

1 **Different water relations between flower and leaf periods: a case**
2 **study in flower-before-leaf-emergence *Magnolia* species**

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24

25 **Abstract.** The differing water relations between flowers and leaves on a plant reflect
26 the lack of coordination between reproductive and vegetative organs during the
27 evolution of angiosperm species. Although the amount of water that flowers consume
28 has been reported to vary across species, accurate measurements of flower water
29 relations compared to that of leaves at the branch level are lacking, and how flowers
30 regulate their hydraulic function and structure to maintain water balance remains
31 unclear. To explore the ecophysiological basis underpinning the differences between
32 flowers and leaves, we measured hydraulic and morphological traits and monitored
33 sap flow in flowers and leaves from the same branches of two Magnoliaceae species
34 that flower before leaf emergence (*Magnolia denudata* and *Magnolia soulangeana*).
35 Sap flux density (J_s) of flowers was 22% and 55% of that predicted for leaves in *M.*
36 *denudata* and *M. soulangeana*, respectively. J_s of flowers commenced before
37 predawn and ceased early in the afternoon, reflecting their night-time flowering
38 pattern and a dramatic decrease of J_s with increasing vapour pressure deficit (D)
39 under the high light of midday. Relative to leaves, tepals were thicker and more
40 hydrated, and had bigger but scarcer stomata, leading to lower stomatal conductance
41 (g_s) and transpiration rate (E), less negative water potential (Ψ_{tepal}), and lower
42 hydraulic conductance. This study revealed different hydraulic patterns in the flowers
43 and leaves of the two *Magnolia* species. Although flowers consumed less than half the
44 water that leaves did, they used different strategies to maintain sufficiently high Ψ to
45 sustain hydraulic safety. *Magnolia* flowers retained more hydrated tepals by
46 exhibiting less water loss than leaves via lower hydraulic conductance. In contrast,
47 *Magnolia* leaves maintained high transpiration rates through efficient stomatal
48 responses to environmental changes compared to flowers.

49

50 **Additional keywords:** floral hydraulics, flowering stage, gas exchange, leaf hydraulic
51 conductance, Magnoliaceae, sap flow, stomata, water potential, xylem hydraulic
52 conductivity.

53

54

55 **Introduction**

56 The primary function of flowers is reproduction and their development requires
57 continuous supplies of water, nutrients and carbohydrates, transported via vascular
58 systems from other organs (Galen *et al.* 1999; Chapotin *et al.* 2003; Feild *et al.*
59 2009b). Although flowers assimilate little carbon, they are located along the outer
60 periphery of the tree canopy, an exposure that threatens desiccation. Thus to attract
61 pollinators, flowers must maintain water balance and turgor to prevent wilting,
62 although they may still transpire significant amounts of water and compete for
63 resources with leaves (Roddy and Dawson 2012; Teixido and Valladares 2014). The
64 coordination of activities between reproductive and vegetative organs within a plant is
65 a fascinating topic (Gross and Soule 1981; Reekie and Bazzaz 1987; Lambrecht and
66 Dawson 2007), yet virtually unknown from a hydraulic perspective. The water
67 transport capacity of petals and leaves of angiosperm species evolved independently,
68 as the vein length per area (VLA) of petals are consistent from basal to more derived
69 lineages (Roddy *et al.* 2013), while VLA of leaves increased nearly threefold during
70 angiosperm evolution (Brodribb and Feild 2010). Although pollinators impose
71 important selection pressures on floral functional traits (Thien *et al.* 2009), the need to
72 survive water limitation must surpass the need to attract pollinators (Feild *et al.*
73 2009a), and water relation traits are directly linked to floral maintenance. For
74 example, a recent study of 11 orchid species reported that greater floral longevity
75 required higher floral dry mass per area and more negative turgor loss points, but the
76 morphological traits of flowers and leaves were independent (Zhang *et al.* 2017).
77 They also found that flowers had more negative P50 (water potentials inducing 50%
78 embolism of veins) than neighbouring leaves, a difference that was significant for two
79 woody species but not two herbaceous ones (Zhang and Brodribb 2017). Therefore,
80 the differing evolutionary trajectories of flowers and leaves suggest contrasting water
81 relation strategies in the two organs, yet the differing amount of water consumption
82 and underlying ecophysiology between flowers and leaves remain unclear.

83 The few studies that address water consumption in flowers indicate that this trait is
84 highly variable across and within species (Whiley *et al.* 1988; Blanke and Lovatt
85 1993; Galen *et al.* 1999; Lambrecht *et al.* 2011; Lambrecht 2013; Roddy *et al.* 2016).
86 For instance, Whiley *et al.* (1988) found that transpiration rate (E) of avocado (*Persea*
87 *americana*) flowers was ~60% that of nearby leaves, while cuticular conductance was
88 similar between flowers and leaves. However, another study found that E of avocado

89 flowers was higher than leaves, which was attributed to largely closed stomata and the
90 waxy surfaces of avocado leaves, as well as the small, low density stomata on the
91 flower petals (Blanke and Lovatt 1993). A delicate study using miniature sap flow
92 sensors to separately quantify water use in single flowers and leaves found two
93 understory species with nearly no sap flow to flowers, while water flow to flowers of
94 two sun-exposed species was 30~50% that of nearby leaves (Roddy and Dawson
95 2012). However, all of these studies were based on species that simultaneously produce
96 flowers and leaves by comparing E or sap flow at the tepal (*i.e.*, a collective name for
97 flower parts that cannot easily be divided into sepals and petals) or leaf level, and
98 accurate estimations of water use by flowers and leaves throughout entire trees has
99 never been reported.

100 Determining separate flower and leaf traits across an entire tree is traditionally
101 difficult. For example, estimates of total flower area are confounded when a large
102 number of the flowers are unevenly distributed, and the tree has a dynamic flowering
103 stage with different flowers continuously opening and fading quickly. One approach
104 to separately estimate sap flow to each organ requires removing leaves during
105 blossom time, but this method may redirect water to the remaining organs and
106 increase both hydraulic conductance and E per area (Meinzer and Grantz 1990) and,
107 as such, would not capture the actual flow partitioning between flowers and leaves in
108 intact plants. By contrast, species with a natural flower-before-leaf-emergence (FBL)
109 characteristic are ideal to study flower water consumption, as they can be directly
110 measured and then later compared with water consumed by leaves on the same branch
111 once leaves emerge.

112 There are over 70 FBL species commonly observed in China, most of which
113 aggregated in large families such as the Magnoliaceae (esp. section *Yulania*),
114 Rosaceae (esp. *Prunus*), and Fabaceae (esp. *Cercis*), while other FBL species are
115 randomly distributed in different families (literature surveyed by the first author).
116 FBL and early flowering are important strategies to occupy the cold early spring niche.
117 Based on analyses of global datasets, selection favoured early flowering plants, and
118 this selection pressure was stronger in temperate than tropical flora (Munguía-Rosas
119 *et al.* 2012). In insect-pollinated species, early flowering and the thermogenesis of
120 large flowers or inflorescences can attract more insects to achieve higher reproduction
121 efficiency (Dieringer 1999; Seymour *et al.* 2003). Furthermore, due to their high

122 ornamental value, FBL species have been cultivated widely to produce larger, more
123 fragrant and colourful flowers (Azuma *et al.* 1999).

124 The Magnoliaceae family is commonly used to study the evolution of flowering
125 plants, with focuses on floral anatomy (Xu and Rudall 2006), pollination biology
126 (Thien 1974; Azuma *et al.* 1999; Thien *et al.* 2000), and phylogenetics and
127 geographical distributions (Qiu *et al.* 1999; Azuma *et al.* 2001; Kim and Suh 2013;
128 Liu *et al.* 2016). Since Magnoliaceae species emerged prior to bee pollinators, their
129 large flower size and floral thermogenesis co-evolved with beetle pollination (Thien
130 1974; Dieringer 1999; Gottsberger *et al.* 2012; Wang *et al.* 2014). FBL species in the
131 Magnoliaceae only exist in sections *Yulania* and *Michelia* (subgenus *Yulania*) within
132 the genus *Magnolia* (Figlar and Nooteboom 2004), and the flowering period of
133 *Yulania* species are the earliest (February) among all the Magnoliaceae lineages (Law
134 2004). *Yulania* species also have very large flowers (*e.g.*, single tepal length and
135 width are about 10 and 5 cm, respectively) compared with most flowering species and
136 other FBL species (Dandy 1927; Law 2004). For these reasons, we chose to focus on
137 section *Yulania* species in this study.

138 The two *Yulania* study species were grown in close proximity and flowered
139 concomitantly in the South China Botanical Garden in Guangzhou, China. We
140 monitored sap flow of branches in both species and microclimate conditions
141 throughout flowering, leaf expansion and maturation periods, as well as daily gas
142 exchange and water potential of tepals and leaves, and morphological and hydraulic
143 traits associated with water transport. This research aimed to: (1) accurately quantify
144 the water consumption by flowers and leaves of two *Yulania* species, taking
145 advantage of the distinctive flower and leaf phenology of FBL species; and (2)
146 investigate the water relations for flowers and leaves by integrating floral, leaf and
147 stem hydraulic measurements. We hypothesized that (1) flowers would use less water
148 per area than leaves of our study species, considering the lower temperatures during
149 the flowering than vegetative period and previous findings that tepals have sparser
150 stomata and lower *E* and hydraulic conductance than leaves in *Magnolia grandiflora*
151 (Feild *et al.* 2009b); and (2) although flowers can regulate water loss by reducing
152 stomatal conductance and tepal and stem water conductivities, these traits might be
153 particularly sensitive to environmental change, causing flowers to avoid dehydration
154 less efficiently than leaves.

155

156 **Material and methods**

157 *Study site and species*

158 Experiments were carried out in the South China Botanical Garden (SCBG) (23°11'N,
159 113°21'E, 20 m altitude) in Guangzhou, China, located in the low-subtropical
160 monsoon climatic region where mean annual temperature is 21.2°C, spanning 13.6°C
161 in January to 28.9°C in July. Mean annual precipitation is ~1700 mm, 80% of which
162 occurs in the wet season between April and September.

163 The study species included *Magnolia denudata* Desr., a famous ornamental species
164 with large white flowers and *Magnolia soulangeana* Soul.-Bod. 'Zhusha', a hybrid
165 (*Magnolia denudata* Desr. × *Magnolia liliflora* Desr.) bred for ornamental purposes,
166 which exhibits large showy purple flowers. Considering feasibility and the number of
167 flower buds available, four *M. denudata* and eight *M. soulangeana* individual trees
168 were selected for sap flow monitoring. Flowers of both species have 9 tepals arranged
169 in 3 whorls, with many spirally arranged stamens in the center. All sampled
170 individuals were mature trees, growing within 200 m² of the exhibition area in SCBG
171 (Liu *et al.* 1997), ranging from 6 to 10 m in height, and 12 to 17 cm in diameter at
172 breast height (DBH).

173

174 *Flowering stage records, tepal and leaf area calculation*

175 Flowering stage was recorded on six and ten branches from four and eight trees for *M.*
176 *denudata* and *M. soulangeana*, respectively. Every day during the flowering period,
177 we recorded the number of flowers on each branch in five custom classified stages:
178 buds with bracts sealed, buds with bracts open, half-open flowers with bracts dropped,
179 fully-open flowers, and faded flowers. We calculated the ratio of open flowers (*i.e.*,
180 number of half and fully-open flowers/total number of flowers on a branch), and
181 flower fading speed (*i.e.*, number of faded flowers/total number of flowers on a
182 branch).

183 Allometric relationships between the basal stem diameter of a branch and the total
184 flower or leaf area on that branch were evaluated using power functions. Because it is
185 prohibited to prune large branches of these ornamental garden trees, we could only
186 measure hydraulic traits on small branches (diameter ~10 mm) and then build models
187 to predict flower and leaf areas on the large branches that we monitored. Total flower
188 area on each branch was calculated as the total number of flowers × mean area of a
189 single flower, which was the average value based on 15 fully-open flowers from

190 nearby branches for each species. Leaf areas on small branches (diameter <10 mm)
191 were measured by a leaf area meter (Li-3000A; Li-Cor, Lincoln, NE, USA), and stem
192 diameters were measured with a calliper. We also selected 15 large branches
193 (diameter 10~40 mm) for each species, and measured the number and diameter of all
194 small branches on them, such that total areas of leaves could be calculated from stem
195 diameters for branches used for sap flow monitoring. We also recorded the average
196 individual tepal and leaf areas, and thickness of leaves and tepals (*i.e.*, at the thickest
197 and thinnest parts, since the base of a tepal is very thick and tapers to the upper
198 margin).

199

200 *Sap flow and environment monitoring*

201 Sap flow was monitored on the same branches that we used to record flowering stage,
202 using the heat balance method (Sakuratani 1981) with the Dynagage Flow32-1K
203 system (Dynamax, Houston, TX, USA). Constrained by branches of a suitable
204 diameter, length, and available straight segment without small branches, gauges were
205 installed at different heights and directions along the trees within a 50 m diameter
206 circle from each data logger. Every gauge and cable connection were waterproofed to
207 avoid rainfall damage. The thermal conductance constant (K_{sh}) for each gauge was
208 calibrated with the heat balance function between 01:00 and 05:00 on 2 to 3 days with
209 heavy cloud or rain, when no sap flow was assumed to occur before the sunrise.
210 Gauge outputs were measured every 60 s and recorded as 10-min means with a
211 CR1000 data logger. The original data were sap flow (g hr^{-1}), which were transformed
212 into sap flux density (J_s , $\text{g m}^{-2} \text{s}^{-1}$) by dividing sapwood area for each of the 16
213 branches. We modelled the relationships between sapwood area and stem diameter for
214 the two species, based on data of smaller branches (diameter<15 mm) during the
215 measurement of hydraulic conductivity, and data of larger branches (diameter 15~60
216 mm) from cores collected by a tree growth cone after removing the equipment to get
217 accurate estimations for each branch. Monitoring occurred between Feb-19 and Mar-
218 27, 2015, which encompassed the entire flowering (Feb-15 to Mar-10) and leaf
219 growth (Mar-2 to Mar-20) periods. However, branches with fewer than five flowers
220 showed sap flow values near zero during most of the flowering period, with the
221 exception of some irregular high points. Only six larger branches showed regular
222 daily dynamics (three *M. denudata* and three *M. soulangeana*), and were used in
223 further analysis of sap flow during the flowering period.

224 An automatic weather station (ECH2O Utility, Decagon Devices Inc. WA, USA)
225 was setup on the third floor roof about 100 m away from the experimental site,
226 monitoring the environment every 60 s, and recording it as 10-min means.
227 Meteorological data included air temperature (T , °C), relative humidity (RH, %), solar
228 radiation (SR, $W\ m^{-2}$), and rainfall (mm) during the experimental period, with vapour
229 pressure deficit (D , kPa) calculated as $a \times \exp[b \times T / (T + c)] \times (1 - RH)$, where a , b , and c
230 are fixed parameters as 0.611 kPa, 17.502 (unitless) and 240.97 °C, respectively.

231

232 *Gas exchange and water potential*

233 Gas exchange was measured on tepals and leaves over two sunny days; one in the
234 middle of the flowering period (Feb-24, 11:00 and 16:00), and the other after most
235 leaves had expanded (Mar-26, 7:00, 10:30, 13:00, 16:30 and 18:00). On five trees per
236 species, we cut off one half-open and one fully-open flower from each tree using a
237 tree pruner, avoiding flowers on the branches where we monitored sap flow. Flower
238 stalks were immediately transferred into water and gas exchange rates were measured
239 on tepals from three whorls (1st, outer whorl; 2nd, middle whorl; 3rd, inner whorl). The
240 sun-exposed branches were bent downward to access leaves for measurements. Five
241 trees for each species were chosen, and four leaves on each tree were measured. The
242 two species we studied have clusters of four leaves each in one of four growth stages
243 (1st, half-expanded leaves; 2nd, fully-expanded leaves; 3rd, mature leaves; 4th, older
244 basal leaves), thus we measured one representative leaf from each stage on each tree.

245 Stomatal conductance (g_s , $mol\ m^{-2}\ s^{-1}$) and transpiration rate (E , $mmol\ m^{-2}\ s^{-1}$) of
246 tepals and leaves were measured with an open leaf gas exchange system (LI-6400, LI-
247 COR, Lincoln, NE, USA). For daily dynamics, a chamber with a transparent lid was
248 used to measure natural light conditions, while CO_2 concentration, T , RH, and D
249 uncontrolled in the chamber, in order to calculate hydraulic conductance based on the
250 real-time E . During gas exchange measurements, water potentials (Ψ , MPa) of tepals
251 taken from the same flower, and of leaves taken from the same twig were measured
252 using a pressure chamber (PMS, Corvallis, OR, USA). Stem water potential (Ψ_{stem} ,
253 MPa) was also measured, using leaves that were wrapped with foil and sealed in
254 plastic bags the evening before measurement day.

255

256 *Stem hydraulic conductivity*

257 Early in the morning, terminal branches (8~10 mm in diameter) from five trees per
258 species were excised. All stems were immediately recut under water and leaves were
259 misted with water, before samples were sealed in black plastic bags with moist towels
260 to prevent transpiration and quickly transported to the laboratory. A stem segment
261 20~30 cm in length was cut under water from each branch, and both cut ends were
262 trimmed with a razor blade. Branch segments were first flushed with filtered and
263 degassed 20 mmol KCl solution at a pressure of 0.1 MPa for 10 min to remove air
264 embolism. Then hydrostatic pressure generated by a 50 cm hydraulic head drove
265 water flow through the segments. The downstream end of each segment was
266 connected to a pipette and the time for fluid in the pipette to cross a certain graduation
267 was recorded. Hydraulic conductivity (K_h , $\text{kg m s}^{-1} \text{MPa}^{-1}$) was calculated as water
268 flux through the segment divided by the pressure gradient driving the flow. Sapwood
269 specific hydraulic conductivity (K_s , $\text{kg m}^{-1} \text{s}^{-1} \text{MPa}^{-1}$) was calculated as K_h divided by
270 the sapwood cross section area (A_s). Leaf specific hydraulic conductivity (K_L , kg m^{-1}
271 $\text{s}^{-1} \text{MPa}^{-1}$) is the ratio of K_h to the total leaf area attached to the stem segment (A_L). A_L
272 was measured by a leaf area meter to calculate the leaf to sapwood area ratio (A_L/A_s ,
273 $\text{m}^2 \text{cm}^{-2}$). Sapwood samples with bark removed were saturated in water overnight,
274 then after wiping the surface dry, the sapwood fresh volume was measured by the
275 water displacement method. These samples were then oven-dried at 70 °C for 72 h and
276 weighed to obtain dry mass. Sapwood density (WD, g cm^{-3}) was calculated as the
277 ratio of dry mass to fresh volume from the same branches used for K_h measurements.

278

279 *Tepal and leaf turgor loss point (Ψ_{tlp})*

280 Pressure volume (PV) curve analysis, based on the bench drying method, was used to
281 calculate turgor loss point (Ψ_{tlp}) for both tepals and leaves (Tyree and Hammel 1972).
282 Terminal branches that contained tepals or leaves were excised from three to five trees
283 per species, recut underwater, and rehydrated until water potential was greater than -
284 0.05 MPa. Tepal and leaf weight, and Ψ were measured periodically during
285 desiccation. After pressure-weight measurements, samples were oven-dried at 70 °C
286 for 72 h, dry weight was used to calculate leaf (or tepal) dry matter content
287 (LDMC, %), and Ψ_{tlp} was determined according to PV models with leaf relative water
288 content (RWC) and $-\Psi^{-1}$ (Schulte and Hinckley 1985). The hydraulic safety margin
289 (HSM, MPa) was calculated as the difference between minimum water potential (*i.e.*,

290 Ψ_{pm}) and Ψ_{tp} . Relative capacitance at full turgor (C_{ft0} , MPa^{-1}) was calculated as Δ
291 $\text{RWC} / \Delta \Psi$ between full turgor and turgor loss point. Leaf (or tepal) area specific
292 capacitance at full turgor (C_{ft} , $\text{mol m}^{-2} \text{MPa}^{-1}$) was standardized as $C_{ft0} \times (\text{leaf turgor}$
293 $\text{mass} - \text{leaf dry mass}) / \text{leaf area}$ (Sack *et al.* 2003).

294

295 *Tepal and leaf hydraulic conductance (K_{tepal} ; K_{leaf})*

296 Although there are different methods to measure hydraulic conductance of detached
297 tepals and leaves (Sack *et al.* 2002), our preliminary experimentation showed that the
298 high-pressure method was not suitable for K_{tepal} measurement, since large amounts of
299 mucilage in tepals may contribute to capacitance but may not increase conductance,
300 which would result in unusually high K_{tepal} values. Thus we estimated K_{tepal} and K_{leaf}
301 ($\text{mmol m}^{-2} \text{s}^{-1} \text{MPa}^{-1}$) based on the real-time transpiration and water potential data
302 (*i.e.*, K_{tepal} and $K_{leaf} = E / (\Psi_{stem} - \Psi)$), which we used to represent the hydraulic
303 conductance of tepals and leaves under natural conditions (Brodribb and Holbrook
304 2003).

305

306 *Specific leaf (or tepal) area (SLA), nutrients and stomatal traits*

307 Specific leaf area (SLA, $\text{cm}^2 \text{g}^{-1}$) was calculated as leaf area divided by leaf dry mass.
308 For each species, 20 tepals and leaves were scanned using a leaf area meter then oven-
309 dried at 70 °C for 72 h. Dried samples were ground and homogenized for nutrient
310 measurements. Total nitrogen content (N, %) was determined by Kjeldahl analysis
311 after digestion with concentrated H_2SO_4 . Total phosphorus content (P, %) was
312 analyzed by atomic absorption spectrum photometry (UV-6000; Metash, Shanghai,
313 China).

314 Epidermal peels of fresh tepals and leaves were extracted using a sharp razor blade,
315 then observed under a microscope equipped with a digital camera (Optec Instrument,
316 Chongqing, China) and a computerized image analysis system (OPTPro2012 version
317 4.0, Optec software). Three epidermal peels from each of three flower whorls and four
318 leaf growth stages were analyzed per species and, on each peel, three images were
319 randomly chosen as replicates. Guard cell length (GL) and width (GW) were
320 measured, and stomatal density (SD) was counted. The stomatal pore area index
321 (SPI, %) indicated stomata pore area per leaf area, which equaled $\text{SD} \times \text{GL}^2$ (Sack *et al.*
322 2003). The maximum diffusive conductance to water vapour (g_{max}) was estimated as

323 transpiration potential, calculated as $(d/v) \times SD \times a_{\max} / [(l + \pi/2) \times \sqrt{(a_{\max}/\pi)}]$ (Brown and
324 Escombe 1900; Franks and Beerling 2009); where d is the diffusivity of water vapour
325 in air at 25 °C ($\text{m}^2 \text{s}^{-1}$); v is the molar volume of air at 25 °C ($\text{m}^3 \text{mol}^{-1}$); SD is
326 stomatal density; a_{\max} is the maximum area of the open stomatal pore, estimated as
327 $\pi \cdot (p/2)^2$ where p is stomata pore length and was approximated as $GL/2$ as in Franks
328 and Beerling (2009); l is stomata depth for fully open stomata, approximated as $GW/2$;
329 and π is the geometric constant. In *Magnolia* species, stomata exist on both the
330 adaxial and abaxial surfaces of tepals, but only on the abaxial surface of leaves. Thus
331 we combined the calculated SPI and g_{\max} of both tepal surfaces to obtain total SPI and
332 g_{\max} values.

333

334 *Data analyses*

335 All data were analysed in R v3.0.3 (R Development Core Team 2013). First, we tested
336 whether the tepal or leaf traits differed among the three flower whorl types or among
337 the four leaf growth stages using one-way ANOVAs, such that values that differed
338 significantly among flower whorls or leaf stages were then analysed using multiple
339 comparisons (Tukey HSD) in the daily dynamic dataset. Next, the differences
340 between flowers and leaves were tested using t -tests for each species separately. In the
341 above tests, data were natural log-transformed to fulfil the requirement of normal
342 distribution, using absolute value for traits with negative values (e.g. Ψ_{tip}).

343 To quantify the relationships between J_s and D , we performed boundary line
344 analyses (Chambers *et al.* 1985; Ewers *et al.* 2005). We used J_s data from days when
345 flower opening ratios were stable and all leaves were expanded, filtering out data
346 collected under limiting light ($SR=0 \text{ W m}^{-2}$) and during low D ($<0.1 \text{ kPa}$) when
347 empirical relationships between canopy stomatal conductance (G_s) and D were not
348 well constrained (Oren *et al.* 1999). This will enable the resulting boundary line to
349 give the best estimate of hydraulic limitation to water flux because the boundary line
350 occurred during conditions that lead to the highest G_s at any given D . Next, the
351 relationships between J_s and D were examined using the boundary line analysis
352 independently for data grouped by four (0~200, 200~400, 400~600, 600~800 W m^{-2})
353 and two (0~400, 400~800 W m^{-2}) light gradients, in order to examine light effects.
354 We found that both flowers and leaves showed significantly different relationships
355 between the two light gradients and, as such, we used low light (LL, 0~400 W m^{-2})
356 and high light (HL, 400~800 W m^{-2}) in the final analyses. We used log-linear models

357 to predict J_s from $\ln D$, which could indicate the sensitivity of sap flow response to
358 changes in D . Furthermore, considering similar SR conditions, and the range of D on
359 Mar-26 (when in leaf) encompassed that measured on Feb-24 (when in flower), we
360 predicted J_s of leaves based on the relationships between J_s of leaves and D for *M.*
361 *denudata* and *M. soulangeana*, in order to directly compare J_s of flowers and leaves.

362 To quantify the sensitivity of g_s to D from the daily dynamic data, we selected
363 morning (10:30-11:00) and afternoon (13:00-16:30) periods to compare flowers and
364 leaves. According to Lohammar's function $g_s = -k \times \ln D + b$, where k is the sensitivity
365 index and, b is a constant (Lohammar *et al.* 1980), we built models for each species in
366 each time period. The relationships between g_s and Ψ for the daily dynamic data were
367 also tested, but clear patterns were not found.

368

369 **Results**

370 *Environments and sap flux density during flowering and vegetative periods*

371 Flowering period had lower daily average D and T, but similar SR compared to the
372 vegetative period (Fig. 1a, b). While there were several rainfall events that
373 distinctively affected D and T, sunny days in both flowering and vegetative periods
374 enabled the daily dynamic measurements of gas exchange and water potential. During
375 the main flowering period, the average sapwood area based J_s was about 234 and 750
376 $\text{kg m}^{-2} \text{ day}^{-1}$ for *M. denudata* and *M. soulangeana*, respectively. In both species, J_s
377 was clearly lower in the flowering period than the vegetative one (Fig. 1c, d),

378 The sunny day with only flowers (Feb-24) or leaves (Mar-26) on the tree elicited
379 very different responses (Fig. 2). Daily D peaked at 13:00 during the flowering period
380 and 16:00 in the vegetative period (Fig. 2a), due to T and RH patterns. Specifically, T
381 was consistently 6.34 ± 0.11 °C lower in the flowering than vegetative day, and RH
382 decreased from 90% at 6:00 to a minimum of 60% at 13:00 in the flowering day,
383 while RH in the vegetative day decreased from 95% at 6:00 to a minimum of 52% at
384 16:00. SR was similar in the mornings of flowering and vegetative periods, but was
385 slightly lower after 13:00 in the flowering period (Fig. 2b). J_s was lower during the
386 flowering period than the vegetative one, a pattern that was more dramatic in *M.*
387 *denudata* than *M. soulangeana* (*i.e.*, daily accumulated floral water consumption was
388 17% and 53% that of leaves for *M. denudata* and *M. soulangeana*, respectively). J_s
389 peaked around 10:00 during flowering period and around 14:00 during the vegetative
390 period (Fig. 2c, d). Furthermore, although D was higher in vegetative than flowering

391 period (Fig. 2a), J_s of flowers was still smaller than that predicted for leaves in the
392 flowering period, and daily accumulated floral water consumption was 22% and 55%
393 that of leaves for *M. denudata* and *M. soulangeana*, respectively (Fig. S1).

394

395 *Flowering stages*

396 We selected periods with stable ratios of opening and fading floral stages for sap flow
397 data analyses to avoid the variance brought by changing flower number. *M. denudata*
398 flowered quickly and maintained a high open flower ratio (i.e., around 70%) for six
399 days, after which the flowers all dramatically faded within four days (Fig. 3a, c).
400 Meanwhile, the flowering stage of *M. soulangeana* was slow, maintaining only 30%
401 open flowers for about a week. Although *M. soulangeana* then remained with a 40%
402 open flower ratio after the initial seven days, the fading stage had already commenced
403 and the majority of flowers (70%) quickly faded within three days (Fig. 3b, d).

404

405 *Effects of vapour pressure deficit on sap flux density and stomatal conductance*

406 J_s of flowers was more vulnerable to high light than J_s of leaves (Fig. 4). Under low
407 light (LL), J_s of flowers initially increased, followed by a slight decrease with $\ln D$,
408 while under high light (HL), J_s of flowers decreased with $\ln D$ for both species (Fig.
409 4a, b). On the other hand, J_s of leaves increased with rising $\ln D$ at both light levels,
410 with higher J_s under HL than LL (Fig. 4c, d). J_s of *M. denudata* leaves was much
411 higher than that of its flowers, while the maximum J_s of *M. soulangeana* flowers was
412 even higher than that of *M. soulangeana* leaves (Fig. 4).

413 In general, g_s of leaves was significantly higher than that of flowers, and leaf g_s
414 was also more sensitive to changes in D (Fig. 5). In the morning, g_s in both flowers
415 and leaves reached higher maximum values and decreased more dramatically with
416 increasing $\ln D$ than in the afternoon. In both morning and afternoon measurements, *M.*
417 *soulangeana* showed higher sensitivity in tepal g_s to $\ln D$, but lower sensitivity of leaf
418 g_s to $\ln D$, compared to *M. denudata* (Fig. 5).

419

420 *Comparisons of plant traits between flowers and leaves*

421 Flowers and leaves differed significantly in nearly all of the measured traits, with the
422 exception of K_s and N and P contents (Table 1). Both single leaf area and total leaf
423 area were greater than those of flower tepals, on branches at the same diameter scale
424 (A_L/A_s in Table 1; Fig. S2). Leaves were thinner than even the thinnest parts of tepals,

425 with higher SLA and LDMC. Thus the averaged total water content amount for tepals
426 and leaves standardized by sapwood area showed that: flowers stored more water than
427 leaves on the same diameter branch (101.6 g cm⁻² and 88.3 g cm⁻² for *M. denudata*
428 with flowers and leaves, respectively; 103.1 g cm⁻² and 90.2 g cm⁻² for *M.*
429 *soulangeana* with flowers and leaves, respectively). Tepals had much larger but also
430 rarer, stomata than leaves, which resulted in SPI and g_{\max} of tepals to be only 3% and
431 2% that in leaves, respectively. However, the measured g_s and E of tepals were about
432 27% and 22% that of leaves for *M. denudata*, respectively, and up to 65% and 55%
433 that of leaves for *M. soulangeana*, respectively. Compared to tepals, leaves had more
434 negative Ψ_{am} , Ψ_{pm} and Ψ_{tlp} , and higher HSM in *M. denudata* but lower HSM in *M.*
435 *soulangeana* (all the HSM>0). Leaves also had much lower C_{ft} , much higher K_{leaf} and
436 smaller K_L than tepals (Table 1).

437 In addition, several traits differed by flower whorls or leaf growth stages in both
438 study species, including leaf area, thickness, flower LDMC, g_s , E , flower Ψ , HSM and
439 K_{leaf} (K_{tepal}). In contrast, single tepal area, Ψ_{tlp} and C_{ft} differed among flower whorls
440 only in *M. soulangeana*. The remaining traits did not differ among whorls or stages
441 (Table 1; Table S1). Specifically, single leaf area was smallest in the half-expanded or
442 older basal leaves and largest in mature leaves. Tepal thickness of the 1st whorl was
443 the thinnest and gradually increased from the 2nd to the 3rd whorl, while half-expanded
444 leaves were thinner than other mature leaves. For LDMC of tepals, the 1st whorl had
445 the highest values, followed by the 2nd and 3rd whorls, while the HSM of tepals was
446 smallest in the 1st whorl. K_{tepal} increased between the 1st, 2nd and 3rd whorls, while K_{leaf}
447 was lowest in the half-expanded leaves and highest in the full-expanded leaves, with
448 mature and older leaves showing intermediate values (Table S1). For *M. soulangeana*,
449 single tepal area was largest in the 2nd whorl, followed by the 3rd and 1st whorls, and
450 both Ψ_{tlp} and C_{ft} increased from the 1st and 2nd whorls to the largest 3rd whorl. E
451 among flower whorls and leaf stages showed the same pattern as g_s analysed below.

452 Further investigations on the daily changes in g_s and Ψ showed that: (1) half-open
453 flowers had generally higher g_s than fully-open flowers, and tepals of half-open
454 flowers in the 3rd whorl had higher g_s than those of the 1st and 2nd whorls (Fig. S3a, b).
455 (2) Leaf g_s initially increased over the morning, peaked around 10:30, and then
456 decreased to near zero for the remainder of the day. Younger leaves (1st leaf) showed
457 higher g_s than mature leaves (Fig. S3c, d). (3) Ψ_{tepal} of half-open flowers was
458 remarkably variable and lacked clear patterns compared to those of fully-open flowers.

459 Ψ_{tepal} of the 1st whorl was more negative than those of the 2nd and 3rd whorls. There
460 were no differences of Ψ_{tepal} between morning and afternoon, or between the two
461 studied species (Fig. S4a, b). (4) Ψ_{leaf} was nearly -0.1 MPa at 7:00, reached its most
462 negative at 13:00, and then returned to around -0.2 MPa at 18:00. There were no
463 differences of Ψ_{leaf} among the four growth stages (Fig. S4c, d). Overall, average Ψ_{leaf}
464 values were more negative than Ψ_{tepal} in both species, and all the Ψ values were above
465 Ψ_{tjp} . Although minimum Ψ in *M. soulangeana* approached average Ψ_{tjp} , specific
466 HSMs remained above zero (Fig. S4, Table 1).

467

468 Discussion

469 *Sap flow and stomatal conductance patterns differ between flowers and leaves*

470 Sap flow in the *Magnolia* flowers that we measured showed distinct daily dynamic
471 patterns compared with leaves, with J_s starting early at predawn (or even from 4:00
472 for *M. soulangeana*), quickly peaking midmorning, then decreased the remainder of
473 the day, despite a continuous increase in D until 13:00. In contrast, leaf J_s remained
474 linked to D throughout the day (Fig. 2). Flowers of most Magnoliaceae species open
475 at night (Dieringer, 1999), probably because their main pollinators are beetles, which
476 are active during the night, while only their secondary pollinators (*i.e.*, bees) are
477 active during the day (Thien 1974). Although high J_s of flowers in the morning was
478 assumed to be associated with low Ψ_{tepal} (Ortuno *et al.* 2006), we show that this is not
479 the case for *Magnolia* species, as Ψ_{tepal} remained high throughout the day (-0.05 ~ -0.2
480 MPa) and did not show dramatically daily changes as in Ψ_{leaf} (-0.1 ~ -0.8 MPa) (Fig.
481 S4). This is perhaps due to lower stomatal or cuticular conductances in tepals
482 compared to leaves, or much higher C_{ft} in tepals than leaves, which could maintain
483 water above turgor (Chapotin *et al.* 2003). At the branch level, we also found that
484 flowers store more water than leaves on the same branches, so that branches do not
485 require high J_s to maintain water balance during flowering period. The buffering
486 effects of water stored in stems, which provided ~10% daily water consumption
487 independent of tree size (Meinzer *et al.* 2004), may similarly explain the low ratio of
488 J_s in flowers to that predicted for leaves (22% and 55% for *M. denudata* and *M.*
489 *soulangeana*, respectively). Therefore, we speculate that the driving forces behind
490 floral J_s might come not only from tepal E or Ψ_{tepal} changes during the day, but also
491 from flower opening forces at night and predawn. These forces may include the apical
492 growth (osmotic potential brought by carbohydrates decomposition) during floral

493 development (Xu and Rudall 2006), floral cuticular conductance brought by
494 thermogenesis (Dieringer 1999; Wang *et al.* 2014), and water needed for the physical
495 expansion of tepals (Wada *et al.* 2004; Azad *et al.* 2007). As we did not measure these
496 physiological activities directly here, we recommend that they be investigated in
497 future studies on floral hydraulics.

498 The *Magnolia* flowers in our study were more vulnerable to environmental
499 fluctuations than leaves, with floral J_s and g_s presenting different responses to
500 changes in D and light (Fig. 4-5). Under low light, flower J_s remained very low and
501 did not respond to increases in D , which might result from the buffering effects of
502 stored water within the tepals, as reported for mango inflorescences (Higuchi and
503 Sakuratani 2005). In contrast, the high light of the afternoon caused the J_s of flowers
504 to decrease quickly as D increased (Fig. 4), because the higher tepal C_{ft} indicates
505 greater water loss under the same D and light stress, i.e., flowers are much more
506 vulnerable to desiccation than leaves. We also noticed that some fully-open flowers
507 started to wilt in the afternoon due to high light or temperature, which caused high D
508 and allowed J_s to decrease, leaving water for the half-open flowers and buds the
509 following day. Together, this helps to define the overall flowering phenology at the
510 tree level. Furthermore, in our study species, low LDMC and the high Ψ_{tlp} and C_{ft} of
511 the tepals indicates large vacuoles in their parenchyma cells and high vulnerability to
512 desiccation, similar to orchids flowers (Zhang *et al.* 2017). Then the tepals produce
513 few stomata to help maintain low g_s and Ψ_{tepal} to sustain high HSM and avoid
514 desiccation under normal water conditions. Therefore, due to higher water storage and
515 lower water loss, we found that the absolute value of tepal g_s was only 27~65% that of
516 leaves, and had a shallower slope with $\ln D$ than leaves (Fig. 5). We also found that the
517 inner whorl of half-open flowers is the primary driver of flower water consumption
518 (i.e., higher Ψ_{tepal} and g_s than the other two whorls, Fig. S3-4; Table S1). While these
519 Ψ_{tepal} findings are consistent with those of *Magnolia grandiflora*, our g_s findings differ
520 such that the 1st whorl of *M. grandiflora* had higher g_s than the 3rd whorl (Feild *et al.*
521 2009b). One possible reason for this discrepancy may be due to the fully-open flowers
522 that they used, as the g_s in our study showed no differences between the 1st and 3rd
523 whorls for fully-open flowers (Fig. S3), indicating that water consumption strongly
524 depends on flowering stage.

525 In leaves, D and water transpired through gas exchange were clearly the main
526 drivers of water transportation and sap flow, as confirmed by the congruent pattern of

527 daily leaf J_s , g_s and D (Fig. 1, 2, 4, 5). Many studies address hydraulic regulation as a
528 method to prevent xylem embolism under water stress brought on by atmospheric
529 dryness (high evaporative demand) and/or soil drought (Tyree and Sperry 1989;
530 Nardini *et al.* 2012). Because our study had sufficient soil and stem water supplies,
531 modest increases in D would initially enhance evaporation, E and K_{leaf} . However, Ψ_{leaf}
532 may slightly drop and a continuous decrease in Ψ_{leaf} would cause stomata closure,
533 leading to lower g_s , E and K_{leaf} , such that xylem tensions in the stems could remain
534 within a safe range (Meinzer and Grantz 1990; Brodribb and Holbrook 2004; Franks
535 2004). Studies at the stand scale show that canopy stomata respond to D via the
536 regulation of g_s and Ψ_{leaf} (Granier and Loustau 1994; Oren *et al.* 1999; Oren *et al.*
537 2001), which is important to understand water balance within the whole ecosystem.
538 Therefore, co-regulation of Ψ_{leaf} , K_{leaf} , and J_s is the result of the hydraulic-
539 photosynthetic coordination of leaves.

540

541 *Ecophysiology underpinning the different water relations between flowers and leaves*
542 Flowers of the two *Magnolia* species consumed less water per area (lower E and J_s)
543 than the leaves, while tepals showed lower K_{tepal} but higher K_L than leaves, due to
544 their specific structures. As assimilation organs, we found that leaves had higher
545 LDMC, indicating greater investments in veins and photosynthetic structures than
546 tepals, as is the case for most angiosperm species (Roddy *et al.* 2013). This allocation
547 leads to lower internal resistance and higher intrinsic K_{leaf} , and enables higher rates of
548 transpiration and photosynthesis in leaves (Brodribb *et al.* 2007). Our results were
549 consistent with this hypothesis in LDMC, K_{leaf} or K_{tepal} , and gas exchange traits.
550 Although thick and well-hydrated tepals led to less negative Ψ_{tepal} , their much lower E
551 was more decisive in K_{tepal} compared with K_{leaf} , showing similar K_{leaf} or K_{tepal} values,
552 as was also reported in *Magnolia grandiflora* (Feild *et al.* 2009b). Large, thick, and
553 hydrated tepals are commonly found in Magnoliaceae species that evolved in
554 relatively moist environments (Feild *et al.* 2009a). These tepal phenotypes may
555 effectively protect stamens and gynoecia, attract pollinators (mainly beetles) by colour,
556 fragrance, and thermogenesis under low air temperature (Azuma *et al.* 1999;
557 Dieringer 1999; Wang *et al.* 2014), or even provide food for pollinators (Thien 1974;
558 Gottsberger *et al.* 2012). Moreover, we found that K_S was similar in the flowers and
559 leaves of our study species, but that flower K_L was higher than that of leaves due to
560 the considerably lower A_L/A_S of flowers (Table 1). These findings confirm that stems

561 are hydraulically built to accommodate the high transpiration by leaves and, as such,
562 are hydraulically overbuilt for flowers. Stem xylem conduits are the structural basis of
563 K_S (Sperry *et al.* 2008), and these should not change appreciably during our two-
564 month experimental period. As the maximum hydraulic conductivity, K_S is suitable to
565 compare hydraulic conductivity potential rather than water transport situation *in situ*.
566 Therefore, while K_S and K_L values only showed different maximum hydraulic
567 conductivity between tepals and leaves, the *in situ* hydraulic differences could be
568 represented by K_{leaf} or K_{tepal} , g_s , E , Ψ , and J_S at leaf or tepal and branch levels, with
569 Ψ_{tip} as a reference to assess HSM, which was always positive under our study
570 conditions.

571 In the two *Magnolia* species studied here, stomata were larger and lower density on
572 the tepals than the leaves, which constrains stomatal conductance, leading to very low
573 absolute values of g_s and E in the flowers. This prevents water loss and helps to
574 maintain the water balance of flowers through stomatal adjustments (Franks and
575 Beerling 2009). Thus under naturally varying environmental conditions, all tepals of
576 fully-open and half-open flowers experienced water potentials higher than Ψ_{tip} (i.e.,
577 positive HSM in Fig. S4). Meanwhile, floral J_S was much less than leaves based on
578 both experimental data (Fig. 2) and simulated values (Fig. S1). Previous studies found
579 that floral stomata of several orchid species were dysfunctional and did not transpire
580 (Hew *et al.* 1980). However, our study found higher opening ratios in tepals than
581 leaves and that tepal g_s was about 27~65% that of leaves, firmly indicating the
582 functionality of tepal stomata. The relatively high g_s might also be affected by
583 evaporation through the epidermis and cuticle in the leaf chamber during gas
584 exchange measurements, which is likely much higher in flowers (30-90 mmol m⁻² s⁻¹
585 for magnoliids) than in leaves (Roddy *et al.* 2016). Consistent with our findings (Fig.
586 S3), E of avocado flowers is 60-80% of nearby leaves, peaking in the early morning
587 and dramatically declining midday (Whiley *et al.* 1988; Blanke and Lovatt 1993).

588 Considering the brief flowering period (7~10 days) and remarkably short lifespan
589 of each tepal (2~3 days) in *Magnolia* species, it should be more economical for the
590 whole plant to invest less water and carbon in the non-photosynthetic tepals (per unit
591 area). This was supported by our study, which found that flowers had lower J_S , E , and
592 LDMC in flowers than leaves. The strong selection pressures for greater hydraulic
593 conductance in leaves within developed angiosperm families did not exist for flowers
594 (Brodribb and Feild 2010; Roddy *et al.* 2013), especially in basal angiosperms like the

595 Magnoliaceae that evolved in wet habitats lacking hydraulic limitations (Feild *et al.*
596 2009a). This is consistent with a recent study that found basal angiosperm flowers
597 maintain higher K_{flower} due to traits related with high rates of water loss and supply
598 (Roddy *et al.* 2016).

599

600 **Conclusion**

601 This study demonstrated different water relations for flowers and leaves of two
602 flower-before-leaf-emergence *Magnolia* species. The ratio of J_S in flowers to that
603 predicted for leaves during the flowering period was 22% and 55% for *M. denudata*
604 and *M. soulangeana*, respectively. J_S in flowers began before predawn and ceased
605 early in the afternoon due to night-flowering and high sensitivity of g_s to D , indicating
606 that stomata closed early to save water before cavitation occurred. Thus, we propose
607 that the strongest driving forces of flower J_S might include Ψ_{tepal} and/or transpiration,
608 as well as other physiological processes during flowering, such as apical growth,
609 thermogenesis, and tepal expansion. In addition, flower water loss happened mainly in
610 the center of the flower and greatly depended on flowering stages. We then explored
611 the ecophysiological basis of the differences in water relations between leaves and
612 flowers, finding that tepals were thicker, more hydrated, had lower LDMC, and had
613 larger and less dense stomata, which lead to lower g_s , g_{max} , E , and K_{tepal} , less negative
614 Ψ_{tepal} and Ψ_{tlp} , and higher K_L than these traits in leaves. This study showed that to
615 keep constant Ψ and avoid losing water before cavitation, tepals maintain lower
616 hydraulic conductance than leaves, while leaves had more efficient stomatal responses
617 to D than tepals. Consequently, flowers consumed less than half the water that leaves
618 did at both the tepal, leaf, and branch levels for both species. Our study examined
619 water consumption and the ecophysiological basis between flowers and leaves in two
620 *Magnolia* species, which we hope will inspire future investigations on floral
621 hydraulics.

622

623 **Appendix**

624 An appendix is available online and consists of the following:

625 Table S1: Morphological and ecophysiological traits with significant differences
626 among three tepal whorls or four leaf growth stages of *M. denudata* and *M.*
627 *soulangeana*.

628 Fig. S1: Predicted J_S of leaves during the flowering period based on the relationships
629 between J_S and D , using D from Feb-24 for *M. denudata* and *M. soulangeana*.

630 Fig. S2: Flower or leaf areas versus stem diameters for *M. denudata* and *M.*
631 *soulangeana*.

632 Fig. S3. Daily changes in flower and leaf stomatal conductance (g_s) of *M. denudata*
633 and *M. soulangeana* during two sunny days with either only flowers or leaves on the
634 tree, respectively.

635 Fig. S4. Daily changes in flower and leaf water potential (Ψ) of *M. denudata* and *M.*
636 *soulangeana* during two sunny days with either only flowers or leaves on the tree,
637 respectively.

638

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647

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803 **Table 1. Morphological and ecophysiological traits of flowers and leaves of *Magnolia denudata* and *Magnolia soulangeana*, with results**
804 **of *t*-tests for each trait.** Data are mean \pm SEM, and natural log-transformed in models. Sample sizes (*n*) of flower or leaf traits are the same for
805 *M. denudata* and *M. soulangeana*, therefore only sample sizes for *M. denudata* are given in brackets. Differences between flowers and leaves for
806 each trait were analysed using *t*-tests (* $P < 0.05$; ** $P < 0.01$; *** $P < 0.001$; ns, not significant), “-” indicates *t*-tests are not applicable, “†”
807 indicates significant differences among three whorls of flowers, or four leaf growth stages by ANOVA, which are reported and further analyzed
808 in the Appendix. Abbreviations: DBH, diameter at breast height; WD, sapwood density; A_L/A_S , leaf to sapwood area ratio; SLA, specific leaf (or
809 tepal) area; LDMC, leaf (or tepal) dry matter content; SPI, stomatal pore area index; g_{max} , maximum stomatal conductance to water vapour; g_s ,
810 stomatal conductance; E , transpiration rate; Ψ_{am} , leaf (or tepal) water potential at 10:30~11:00; Ψ_{pm} , leaf (or tepal) water potential at 16:00~16:30;
811 Ψ_{tlp} , turgor loss point; HSM, hydraulic safety margin; C_{ft} , capacitance at full turgor; K_{leaf} or K_{tepal} , leaf (or tepal) hydraulic conductance; K_s ,
812 sapwood specific hydraulic conductivity; K_L , leaf (or tepal) specific hydraulic conductivity.

	<i>Magnolia denudata</i>			<i>Magnolia soulangeana</i>		
	Flower (<i>n</i>)	Leaf (<i>n</i>)	<i>t</i> -test	Flower	Leaf	<i>t</i> -test
Tree height (m)	8.48 \pm 0.51 (5)		-	7.42 \pm 0.68		-
DBH (cm)	16.63 \pm 0.44 (5)		-	12.91 \pm 0.62		-
WD (g cm ⁻³)	0.42 \pm 0.02 (5)		-	0.47 \pm 0.01		-
Single tepal or leaf area (cm ²)	20.62 \pm 1.07 (18)	58.35 \pm 3.61 (24) †	***	42.57 \pm 3.61 †	58.69 \pm 4.12 †	***
A_L/A_S (m ² cm ⁻²)	0.17 \pm 0.04 (5)	0.62 \pm 0.05 (5)	***	0.22 \pm 0.03	0.71 \pm 0.06	***
Tepal or leaf thickness (mm)	2.12 \pm 0.17 (18) †	0.15 \pm 0.00 (24) †	***	2.17 \pm 0.27 †	0.14 \pm 0.00 †	***
Tepal thinnest thickness (mm)	0.20 \pm 0.01 (18)		-	0.21 \pm 0.02 †		-

SLA (cm ² g ⁻¹)	258.23 ± 24.12 (18)	335.31 ± 6.43 (12)	***	323.34 ± 24.22	352.43 ± 12.01	*
LDMC (%)	6.09 ± 0.15 (18) †	17.32 ± 0.31 (12)	***	6.24 ± 0.11 †	18.28 ± 0.45	***
N (%)	2.97 ± 0.02 (9)	2.86 ± 0.08 (12)	ns	2.62 ± 0.13	2.62 ± 0.06	ns
P (%)	0.38 ± 0.02 (9)	0.35 ± 0.01 (12)	ns	0.36 ± 0.01	0.34 ± 0.02	ns
Abaxial stomatal size (μm ²)	875.62 ± 47.05 (18)	561.22 ± 12.89 (24)	***	864.48 ± 36.46	513.65 ± 13.55	***
Adaxial stomatal size (μm ²)	998.68 ± 57.32 (18)		-	918.11 ± 44.60		-
Stomatal density (number mm ⁻²)	3.08 ± 0.31 (36)	277.33 ± 8.09 (24)	***	2.11 ± 0.22	244.67 ± 7.05	***
SPI (%)	1.00 ± 0.06 (18)	30.49 ± 1.00 (24)	***	0.65 ± 0.04	22.80 ± 0.63	***
<i>g</i> _{max} (mol m ⁻² s ⁻¹)	0.07 ± 0.00 (18)	2.93 ± 0.09 (24)	***	0.05 ± 0.00	2.31 ± 0.06	***
<i>g</i> _s (mol m ⁻² s ⁻¹)	0.021±.002 (120) †	0.078±.008 (120) †	***	0.041 ± .003 †	0.063 ± .003 †	***
<i>E</i> (mmol m ⁻² s ⁻¹)	0.28 ± 0.02 (120) †	1.30 ± 0.10 (120) †	***	0.51 ± 0.03 †	0.93 ± 0.05 †	***
<i>Ψ</i> _{am} (MPa)	-0.15 ± 0.02 (18) †	-0.35 ± 0.05 (12)	***	-0.13 ± 0.03 †	-0.53 ± 0.05	***
<i>Ψ</i> _{pm} (MPa)	-0.11 ± 0.02 (18) †	-0.57 ± 0.05 (12)	***	-0.08 ± 0.01 †	-0.76 ± 0.02	***
<i>Ψ</i> _{tlp} (MPa)	-0.27 ± 0.02 (18)	-0.82 ± 0.02 (12)	***	-0.22 ± 0.03 †	-0.77 ± 0.03	***
HSM (MPa)	0.12 ± 0.01 (18) †	0.24 ± 0.02 (12)	***	0.11 ± 0.03 †	0.02 ± 0.01	***
<i>C</i> _{fit} (mol m ⁻² MPa ⁻¹)	6.01 ± 0.48 (18)	0.78 ± 0.03 (12)	***	6.77 ± 1.45 †	0.60 ± 0.04	***
<i>K</i> _{leaf} or <i>K</i> _{tepal} (mmol m ⁻² s ⁻¹ MPa ⁻¹)	4.13 ± 0.70 (18) †	14.46 ± 1.23 (12) †	***	4.81 ± 0.83 †	12.11 ± 1.54 †	***
<i>K</i> _S (kg m ⁻¹ s ⁻¹ MPa ⁻¹)	4.37 ± 0.34 (10)	3.73 ± 0.44 (10)	ns	2.16 ± 0.42	2.33 ± 0.35	ns
<i>K</i> _L (10 ⁻⁴ kg m ⁻¹ s ⁻¹ MPa ⁻¹)	30.60 ± 6.24 (10)	4.42 ± 0.58 (10)	***	8.75 ± 1.24	3.76 ± 0.82	***

813 **Figure Legends**

814 **Fig. 1.** Daily changes of (a) vapour pressure deficit (D , closed circles) and solar
815 radiation (SR, open circles), (b) temperature (black triangles) and rainfall (black bars),
816 sap flux density (J_s) of (c) *Magnolia denudata* and (d) *Magnolia soulangeana*,
817 indicating the flowering and vegetative periods as grey areas in Feb and Mar,
818 respectively. The day that we carried out daily change measurements are marked as
819 D1 and D2 in panels (c) and (d).

820

821 **Fig. 2.** Daily curves of (a) vapour pressure deficit (D), (b) solar radiation (SR), and
822 sap flux density (J_s) of (c) *M. denudata* and (d) *M. soulangeana* on two sunny days
823 with only flowers (Feb-24, white dots) or only leaves (Mar-26, black dots) on the tree.

824

825 **Fig. 3.** Flower opening (a, b) and fading (c, d) stages for *M. denudata* and *M.*
826 *soulangeana*, respectively. Flower number records are based on the 16 branches used
827 for sap flow monitoring ($n = 6$ for *M. denudata*; $n = 10$ for *M. soulangeana*), data are
828 mean \pm SEM. Grey areas in (a) and (b) indicate flowering periods with stable ratios
829 for both opening and fading stages.

830

831 **Fig. 4.** Sap flux density (J_s) in relation to daytime vapour pressure deficit (D) during
832 the flowering (a, b) and vegetative (c, d) periods for *M. denudata* and *M. soulangeana*,
833 respectively. Grey crosses show raw data in ten minutes intervals from days when
834 flower opening ratios were stable and all leaves were expanded, as indicated by grey
835 areas in Figs 1 and 3, with data from rainy days, under limiting light (SR=0 W m⁻²)
836 and during low D (<0.1 kPa) filtered out. Boundary line analyses give the maximum
837 J_s at different SR gradients as low light (LL, black triangles/circles, solid lines,
838 SR=0~400 W m⁻²) and high light (HL, white triangles/circles, dash lines,
839 SR=400~800 W m⁻²). The relationships between J_s and $\ln D$ are: (a) *M. denudata*
840 flower, LL, not modelled; HL, $J_s=15.17-24.70 \times \ln D$; (b) *M. soulangeana* flower, LL,
841 not modelled; HL, $J_s=47.31-80.38 \times \ln D$; (c) *M. denudata* leaf, LL,
842 $J_s=50.41+28.25 \times \ln D$; HL, $J_s=96.87+42.10 \times \ln D$; and (d) *M. soulangeana* leaf, LL,
843 $J_s=39.94+18.43 \times \ln D$; HL, $J_s=73.39+4.45 \times \ln D$.

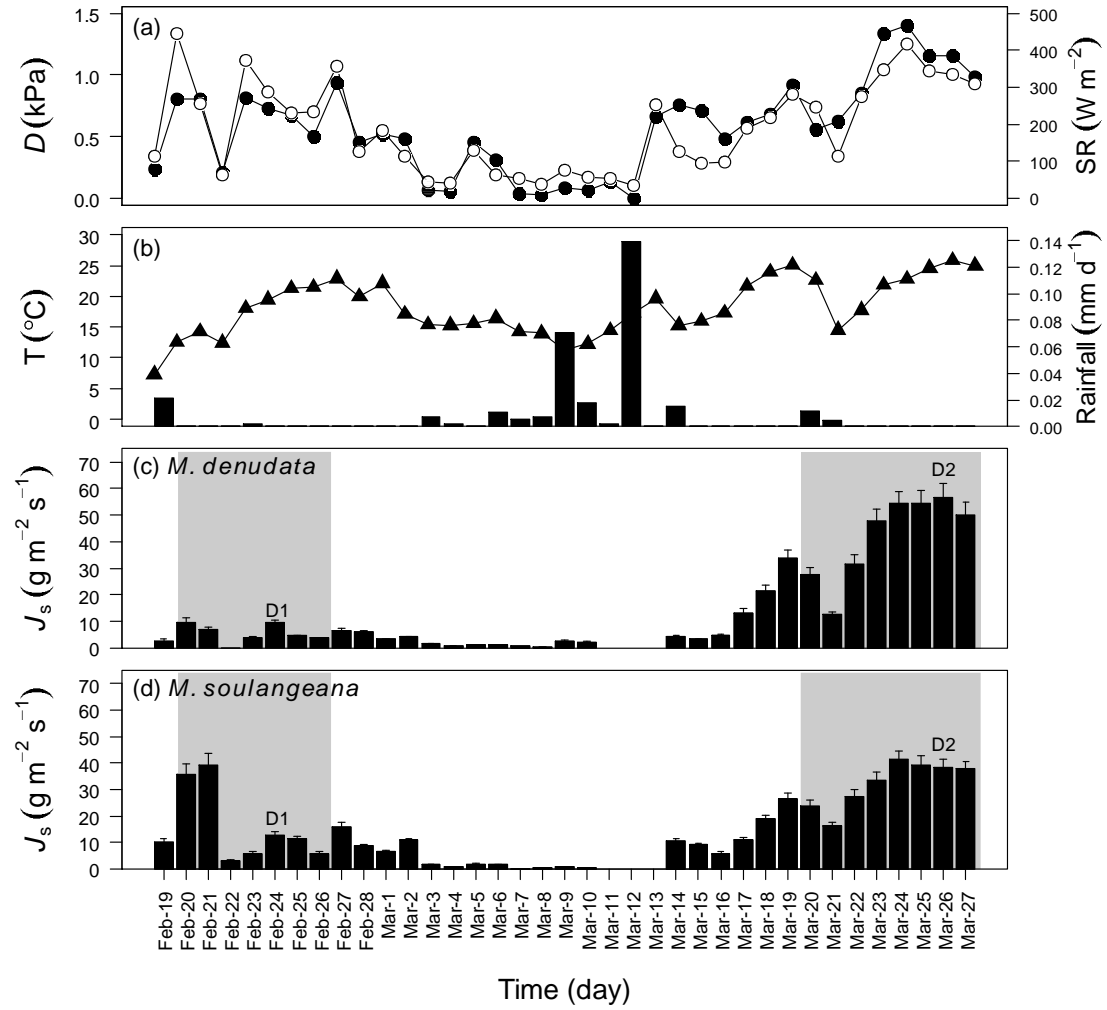
844

845 **Fig. 5.** Stomatal conductance (g_s) of flower (a, b) and leaf (c, d) in relation to air
846 vapour pressure deficit (D) in the morning and afternoon of two sunny days,

847 respectively. The relationships between g_s and $\ln D$ are modelled for *M. denudata*
848 (white triangles/circles, dashed lines) and *M. soulangeana* (black triangles/circles,
849 solid lines) separately: (a) *M. denudata*, $g_s=0.05-0.09 \times \ln D$; *M. soulangeana*, $g_s=0.08-$
850 $0.18 \times \ln D$; (b) *M. denudata*, $g_s=0.05-0.11 \times \ln D$; *M. soulangeana*, $g_s=0.09-0.21 \times \ln D$;
851 (c) *M. denudata*, $g_s=0.26-0.54 \times \ln D$; *M. soulangeana*, $g_s=0.10-0.19 \times \ln D$; and (d) *M.*
852 *denudata*, $g_s=0.34-0.33 \times \ln D$; *M. soulangeana*, $g_s=0.12-0.10 \times \ln D$. Note the axes
853 scales differ in each figure.

854

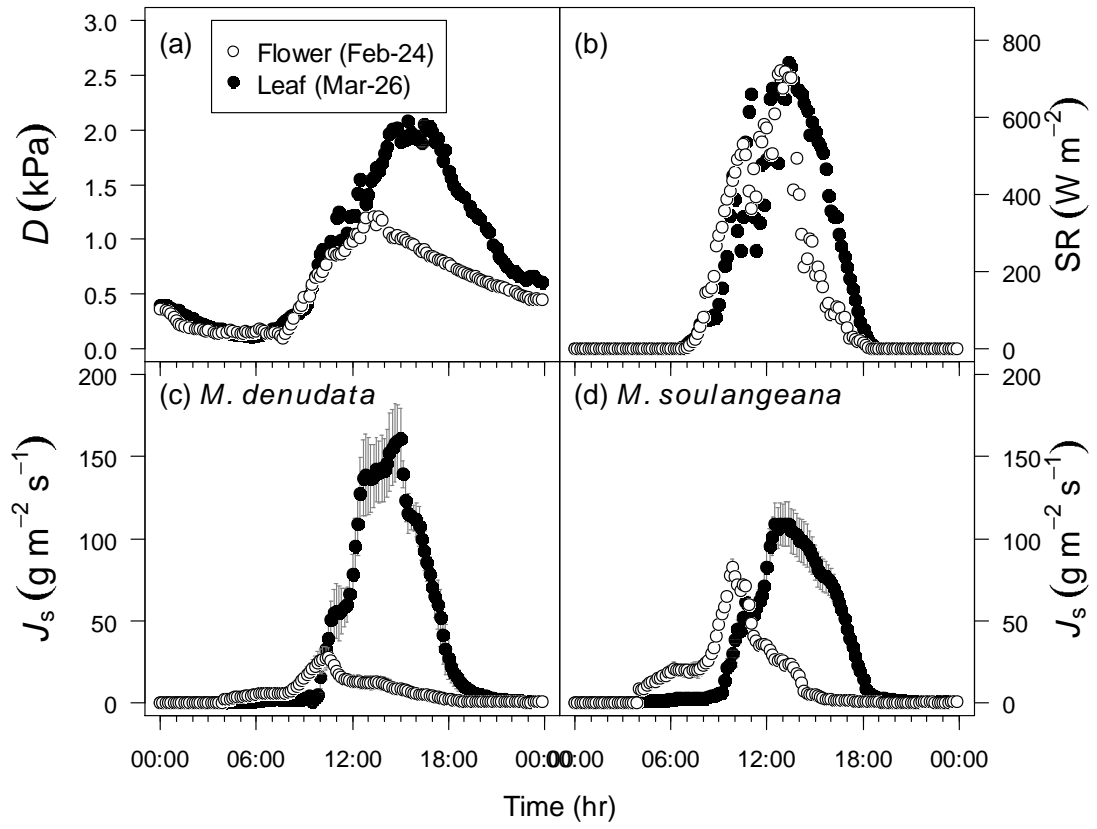
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857 **Fig. 1. Liu et al.**

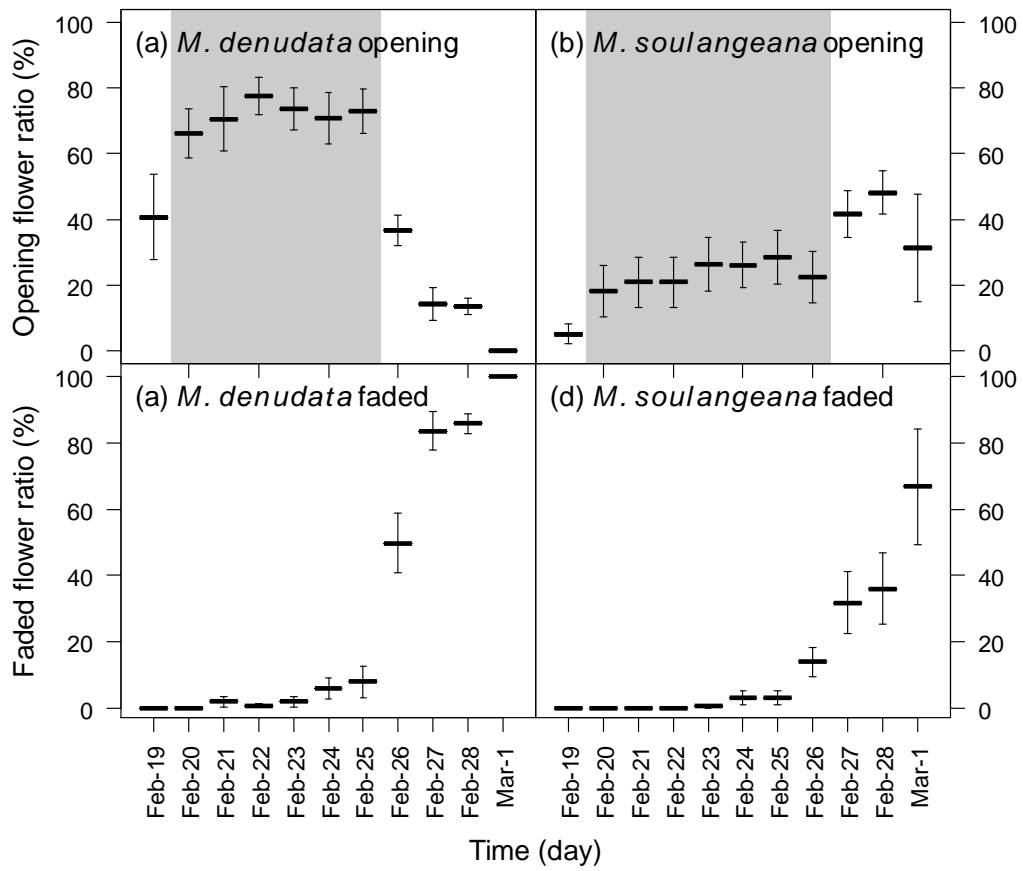
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860 **Fig. 2. Liu *et al.***

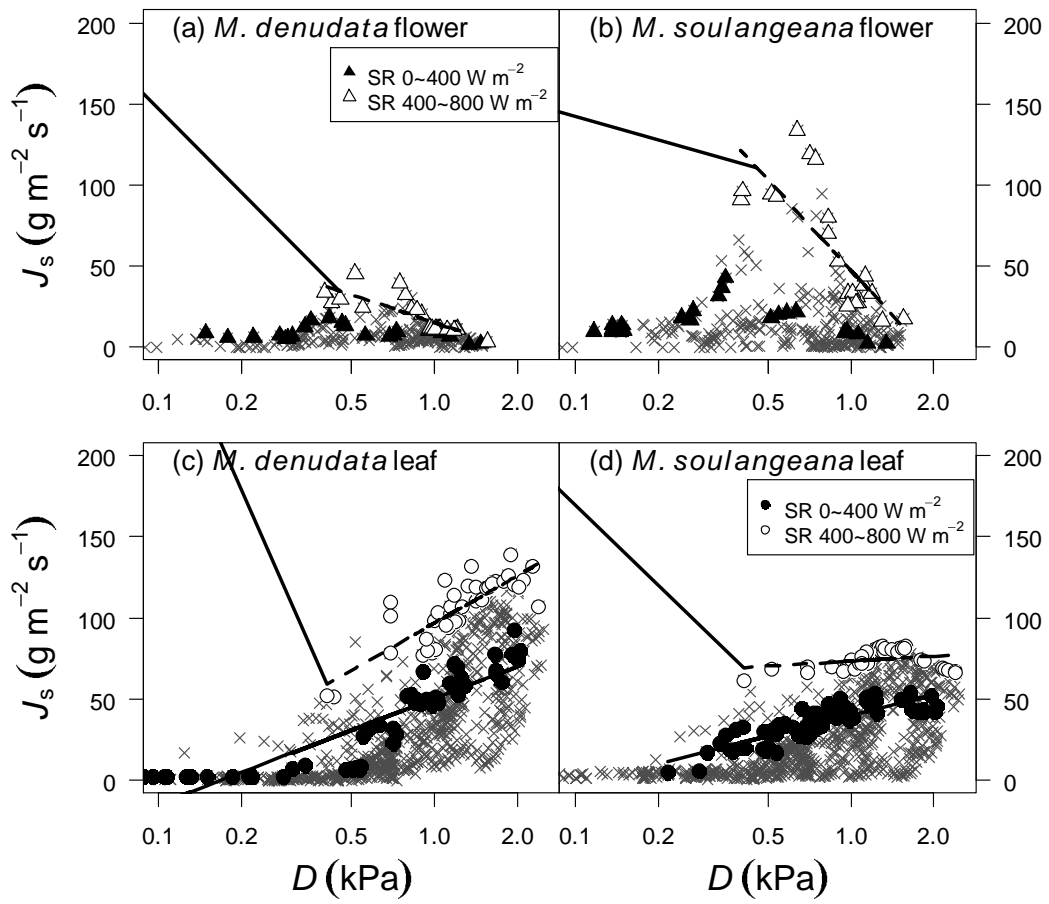
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862

863 **Fig. 3. Liu *et al.***

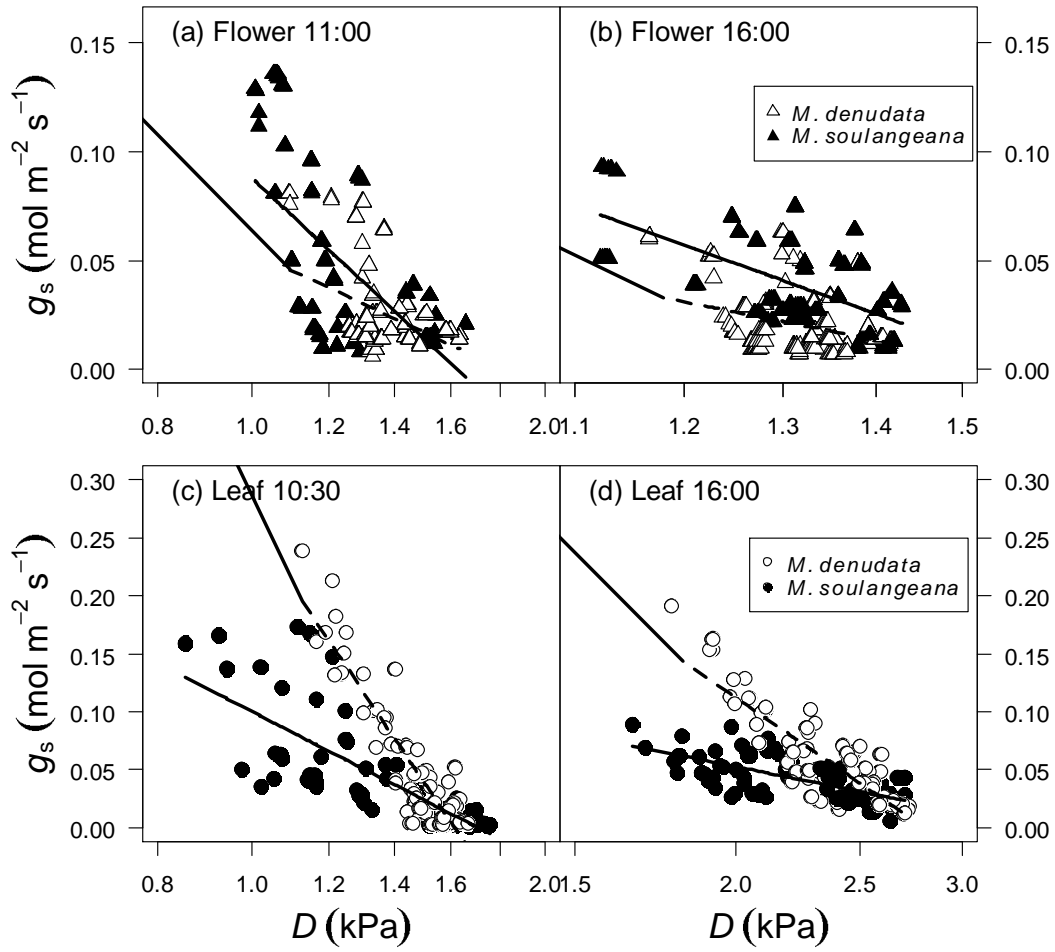
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866 **Fig. 4. Liu *et al.***

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868

869 **Fig. 5. Liu et al.**

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