moisture, and wind velocity for the three species.

Fig. S8. Seasonal variations in daily maximum sap-flux density with depth of the xylem for the three species.

Fig. S9. Boxplots of the seasonal variation in percentage contribution of sap flow at different sapwood depths to daily whole-tree sap flow for the three species.

Fig. S10. Boxplots showing monthly variability in R_{NS} as a parameter of azimuthal variability in sap flux density.

Fig. S11. Monthly variability in the contribution of nocturnal sap flow to daily sap flow.

Fig. S12. Comparison of diel patterns between observed whole-tree sap flow and predictions for the three species based on GLS linear regression models.

Table. S1. Slopes and r^2 values of the linear regression models without intercept developed for estimating sap flow in replicate trees using Equation 3.

Table. S2. Results from multiple linear regression models for the day and night periods to assess the drivers of radial variability in sap flow across the three species.

Table. S3. Results from multiple linear regression models for the day and night periods to assess the drivers of azimuthal variability across the three species.

Table. S4. Results from multiple linear regression models to assess the drivers of nocturnal sap flow across the three species.

Acknowledgements

The authors thank the Department of Forests, Environment and Wildlife Management, Rural Management and Development Department, the Home Department, Govt. of Sikkim, India, and the Indian Army for research permits and field support in Sikkim. The study would not have been possible without crucial field support from Mr Passang Tamang, Mr Girish Varma, Mr Naresh Rai, and the staff of the Fambong-Loh Wildlife Sanctuary, East Sikkim. We also thank Dr Nachiket Kelkar and Dr Aniruddha Marathe for their useful comments on the statistical analysis, and Mr Lalit Kumar Rai and Dr Radhika Kanade for their valuable insights on forest ecology. We are indebted to Prof. Nathan G. Phillips at Boston University, USA, Prof. Frederick Meinzer at Oregon State University, USA, the Forests Science Laboratory (Corvallis), USA, and the Center for Embedded Network Sensing, University of California, USA, for generously sharing the original design of the thermal dissipation probe assembly. We thank Mr Mayank Singh for designing the graphical abstract. We acknowledge the logistical support provided by the ATREE Regional Office, Gangtok, during the study.

Author contributions

MK, GJ, and JK: conceptualization and methodology; MK: manufacturing of instruments, developing software, formal analysis and visualization of the data, and writing the original draft; MK and GJ: testing and modification of the sap-flow probes; MK and YB: performing the experiments; YB, GJ, and JK: reviewing and editing the manuscript; JK: supervision, project administration, and funding acquisition.

Conflict of interest

The authors declare that they have no conflicts of interest in relation to this work.

Funding

This work was supported by grants from the Department of Biotechnology, Govt. of India (GoI) (grant no. BT/01/NE/PS/NCBS/09) and the National Mission for Himalayan Studies (NMHS),

Ministry of Environment, Forest and Climate Change, GoI (grant no. GBPI/NMHS/HF/RA/2015-16/).

Data availability

The data supporting the findings of this study are available from the corresponding author, Manish Kumar, on request. A sample subset of the data supporting the findings of this study is also available in the Dryad Digital Repository at https://doi.org/10.5061/dryad.47d7wm3cg; Kumar et al. (2023).

References

Allen MF. 2014. How oaks respond to water limitation. In: Standiford RB, Purcell KL, eds. Proceedings of the seventh California oak symposium: managing oak woodlands in a dynamic world. General Technical Report PSW-GTR-251. Berkeley, CA: USDA Forest Service, Pacific Southwest Research Station, 13–22.

Aparecido LMT, Miller GR, Cahill AT, Moore GW. 2016. Comparison of tree transpiration under wet and dry canopy conditions in a Costa Rican premontane tropical forest. Hydrological Processes **30**, 5000–5011.

Asbjornsen H, Goldsmith GR, Alvarado-Barrientos MS, et al. 2011. Ecohydrological advances and applications in plant–water relations research: a review. Journal of Plant Ecology **4**, 3–22.

Barbeta A, Ogaya R, Peñuelas J. 2012. Comparative study of diurnal and nocturnal sap flow of *Quercus ilex* and *Phillyrea latifolia* in a Mediterranean holm oak forest in Prades (Catalonia, NE Spain). Trees **26**, 1651–1659.

Basnett S, Devy SM. 2021. Phenology determines leaf functional traits across *Rhododendron* species in the Sikkim Himalaya. Alpine Botany **131**, 63–72.

Berdanier AB, Miniat CF, Clark JS. 2016. Predictive models for radial sap flux variation in coniferous, diffuse-porous and ring-porous temperate trees. Tree Physiology **36**, 932–941.

Berry ZC, Gotsch SG, Holwerda F, Muñoz-Villers LE, Asbjornsen H. 2016. Slope position influences vegetation—atmosphere interactions in a tropical montane cloud forest. Agricultural and Forest Meteorology **221**, 207–218.

Berry ZC, Looker N, Holwerda F, Gómez Aguilar LR, Ortiz Colin P, González Martínez T, Asbjornsen H. 2018. Why size matters: the interactive influences of tree diameter distribution and sap flow parameters on upscaled transpiration. Tree Physiology **38**, 264–275.

Bhutia Y, Gudasalamani R, Ganesan R, Saha S. 2019. Assessing forest structure and composition along the altitudinal gradient in the state of Sikkim, Eastern Himalayas, India. Forests **10**, 6331–6317.

Bond BJ, Jones J, Moore G, Phillips N, Post D, McDonnell JJ. 2002. The zone of vegetation influence on baseflow revealed by diel patterns of streamflow and vegetation water use in a headwater basin. Hydrological Processes **16**, 1671–1677.

Bretfeld M, Ewers BE, Hall JS. 2018. Plant water use responses along secondary forest succession during the 2015–2016 El Niño drought in Panama. New Phytologist **219**, 885–899.

Brodersen CR, Roddy AB, Wason JW, McElrone AJ. 2019. Functional status of xylem through time. Annual Review of Plant Biology **70**, 4071–4433.

Brown S, Lugo AE. 1990. Tropical secondary forests. Journal of Tropical Ecology **6**, 1–32.

Bruijnzeel LA, Mulligan M, Scatena FN. 2011. Hydrometeorology of tropical montane cloud forests: emerging patterns. Hydrological Processes **25**, 465–498.

Bugmann H, Bigler C. 2011. Will the CO_2 fertilization effect in forests be offset by reduced tree longevity? Oecologia **165**, 533–544.

Burgess SSO, Dawson TE. 2008. Using branch and basal trunk sap flow measurements to estimate whole-plant water capacitance: a caution. Plant and Soil **305**, 5–13.

- Carminati A, Passioura JB, Zarebanadkouki M, Ahmed MA, Ryan PR, Watt M. Delhaize E. 2017. Root hairs enable high transpiration rates in drying soils. New Phytologist 216, 771-781.
- Čermák J. Nadezhdina N. 2011. Instrumental approaches for studving tree-water relations along gradients of tree size and forest age. In: Meinzer F, Lachenbruch B, Dawson T. eds. Size- and age-related changes in tree structure and function. Tree physiology, vol. 4. Dordrecht: Springer, 385-426.
- Chan AM, Bowling DR. 2017. Assessing the thermal dissipation sap flux density method for monitoring cold season water transport in seasonally snow-covered forests. Tree Physiology 37, 984-995.
- Chand DB, Poudyal K, Jha PK. 2017. Shifts in leaf phenology of three Himalayan oak species: role of wood water properties. Ecoprint: an International Journal of Ecology 24, 29-36.
- Chave J. 2006. Measuring wood density for tropical forest trees: a field manual. Sixth framework programme (2002–2006) PAN-AMAZONIA Project, European Commission.
- 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.
- Chen X, Gao J, Zhao P, McCarthy HR, Zhu L, Ni G, Ouyang L. 2018. Tree species with photosynthetic stems have greater nighttime sap flux. Frontiers in Plant Science 9, 30,
- Chettri N, Sharma E, Deb DC, Sundriyal RC. 2002. Impact of firewood extraction on tree structure, regeneration and woody biomass productivity in a trekking corridor of the Sikkim Himalaya. Mountain Research and Development
- Chettri, N., Shakya, B., Thapa, R., & Sharma, E. (2008). Status of a protected area system in the Hindu Kush-Himalayas: An analysis of PA coverage. The International Journal of Biodiversity Science and Management, 4(3), 164–178.
- Chiariello NR, Field CB, Mooney HA. 2006. Midday wilting in a tropical pioneer tree. Functional Ecology 1, 3-11.
- Chiu CW, Komatsu H, Katayama A, Otsuki K. 2016. Scaling-up from tree to stand transpiration for a warm-temperate multi-specific broadleaved forest with a wide variation in stem diameter. Journal of Forest Research 21, 161-169.
- Chu CR, Hsieh CI, Wu SY, Phillips NG. 2009. Transient response of sap flow to wind speed. Journal of Experimental Botany 60, 249-255.
- Dahal N, Lamichhaney S, Kumar S. 2021. Climate change impacts on Himalayan biodiversity: evidence-based perception and current approaches to evaluate threats under climate change. Journal of the Indian Institute of Science **101**. 195-210.
- Daniel D, Anandhi A, Sen S. 2021. Conceptual model for the vulnerability assessment of springs in the Indian Himalayas. Climate 9, 121.
- Davis TW, Kuo C-M, Liang X, Yu P-S. 2012. Sap flow sensors: construction, quality control and comparison. Sensors 12, 954-971.
- Delzon S, Sartore M, Granier A, Loustau D. 2004. Radial profiles of sap flow with increasing tree size in maritime pine. Tree Physiology 24, 1285–1293.
- Dimri AP, Kumar D, Srivastava M. 2018. Regional climate changes over Northeast India: present and future. In Singh A, Punia M, Haran N, Singh T. eds. Development and disaster management. Singapore: Palgrave Macmillan, 41-63
- De Dios VR, Roy J, Ferrio JP, Alday JG, Landais D, Milcu A, Gessler A. 2015. Processes driving nocturnal transpiration and implications for estimating land evapotranspiration. Scientific Reports 5, 10975.
- Dong T, Duan B, Korpelainen H, Niinemets U, Li C. 2019. Asymmetric pruning reveals how organ connectivity alters the functional balance between leaves and roots of Chinese fir. Journal of Experimental Botany 70, 1941–1953.
- Donohue RJ, Roderick ML, McVicar TR. 2007. On the importance of including vegetation dynamics in Budyko's hydrological model. Hydrology and Earth System Sciences 11, 983-995.
- Donohue RJ, Roderick ML, McVicar TR, Yang Y. 2017. A simple hypothesis of how leaf and canopy-level transpiration and assimilation respond to elevated CO₂ reveals distinct response patterns between disturbed and undisturbed vegetation. Journal of Geophysical Research, Biogeosciences 122,
- dos Santos OdO, Mendes KR, Martins SVC, Batista-Silva W, dos Santos MA, de Figueirôa JM, de Souza ER, Fernandes D, Araújo WL, Pompelli MF. 2019. Physiological parameters and plasticity as key factors to

- understand pioneer and late successional species in the Atlantic Rainforest. Acta Physiologiae Plantarum 41. 145.
- Eckert D, Martens HJ, Gu L, Jensen AM. 2021. CO2 refixation is higher in leaves of woody species with high mesophyll and stomatal resistances to CO₂ diffusion. Tree Physiology 41, 1450-1461.
- Fiora A, Cescatti A. 2006. Diurnal and seasonal variability in radial distribution of sap flux density: implications for estimating stand transpiration. Tree Physiology 26, 1217-1225.
- Flo V, Martinez-Vilalta J, Steppe K, Schuldt B, Poyatos R. 2019. A synthesis of bias and uncertainty in sap flow methods. Agricultural and Forest Meteorology 271, 362-374.
- Forrester DI. 2015. Transpiration and water-use efficiency in mixed-species forests versus monocultures: effects of tree size, stand density and season. Tree Physiology **35**, 289–304.
- Forster MA. 2014. How significant is nocturnal sap flow? Tree Physiology 34. 757-765
- Franco AC. Lüttge U. 2002. Midday depression in savanna trees: coordinated adjustments in photochemical efficiency, photorespiration, CO₂ assimilation and water use efficiency. Oecologia 131, 356-365.
- Gao J, Zhao P, Shen W, Niu J, Zhu L, Ni G. 2015. Biophysical limits to responses of water flux to vapor pressure deficit in seven tree species with contrasting land use regimes. Agricultural and Forest Meteorology 200, 258–269.
- Ghimire CP, Lubczynski MW, Bruijnzeel LA, Chavarro-Rincón D. 2014. Transpiration and canopy conductance of two contrasting forest types in the Lesser Himalaya of Central Nepal. Agricultural and Forest Meteorology 197,
- Granier A. 1987. Evaluation of transpiration in a Douglas-fir stand by means of sap flow measurements. Tree Physiology 3, 309-320.
- Gupta HV, Kling H. 2011. On typical range, sensitivity, and normalization of mean squared error and Nash-Sutcliffe efficiency type metrics. Water Resources Research 47. W10601.
- Gurung S, Chettri A. 2019. Vegetation analysis of Oak Forests of Fambong Iho Wildlife Sanctuary in Sikkim Himalayas. International Journal of Basic and Applied Biology 6, 192-197.
- Guyot A, Ostergaard KT, Fan J, Santini NS, Lockington DA. 2015. Xylem hydraulic properties in subtropical coniferous trees influence radial patterns of sap flow: implications for whole tree transpiration estimates using sap flow sensors. Trees 29, 961-972.
- Hashimoto H. Nemani RR. Bala G. et al. 2019. Constraints to vegetation growth reduced by region-specific changes in seasonal climate. Climate 7, 27.
- Hatton TJ. Catchpole EA. Vertessy RA. 1990. Integration of sap flow velocity to estimate plant water use. Tree Physiology 6, 201-209.
- Heimann M. Reichstein M. 2008. Terrestrial ecosystem carbon dynamics and climate feedbacks. Nature 451, 289-292.
- Hernandez-Santana V, Hernandez-Hernandez A, Vadeboncoeur MA, Asbjornsen H. 2015. Scaling from single-point sap velocity measurements to stand transpiration in a multispecies deciduous forest: uncertainty sources, stand structure effect, and future scenarios. Canadian Journal of Forest Research 45,
- Huc R, Ferhi A, Guehl JM. 1994. Pioneer and late stage tropical rainforest tree species (French Guiana) growing under common conditions differ in leaf gas exchange regulation, carbon isotope discrimination and leaf water potential. Oecologia 99, 297-305.
- James S, Clearwater MJ, Meinzer FC, Goldstein G. 2002. Heat dissipation sensors of variable length for the measurement of sap flow in trees with deep sapwood. Tree Physiology 22, 277-283.
- Kamakura M, Tsuruta K, Azuma WA, Kosugi Y. 2021. Hydraulic architecture and internal water storage of Japanese cypress using measurements of sap flow and water potential. Ecohydrology 14, e2325.
- Kambach S, Condit R, Aguilar S, et al. 2022. Consistency of demographic trade-offs across 13 (sub)tropical forests. Journal of Ecology 110, 1485–1496.
- Kanade R, John R. 2018. Topographical influence on recent deforestation and degradation in the Sikkim Himalaya in India; implications for conservation of East Himalayan broadleaf forest. Applied Geography 92, 85-93.
- Kangur O, Steppe K, Schreel JDM, Von Der Crone JS, Sellin A. 2021. Variation in nocturnal stomatal conductance and development of predawn

disequilibrium between soil and leaf water potentials in nine temperate deciduous tree species. Functional Plant Biology 48, 483-492.

Kannenberg SA, Guo JS, Novick KA, Anderegg WRL, 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.

Kavanagh KL, Pangle R, Schotzko AD. 2007. Nocturnal transpiration causing disequilibrium between soil and stem predawn water potential in mixed conifer forests of Idaho. Tree Physiology 27, 621-629.

Kim HK, Park J, Hwang I. 2014. Investigating water transport through the xylem network in vascular plants. Journal of Experimental Botany 65, 1895-1904

Komatsu H. Shinohara Y. Kume T. Tsuruta K. Otsuki K. 2016. Does measuring azimuthal variations in sap flux lead to more reliable stand transpiration estimates? Hydrological Processes 30, 2129-2137.

Krishnan R, Shrestha AB, Ren G, et al. 2019. Unravelling climate change in the Hindu Kush Himalaya: rapid warming in the mountains and increasing extremes. In: Wester P, Mishra A, Mukherji A, Shrestha A. eds. The Hindu Kush Himalaya assessment. Cham, Switzerland: Springer International Publishing,

Krishnaswamy J, Bonell M, Venkatesh B, Purandara BK, Lele S, Kiran MC. Reddy V. Badiger S. Rakesh KN. 2012. The rain-runoff response of tropical humid forest ecosystems to use and reforestation in the Western Ghats of India. Journal of Hydrology 472-473, 216-237.

Krishnaswamy J, John R, Joseph S. 2014. Consistent response of vegetation dynamics to recent climate change in tropical mountain regions. Global Change Biology 20, 203-215.

Kulkarni A, Patwardhan S, Kumar KK, Ashok K, Krishnan R. 2013. Projected climate change in the Hindu Kush-Himalayan region by using the high-resolution regional climate model PRECIS. Mountain Research and Development 33, 142-151.

Kumagai T, Aoki S, Shimizu T, Otsuki K. 2007. Sap flow estimates of stand transpiration at two slope positions in a Japanese cedar forest watershed. Tree Physiology 27, 161-168.

Kumar M. Bhutia Y. Joseph G. Krishnaswamv J. 2022. Transpiration drives diurnal and seasonal streamflow in secondary tropical montane forests of Eastern Himalaya. Authorea. doi: 10.22541/au.167048830.02473076/v1.

Kumar M, Hodnebrog O, Sophie Daloz A, Sen S, Badiger S, Krishnaswamy J. 2021. Measuring precipitation in Eastern Himalaya: ground validation of eleven satellite, model and gauge interpolated gridded products. Journal of Hydrology 599, 126252.

Kumar M, Joseph G, Bhutia Y, and Krishnaswamy J. 2023. Data from: Contrasting sap flow characteristics between pioneer and late-successional tree species in secondary tropical montane forests of Eastern Himalaya, India. [Dataset]. Dryad. https://doi.org/10.5061/dryad.47d7wm3cg

Kumar M, Rathod R, Mukherji A. 2023. Water security and spring conservation in the Himalaya. In Ojha H, Schofield N, Camkin J. eds. Climate risks to water security. Framing effective response in Asia and the Pacific. Palgrave studies in climate resilient societies. Singapore: Palgrave Macmillan, 15-36.

Küppers M, Motzer T, Schmitt D, Ohlemacher C, Zimmermann R, Horna V, Küppers BIL, Mette T. 2008. Stand structure, transpiration responses in trees and vines and stand transpiration of different forest types within the mountain rainforest. In: Beck E, Bendix J, Kottke I, Makeschin F, Mosandl R. eds. Gradients in a tropical mountain ecosystem of Ecuador. Ecological studies, vol. 198. Berlin, Heidelberg: Springer, 243-258.

Lai HR, Craven D, Hall JS, Hui FKC, van Breugel M. 2021. Successional syndromes of saplings in tropical secondary forests emerge from environment-dependent trait-demography relationships. Ecology Letters 24, 1776-1787.

Li Y, Liu J, Chen G, Zhou G, Huang W, Yin G, Zhang D, Li Y. 2013. Wateruse efficiency of four native trees under CO₂ enrichment and N addition in subtropical model forest ecosystems. Journal of Plant Ecology 8, 411-419.

Lu P, Urban L, Ping Z. 2004. Granier's thermal dissipation probe (TDP) method for measuring sap flow in trees: theory and practice. Acta Botanica Sinica 46. 631-646

Lvu J. He QY, Yang J. Chen QW, Cheng RR, Yan MJ, Yamanaka N, Du S. 2020. Sap flow characteristics in growing and non-growing seasons in three tree species in the semiarid Loess Plateau region of China. Trees 34, 943-955.

Manzoni S, Vico G, Porporato A, Katul G. 2013. Biological constraints on water transport in the soil-plant-atmosphere system. Advances in Water Resources 51, 292-304.

Marks CO, Lechowicz M. 2007. The ecological and functional correlates of nocturnal transpiration. Tree Physiology 27, 577-584.

McCulloh KA, Meinzer FC, Sperry JS, Lachenbruch B, Voelker SL, Woodruff DR. Domec JC. 2011. Comparative hydraulic architecture of tropical tree species representing a range of successional stages and wood density. Oecologia 167, 27-37.

McVicar TR, Donohue RJ, Grady APO, Li L. 2010. The effects of climatic changes on plant physiological and catchment ecohydrological processes in the high-rainfall catchments of the Murray-Darling Basin: a scoping study. CSIRO: Water for a Healthy Country National Research Flagship. Canberra: Murray-Darling Basin Authority.

Mencuccini M, Manzoni S, Christoffersen B. 2019. Modelling water fluxes in plants: from tissues to biosphere. New Phytologist 222, 1207-1222.

Mengis N, Keller DP, Eby M, Oschlies A. 2015. Uncertainty in the response of transpiration to CO₂ and implications for climate change. Environmental Research Letters 10, 094001.

Miller G. Cahill T. Moore G. Leung R. 2018. Improving land-surface modeling of evapotranspiration processes in tropical forests. Report no. DOE-TAMU-10654. College Station: Texas A&M Engineering Experiment Station.

Moore GW, Jones JA, Bond BJ. 2011. How soil moisture mediates the influence of transpiration on streamflow at hourly to interannual scales in a forested catchment. Hydrological Processes 25, 3701-3710.

Moore GW, Orozco G, Aparecido LMT, Miller GR. 2018. Upscaling transpiration in diverse forests: insights from a tropical premontane site. Ecohydrology 11, e1920.

Motzer T, Munz N, Anhuf D, Küppers M. 2010. Transpiration and microclimate of a tropical montane rain forest, southern Ecuador. In: Bruijnzeel LA, Scatena FN, Hamilton HLS. eds. Tropical montane cloud forests: science for conservation and management. Cambridge, UK: Cambridge University Press, 447-455.

Motzer T, Munz N, Küppers M, Schmitt D, Anhuf D. 2005. Stomatal conductance, transpiration and sap flow of tropical montane rain forest trees in the southern Ecuadorian Andes. Tree Physiology 25, 1283-1293.

Nachabe M, Shah N, Ross M, Vomacka J. 2005. Evapotranspiration of two vegetation covers in a shallow water table environment. Soil Science Society of America Journal 69, 492-499.

Nogueira A, Martinez CA, Ferreira LL, Prado CHBA. 2004. Photosynthesis and water use efficiency in twenty tropical tree species of differing succession status in a Brazilian reforestation. Photosynthetica 42, 351–356.

O'Brien JJ. Oberbauer SF. Clark DB. 2004. Whole tree xvlem sap flow responses to multiple environmental variables in a wet tropical forest. Plant, Cell & Environment 27, 551-567.

Ohsawa M. 1993. Latitudinal pattern of mountain vegetation zonation in southern and eastern Asia. Journal of Vegetation Science 4, 13-18.

Ohsawa M, Shakya PR, Numata M. 1986. Distribution and succession of West Himalayan forest types in the eastern part of the Nepal Himalaya. Mountain Research & Development 6, 143–157.

Pandey S, Cherubini P, Saurer M, Carrer M, Petit G. 2020. Effects of climate change on treeline trees in Sagarmatha (Mt. Everest, Central Himalaya). Journal of Vegetation Science 31, 1144-1153.

Paudel I, Kanety T, Cohen S. 2013. Inactive xylem can explain differences in calibration factors for thermal dissipation probe sap flow measurements. Tree Physiology **33**, 986–1001.

Pausch RC, Grote EE, Dawson TE. 2000. Estimating water use by sugar maple trees: considerations when using heat-pulse methods in trees with deep functional sapwood. Tree Physiology 20, 217-227.

- Pavlis J, Jeník J. 2000. Roots of pioneer trees in Amazonian rain forest. Trees **14**. 442–455.
- Peters RL, Fonti P, Frank DC, et al. 2018. Quantification of uncertainties in conifer sap flow measured with the thermal dissipation method. New Phytologist 219, 1283-1299
- Phillips N, Nagchaudhuri A, Oren R, Katul G. 1997. Time constant for water transport in lobiolly pine trees estimated from time series of evaporative demand and stem sapflow. Trees 11, 412-419.
- Phillips N, Oren R, Zimmermann R. 1996. Radial patterns of xylem sap flow in non-, diffuse- and ring-porous tree species. Plant, Cell & Environment 19, 983-990.
- Pinheiro JC, Bates D. 2006. Mixed-effects models in S and S-PLUS. New York: Springer Science & Business Media, 250-270.
- Poudyal K, Jha PK, Zobel DB, Thapa CB. 2004. Patterns of leaf conductance and water potential of five Himalayan tree species. Tree Physiology 24,
- Poyatos R, Granda V, Flo V, et al.. 2021. Global transpiration data from sap flow measurements: the SAPFLUXNET database. Earth System Science Data **13**, 2607-2649.
- Rabbel I, Diekkrüger B, Voigt H, Neuwirth B. 2016. Comparing ΔT_{max} determination approaches for granier-based sapflow estimations. Sensor 16, 2042.
- Ramakrishnan PS. Kushwaha SPS. 2001. Secondary forests of the Himalaya with emphasis on the north-eastern hill region of India. Journal of Tropical Forest Science 13, 727-747.
- Resco de Dios V, Díaz-Sierra R, Goulden ML, Barton CVM, Boer MM, Gessler A, Ferrio JP, Pfautsch S, Tissue DT. 2013. Woody clockworks: circadian regulation of night-time water use in Eucalyptus globulus. New Phytologist 200, 743-752.
- Rüger N, Condit R, Dent DH, DeWalt SJ, Hubbell SP, Lichstein JW, Lopez OR, Wirth C, Farrior CE. 2020. Demographic trade-offs predict tropical forest dynamics. Science 368, 165-168.
- Schaap MG, Leij FJ, Van Genuchten MT. 2001. Rosetta: a computer program for estimating soil hydraulic parameters with hierarchical pedotransfer functions. Journal of Hydrology 251, 163-176.
- Sebastian DE, Ganguly S, Krishnaswamy J, Du K, Nemani R, Ghosh S. 2019. Multi-scale association between vegetation growth and climate in India: a wavelet analysis approach. Remote Sensing 11, 2073.
- Sharma A, Goyal MK. 2020. Assessment of the changes in precipitation and temperature in Teesta River basin in Indian Himalayan Region under climate change. Atmospheric Research 231, 104670.
- Sharma CL, Sharma M, Carter MJ, Kharkongor BM. 2011. Inter species wood variation of Castanopsis species of Meghalaya. Journal of the Indian Academy of Wood Science 8. 124-129.
- Shinohara Y, Tsuruta K, Ogura A, Noto F, Komatsu H, Otsuki K, Maruyama T. 2013. Azimuthal and radial variations in sap flux density and effects on stand-scale transpiration estimates in a Japanese cedar forest. Tree Physiology 33, 550-558.
- Shrestha UB. Gautam S. Bawa KS. 2012. Widespread climate change in the Himalayas and associated changes in local ecosystems. PLoS ONE 7, e36741.
- Siddia Z. Cao KF. 2018. Nocturnal transpiration in 18 broadleaf timber species under a tropical seasonal climate. Forest Ecology and Management 418, 47–54.
- Singh SP, Bassignana-Khadka I, Karky BS, Sharma E, Eklabya S, Sharma E. 2011. Climate change in the Hindu Kush-Himalayas. The state of current knowledge. Kathmandu, Nepal: International Centre for Integrated Mountain Development (ICIMOD).
- Singh SP, Tewari A, Singh SK, Pathak GC. 2000. Significance of phenologically asynchronous populations of the central Himalayan oaks in drought adaptation. Current Science 79, 353-357.
- Singh SP, Zobel DB, Garkoti SC, Tewari A, Negi CMS. 2006. Patterns in water relations of central Himalayan trees. Tropical Ecology 47, 159–182.
- Sobrado MA. 2003. Hydraulic characteristics and leaf water use efficiency in trees from tropical montane habitats. Trees 17, 400-406.
- Sudhakar S, Prasad PRC, Arrawatia ML, Sudha K, Babar S, Rajeshwar Rao SKSV. 2008. Landscape analysis in Fambong Lho wildlife sanctuary, east district, Sikkim, India using remote sensing and GIS techniques. Journal of the Indian Society of Remote Sensing 36, 203-216.

- Sundriyal RC, Sharma E. 1996. Anthropogenic pressure on tree structure and biomass in the temperate forest of Mamlay watershed in Sikkim. Forest Ecology and Management 81, 113-134.
- Sundrival RC. Sharma E. 2003. Underutilized edible plants of the Sikkim Himalaya: need for domestication. Current Science 85, 731-736.
- Suzuki M, Noshiro S, Takahashi A, Yoda K, Joshi L. 1991. Wood structure of Himalayan plants, II. In Ohba H, Malla SB. eds. The Himalayan plants, vol. 2. Tokyo: The University of Tokyo, 1-118.
- Tanaka K, Takizawa H, Kume T, Xu J, Tantasirin C, Suzuki M. 2004. Impact of rooting depth and soil hydraulic properties on the transpiration peak of an evergreen forest in northern Thailand in the late dry season. Journal of Geophysical Research 109, D23107.
- Tang CQ. Ohsawa M. 1999. Altitudinal distribution of evergreen broad-leaved trees and their leaf-size pattern on a humid subtropical mountain, Mt. Emei, Sichuan, China. Plant Ecology 145, 221-233.
- Terrer C, Jackson RB, Prentice IC, et al. 2019. Nitrogen and phosphorus constrain the CO₂ fertilization of global plant biomass. Nature Climate Change 9.684-689.
- Tewari A, Bhatt J, Mittal A. 2016. Influence of tree water potential in inducing flowering in Rhododendron arboreum in the central Himalayan region. iForest 9, 842-846.
- Tewari A, Shah S, Singh N, Mittal A. 2018. Treeline species in Western Himalaya are not water stressed: a comparison with low elevation species. Tropical Ecology 59, 313-325.
- Wang H, He K, Li R, Sheng Z, Tian Y, Wen J, Chang B. 2017. Impact of time lags on diurnal estimates of canopy transpiration and canopy conductance from sap-flow measurements of *Populus cathayana* in the Qinghai–Tibetan Plateau. Journal of Forestry Research 28, 481-490.
- Wang J, Turner NC, Feng H, Dyck M, He H. 2023. Heat tracer-based sap flow methods for tree transpiration measurements: a mini review and bibliometric analysis. Journal of Experimental Botany 74, 723-742.
- Wang K, Dickinson RE. 2012. A review of global terrestrial evapotranspiration: observation, modeling, climatology, and climatic variability. Reviews of Geophysics 50, RG2005.
- Wang Q, Gao J, Zhao P, Zhu L, Ouyang L, Ni G, Zhao X. 2018. Bioticand abiotic-driven variations of the night-time sap flux of three co-occurring tree species in a low subtropical secondary broadleaf forest. AoB PLANTS 10, ply025.
- Wiedemann A, Marañón-Jiménez S, Rebmann C, Herbst M, Cuntz M. 2016. An empirical study of the wound effect on sap flux density measured with thermal dissipation probes. Tree Physiology 36, 1471-1484.
- Wohl E, Barros A, Brunsell N, et al. 2012. The hydrology of the humid tropics. Nature Climate Change 2, 655-662.
- Yang Y, Donohue RJ, McVicar TR, Roderick ML, Beck HE. 2016. Longterm CO2 fertilization increases vegetation productivity and has little effect on hydrological partitioning in tropical rainforests. Journal of Geophysical Research, Biogeosciences 121, 2125-2140.
- Zeppel M, Macinnis-Ng CMO, Ford CR, Eamus D. 2008. The response of sap flow to pulses of rain in a temperate Australian woodland. Plant and Soil **305**, 121-130.
- Zeppel MJB, Lewis JD, Medlyn B, et al. 2011. Interactive effects of elevated CO₂ and drought on nocturnal water fluxes in Eucalyptus saligna. Tree Physiology **31**, 932–944.
- Zhang X, Wang Y, Wang Y, Zhang S, Zhao X. 2019. Effects of social position and competition on tree transpiration of a natural mixed forest in Chongging, China. Trees 33, 719-732.
- Zhang Z, Sun G, Chen L, Xu H, Chen S. 2020. Biophysical controls on nocturnal sap flow in plantation forests in a semi-arid region of northern China. Agricultural and Forest Meteorology 284, 107904.
- Zhang Z, Zhao P, Zhao X, Zhou J, Zhao P, Zeng X, Hu Y, Ouyang L. 2018. The tree height-related spatial variances of tree sap flux density and its scale-up to stand transpiration in a subtropical evergreen broadleaf forest. Ecohydrology
- Zobel DB, Garkoti SC, Singh SP, Tewari A, Negi CMS. 2001. Patterns of water potential among forest types of the central Himalaya. Current Science 80, 774-779.