

1 **Interaction of CO₂ concentrations and water stress in semi-**
2 **arid plants causes diverging response in instantaneous water**
3 **use efficiency and carbon isotope composition**

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9 **Abstract.** In the context of global warming attributable to the increasing levels of CO₂, severe drought
10 may be more frequent in areas with chronic water shortages (semi-arid areas). This necessitates research
11 on the interactions between increased levels of CO₂ and drought on plant photosynthesis. It is commonly
12 reported that ¹³C fractionation occurs as CO₂-gas diffuses from the atmosphere to the sub-stomatal cavity.
13 Few researchers have investigated ¹³C fractionation at the site of carboxylation to cytoplasm before
14 sugars are exported outward from the leaf. This process typically progresses in response to variations in
15 environmental conditions (i.e., CO₂ concentrations and water stress), including in their interaction.
16 Therefore, saplings of two typical plant species (*Platycladus orientalis* and *Quercus variabilis*) from
17 semi-arid areas of Northern China were selected and cultivated in growth chambers with orthogonal
18 treatments (four CO₂ concentration ([CO₂]) × five soil volumetric water content (SWC)). The δ¹³C of
19 water-soluble compounds extracted from leaves of saplings was determined for an assessment of
20 instantaneous water use efficiency (WUE_{cp}) after cultivation. Instantaneous water use efficiency derived
21 from gas-exchange measurement (WUE_{ge}) was integrated to estimate differences in δ¹³C signal variation
22 before leaf-level translocation of primary assimilates. The WUE_{ge} in *P. orientalis* and *Q. variabilis* both
23 decreased with increased soil moisture at 35–80% of field capacity (FC), and increased with elevated
24 [CO₂] by increasing photosynthetic capacity and reducing transpiration. Instantaneous water use
25 efficiency (iWUE) according to environmental changes, differed between the two species. The WUE_{ge}
26 in *P. orientalis* was significantly greater than that in *Q. variabilis*, while an opposite tendency was
27 observed when comparing WUE_{cp} between the two species. Total ¹³C fractionation at the site of
28 carboxylation to cytoplasm before sugar export (total ¹³C fractionation) was species-specific, as
29 demonstrated in the interaction of [CO₂] and SWC. Rising [CO₂] coupled with moistened soil generated
30 increasing disparities in δ¹³C between water-soluble compounds (δ¹³C_{WSC}) and estimates based on gas-
31 exchange observations (δ¹³C_{obs}) in *P. orientalis*, ranging between 0.0328–0.0472‰. Differences between
32 δ¹³C_{WSC} and δ¹³C_{obs} in *Q. variabilis* increased as [CO₂] and SWC increased (0.0384–0.0466‰). The ¹³C
33 fractionation from mesophyll conductance (g_m) and post-carboxylation both contributed to the total ¹³C
34 fractionation that was determined by δ¹³C of water-soluble compounds and gas-exchange measurements.
35 Total ¹³C fractionation was linearly dependent on stomatal conductance, indicating post-carboxylation
36 fractionation could be attributed to environmental variation. The magnitude and environmental
37 dependence of apparent post-carboxylation fractionation is worth our attention when addressing
38 photosynthetic fractionation.

39 **Key words:** Post-carboxylation fractionation; Carbon isotope fractionation; Elevated CO₂ concentration;
40 Soil volumetric water content; Instantaneous water use efficiency

41 **1 Introduction**

42 Since the industrial revolution, atmospheric CO₂ concentration has increased at an annual rate of 0.4%,
43 and is expected to increase to 700 μmol·mol⁻¹, culminating in more frequent periods of dryness (IPCC,
44 2014). Increasing atmospheric CO₂ concentrations that exacerbate the greenhouse effect will increase
45 fluctuations in global precipitation patterns, which will probably amplify drought frequency in arid
46 regions and lead to more frequent extreme flooding events in humid regions (Lobell et al., 2014).
47 Accompanying the increasing concentration of CO₂, mean δ¹³C of atmospheric CO₂ is currently being
48 depleted by 0.02‰–0.03‰ year⁻¹ (CU-INSTAAR/NOAACMDL network for atmospheric CO₂;
49 <http://www.esrl.noaa.gov/gmd/>).

50 The current carbon isotopic composition may respond to environmental change and its influence on
51 diffusion via plant physiological and metabolic processes (Gessler et al., 2014; Streit et al., 2013). While
52 depletion of δ¹³C_{CO₂} is occurring in the atmosphere, variations in CO₂ concentration ([CO₂]) may affect
53 δ¹³C of plant organs which, in turn, respond physiologically to changes in climate (Gessler et al., 2014).
54 The carbon discrimination (¹³Δ) in leaves could also provide timely feedback to the availability of soil
55 moisture and atmospheric vapor pressure deficit (Cernusak et al., 2012). Discrimination of ¹³C in leaves
56 relies mainly on environmental factors that affect the ratio of intercellular to ambient [CO₂] (C_i/C_a).
57 Rubisco activities and the mesophyll conductance derived from the difference of [CO₂] between
58 intercellular sites and chloroplasts are also involved (Farquhar et al., 1982; Cano et al., 2014). Changes
59 in environmental conditions affect photosynthetic discrimination, recording differentially in the δ¹³C of
60 water-soluble compounds (δ¹³C_{WSC}) in different plant organs. Several processes during photosynthesis
61 alter the δ¹³C of carbon transported within plants. Carbon-fractionation during photosynthetic CO₂
62 fixation has been reviewed elsewhere (Farquhar et al., 1982; Farquhar and Sharkey, 1982).

63 Post-photosynthetic fractionation is derived from equilibrium and kinetic isotopic effects that
64 determine isotopic differences between metabolites and intramolecular reaction positions. These are
65 defined as “post-photosynthetic” or “post-carboxylation” fractionation (Jäggi et al., 2002; Badeck et al.,
66 2005; Gessler et al., 2008). Post-carboxylation fractionation in plants includes the carbon discrimination
67 that follows carboxylation of ribulose-1, 5-bisphosphate and internal diffusion (RuBP, 27%), as well as
68 related transitory starch metabolism (Gessler et al., 2008; Gessler et al., 2014), fractionation-associated
69 phloem transport, remobilization or storage of soluble carbohydrates, and starch metabolism
70 fractionation in sink tissue (tree rings). In the synthesis of soluble sugars, ¹³C-depletions of triose
71 phosphates occur during export from the cytoplasm, and during production of fructose-1, as does 6-
72 bisphosphate by aldolase in transitory starch synthesis (Rossmann et al., 1991; Gleixner and Schmidt,
73 1997). Synthesis of sugars before transportation to the twig is associated with the post-carboxylation
74 fractionation generated in leaves. Although these are likely to play a role, another consideration is [CO₂]
75 in the chloroplast (C_c), not in the intercellular space, as considered in the simplified equation of
76 Farquhar’s model (Evans et al., 1986; Farquhar et al., 1989) is actually defined as carbon isotope
77 discrimination (δ¹³C). Differences between gas-exchange derived values and online measurements of
78 δ¹³C have often been used to estimate C_i-C_c and mesophyll conductance for CO₂ (Le Roux et al., 2001;
79 Warren and Adams, 2006; Flexas et al., 2006; Evans et al., 2009; Flexas et al., 2012; Evans and von
80 Caemmerer 2013). In this regard, changes in mesophyll conductance could be partly responsible for the
81 differences in the two measurements, as it generally increases in the short term in response to elevated

82 CO₂ (Flexas et al., 2014), but tends to decrease under drought (Hommel et al., 2014; Th eroux-Rancourt
83 et al., 2014). Therefore, it is necessary to avoid confusion between carbon isotope discrimination derived
84 from synthesis of soluble sugars and/or mesophyll conductance. The degree to which carbon
85 fractionation is related to environmental variation has yet to be fully investigated.

86 The simultaneous isotopic analysis of leaves allows determination of temporal variation in isotopic
87 fractionation (Rinne et al., 2016). This will aid in ^{an}the accurate recording of environmental conditions. -X
88 Newly assimilated carbohydrates can be extracted, and these are termed the water-soluble compounds
89 (WSCs) in leaves (Brandes et al., 2006; Gessler et al., 2009). WSCs can also be associated with an
90 assimilation-weighted mean of C_i/C_a (and C_d/C_a) photosynthesized over periods ranging from a few
91 hours to 1–2 days (Pons et al., 2009). However, there is disagreement whether fractionation caused by
92 post-carboxylation and/or mesophyll resistance can alter the stable signatures of leaf carbon and thence
93 influence instantaneous water use efficiency (iWUE). In addition, the manner in which iWUE derived
94 from isotopic fractionation responds to environmental factors, such as elevated [CO₂] and/or soil water
95 gradients, is unknown. largely -X

96 Consequently, we investigated the $\delta^{13}\text{C}$ of the fast-turnover carbohydrate pool in sapling leaves of two
97 tree species, *Platycladus orientalis* (L.) Franco and *Quercus variabilis* Bl., native to semi-arid areas of
98 China. We conducted gas-exchange measurements in controlled-environment growth chambers. One
99 goal is to differentiate the ¹³C fractionation from the site of carboxylation to cytoplasm prior to sugar
100 transportation in *P. orientalis* and *Q. variabilis*, which is the total ¹³C fractionation, determined from the -X
101 $\delta^{13}\text{C}$ of WSCs and gas-exchange measurements. Another goal is to discuss the potential causes for the
102 observed divergence, estimate contributions of post-photosynthesis and mesophyll conductance on these
103 differences, and describe how carbon isotopic fractionation responds to the interactive effects of elevated
104 [CO₂] and water stress.

105 2 Material and Methods

106 2.1 Study site and design

107 *P. orientalis* and *Q. variabilis* saplings, selected as experimental material, were obtained from the
108 Capital Circle forest ecosystem station, a part of Chinese Forest Ecosystem Research Network (CFERN),
109 40°03'45"N, 116°5'45"E, Beijing, China. This region is forested by *P. orientalis* and *Q. variabilis*. We the -X
110 chose saplings of similar basal diameters, heights, and growth class. Each sapling was placed into an
111 individual pot (22 cm diam. × 22 cm high). Undisturbed soil samples were collected from the field, sieved
112 (with particles >10 mm removed), and placed into the pots. The soil bulk density in the pots was
113 maintained at 1.337–1.447 g·cm⁻³. After a 30-day transplant recovery period, the saplings were placed
114 into growth chambers for orthogonal cultivation.

115 The controlled experiment was conducted in growth chambers (FH-230, Taiwan Hipoint Corporation,
116 Kaohsiung City, Taiwan). To reproduce the meteorological conditions of different growing seasons in
117 the research region, daytime and nighttime temperatures in the chambers were set to 25 ± 0.5°C from
118 07:00 to 17:00 and 18 ± 0.5°C from 17:00 to 07:00. Relative humidity was maintained at 60% and 80%
119 during the daytime and nighttime, respectively. The mean daytime light intensity was 200–240 μmol·m⁻²·s⁻¹.
120 The chamber system ^{was} designed to ~~be~~ control and monitor [CO₂]. Two growth chambers (A and -X
121 B) were used in this study. Chamber A maintained [CO₂] at 400 ppm (C₄₀₀) and 500 ppm (C₅₀₀). Chamber
122 B maintained [CO₂] at 600 ppm (C₆₀₀) and 800 ppm (C₈₀₀). The target [CO₂] in each chamber had a
123 standard deviation of ± 50 ppm during plant cultivation and testing.

124 An automatic watering device was used to irrigate the potted saplings to avoid heterogeneity when
125 scheduled watering was not made (Fig. 1). The watering device consisted of a water storage tank, holder,
126 controller, soil moisture sensors, and a drip irrigation component. Prior to use, the tank was filled with
127 water, and the soil moisture sensor was inserted to a uniform depth in the soil. After connecting the
128 controller to an AC power supply, target soil volumetric water content (SWC) could be set and monitored
129 by soil moisture sensors. Since changes in SWC could be sensed by the sensors, this automatic watering
130 device could be regulated to begin watering or stop watering the plants. One irrigation device was
131 installed per chamber. Based on mean field capacity (FC) of potted soil (30.70%), we established
132 orthogonal treatments of four [CO₂] × five SWC (Table. 1). In Table 1, A₁-A₄ denotes [CO₂] of 400
133 (C₄₀₀), 500 (C₅₀₀), 600 (C₆₀₀) and 800 ppm (C₈₀₀) in the chambers; B₁-B₅ denotes 35-45% (10.74-
134 13.81%), 50-60% (15.35-18.42%), 60-70% (18.42-21.49%), 70-80% (21.49-24.56%), and 100% of
135 FC (CK, 27.63-30.70%). Each orthogonal treatment of [CO₂] × SWC for two saplings per species was
136 repeated twice. Each treatment lasted 7 days. One pot was exposed in each of the [CO₂] × SWC
137 treatments. Pots in the chambers were rearranged every two days to promote uniform illumination.

138 2.2 Foliar gas exchange measurement

139 Fully expanded primary annual leaves of the saplings were measured with a portable infrared gas
140 photosynthesis system (LI-6400, Li-Cor, Lincoln, US) before and after the 7-day cultivation. Two
141 saplings per species were replicated per treatment (SWC × [CO₂]). For each sapling, four leaves were
142 sampled and four measurements were conducted on each leaf. Main photosynthetic parameters, such as
143 net photosynthetic rate (P_n) and transpiration rate (T_r), were measured. Based on theoretical
144 considerations of Von Caemmerer and Farquhar (1981), stomatal conductance (g_s) and intercellular [CO₂]
145 (C_i) were calculated by the Li-Cor software. Instantaneous water use efficiency via gas exchange (WUE_{ge})
146 was calculated as the ratio P_n / T_r .

147 2.3 Plant material collection and leaf water-soluble compounds extraction

148 Eight recently-expanded sun leaves were selected per sapling and homogenized in liquid nitrogen after
149 gas-exchange measurements were finished. For extraction of WSCs from the leaves (Gessler et al.,
150 2004), 50 mg of ground leaves and 100 mg of PVPP (polyvinylpyrrolidone) were mixed and
151 incubated in 1 mL distilled water for 60 min at 5 °C in a centrifuge tube. Each leaf sample was replicated
152 twice. The tubes containing the mixture were heated in 100 °C water for 3 min. After cooling to room
153 temperature, the supernatant of the mixture was centrifuged (12000 × g for 5 min) and 10 μL of
154 supernatant was transferred into a tin capsule and dried at 70 °C. Folded capsules were used for δ¹³C
155 analysis of WSCs. The samples of WSCs from leaves were combusted in an elemental analyzer
156 (EuroEA, HEKAtech GmbH, Wegberg, Germany) and analyzed with a mass-spectrometer
157 (DELTA^{plus}XP, ThermoFinnigan).

158 Carbon isotope signatures were expressed in δ-notation (parts per thousand), relative to the
159 international Pee Dee Belemnite (PDB) standard:

$$160 \delta^{13}\text{C} = \left(\frac{R_{\text{sample}}}{R_{\text{standard}}} - 1 \right) \times 1000 \quad (1)$$

161 where δ¹³C is the heavy isotope and R_{sample} and R_{standard} refer to the isotope ratio between the particular
162 substance and the corresponding standard, respectively. The precision of repeated measurements was
163 0.1 ‰.

164 2.4 Isotopic calculation

no indentation is needed except at the beginning of the paragraph

165 2.4.1 ^{13}C fractionation from the site of carboxylation to cytoplasm prior to sugar transportation

166 Based on the linear model of Farquhar and Sharkey (1982), the isotope discrimination, Δ , was
167 calculated as

$$168 \Delta = (\delta^{13}\text{C}_a - \delta^{13}\text{C}_{WSC}) / (1 + \delta^{13}\text{C}_{WSC}), \quad (2)$$

169 where $\delta^{13}\text{C}_a$ and $\delta^{13}\text{C}_{WSC}$ are the isotope signatures of ambient $[\text{CO}_2]$ in chambers and WSCs extracted
170 from leaves, respectively. The $C_i:C_a$ was determined by

$$171 C_i:C_a = (\Delta - a) / (b - a), \quad (3)$$

172 where C_i and C_a are the $[\text{CO}_2]$ within substomatal cavities and in growth chambers, respectively; a is
173 the fractionation occurring CO_2 diffusion in still air (4‰) and b refers to the discrimination during CO_2
174 fixation by ribulose 1,5- biphosphate carboxylase/oxygenase (Rubisco) and internal diffusion (30‰).
175 Instantaneous water use efficiency by gas-exchange measurement (WUE_{ge}) was calculated as

$$176 \text{WUE}_{ge} = P_n:T_r = (C_a - C_i) / 1.6\Delta e, \quad (4)$$

177 where 1.6 is the diffusion ratio of stomatal conductance for water vapor to CO_2 in chambers and Δe is
178 the difference between e_{if} and e_{am} , representing the extra- and intra-cellular water vapor pressure,
179 respectively:

$$180 \Delta e = e_{if} - e_{atm} = 0.611 \times e^{17.502T/(240.97+T)} \times (1 - \text{RH}), \quad (5)$$

181 where T and RH are the leaf-surface temperature and relative humidity, respectively. Combining Eqns.
182 (2, 3 and 4), the instantaneous water use efficiency was determined by the $\delta^{13}\text{C}_{WSC}$ of leaves, defined as:

$$183 \text{WUE}_{cp} = \frac{P_n}{T_r} = (1 - \varphi) (C_a - C_i) / 1.6\Delta e = C_a(1 - \varphi) \left[\frac{b - \delta^{13}\text{C}_a + (b+1)\delta^{13}\text{C}_{WSC}}{(b-a)(1 + \delta^{13}\text{C}_{WSC})} \right] / 1.6\Delta e, \quad (6)$$

184 where φ is the respiratory ratio of leaf carbohydrates to other organs at night (0.3).

185 Then the ^{13}C fractionation from the site of carboxylation to cytoplasm prior to sugar transportation
186 (defined as the total ^{13}C fractionation) was estimated by the observed $\delta^{13}\text{C}$ of WSCs from leaves ($\delta^{13}\text{C}_{WSC}$)
187 and the modeled $\delta^{13}\text{C}$ calculated from gas-exchange measurements ($\delta^{13}\text{C}_{model}$). The $\delta^{13}\text{C}_{model}$ was
188 calculated by Δ_{model} from Eqn. (2); Δ_{model} was determined by combining Eqns. (3 and 4) as

$$189 \Delta_{model} = (b - a) \left(1 - \frac{1.6\Delta e \text{WUE}_{ge}}{C_a} \right) + a, \quad (7)$$

$$190 \delta^{13}\text{C}_{model} = \frac{C_a - \Delta_{model}}{1 + \Delta_{model}}, \quad (8)$$

$$191 \text{Total } ^{13}\text{C fractionation} = \delta^{13}\text{C}_{WSC} - \delta^{13}\text{C}_{model}. \quad (9)$$

192 2.4.2 Method of estimating mesophyll conductance and the contribution of post-carboxylation
193 fractionation

194 CO_2 diffusion into photosynthetic sites includes two main processes. CO_2 first moves from ambient
195 air surrounding the leaf (C_a) through stomata to the sub-stomatic cavities (C_i). From sub-stomatic cavities,
196 CO_2 then moves to the sites of carboxylation within the chloroplast stroma (C_c) of the leaf mesophyll.
197 The latter procedure of diffusion is termed mesophyll conductance (g_m ; Flexas et al., 2008). The carbon
198 isotope discrimination was generated from the relative contribution of diffusion and carboxylation,
199 reflected by C_c to C_a . The carbon isotopic discrimination (Δ) can be presented as (Farquhar et al. 1982):

$$\Delta = a_b \frac{C_a - C_s}{C_a} + a \frac{C_s - C_i}{C_a} + (e_s + a_l) \frac{C_i - C_c}{C_a} + b \frac{C_c}{C_a} - \frac{eR_D + f\Gamma^*}{C_a}, \quad (10)$$

201 where C_a , C_s , C_i , and C_c are the $[\text{CO}_2]$ in the ambient air, at the boundary layer of the leaf, in the
 202 substomatal cavities, and at the sites of carboxylation, respectively; a_b is the CO_2 diffusional fractionation
 203 at the boundary layer (2.9‰); e_s is the discrimination for CO_2 diffusion when CO_2 enters in solution
 204 (1.1‰, at 25°C); a_l is the CO_2 diffusional fractionation in the liquid phase (0.7‰); e and f are carbon
 205 discriminations derived in dark respiration (R_D) and photorespiration, respectively; k is the carboxylation
 206 efficiency, and Γ^* is the CO_2 compensation point in the absence of dark respiration (Brooks and
 207 Farquhar, 1985).

208 When gas in the cuvette is well stirred during gas-exchange measurements, diffusion across the
 209 boundary layer ~~could be neglected~~ and Eqn. (10) can be written as

$$\Delta = a \frac{C_a - C_i}{C_a} + (e_s + a_l) \frac{C_i - C_c}{C_a} + b \frac{C_c}{C_a} - \frac{eR_D + f\Gamma^*}{C_a}. \quad (11)$$

211 There is no consensus about the value of e , although recent measurements estimate it as ranging
 212 from 0-4‰. The value of f has been estimated to range from 8-12‰ (Gillon and Griffiths, 1997;
 213 Igamberdiev et al., 2004; Lanigan et al., 2008). As the most direct factor, b influences the calculation of
 214 g_m , which is thought to be approximately 30‰ in higher plants (Guy et al., 1993).

215 The difference of $[\text{CO}_2]$ between substomatal cavities and chloroplasts is omitted, while diffusion
 216 related to dark-respiration and photorespiration are negligible and Eqn. (11) may be simplified to

$$\Delta_i = a + (b - a) \frac{C_i}{C_a}. \quad (12)$$

218 Eqn. (12) denotes the linear relationship between carbon discrimination and C_i/C_a . ~~That~~ ^{This} underlines
 219 subsequent comparison between expected Δ (originating from gas-exchange, Δ_i , and measured Δ_{obs}), ^{which can}
 220 ~~could~~ ^{can be used to} evaluate the differences of $[\text{CO}_2]$ between intercellular air and sites of carboxylation ^{associated}
 221 ^{with ^{13}C fractionation from mesophyll conductance.} Consequently, g_m is calculated by subtracting the
 222 Δ_{obs} of Eqn. (11) from Δ_i [Eqn. (12)]:

$$\Delta_i - \Delta_{obs} = (b - e_s - a_l) \frac{C_i - C_c}{C_a} + \frac{eR_D + f\Gamma^*}{C_a} \quad (13)$$

224 and P_n from Fick's first law is ~~presented by~~ ^{relates}

$$P_n = g_m(C_i - C_c). \quad (14)$$

226 Substituting Eqn. (14) into Eqn. (13) ~~we obtain~~ ^{gives us}

$$\Delta_i - \Delta_{obs} = (b - e_s - a_l) \frac{P_n}{g_m C_a} + \frac{eR_D + f\Gamma^*}{C_a}, \text{ and} \quad (15)$$

$$g_m = \frac{(b - e_s - a_l) \frac{P_n}{C_a}}{(\Delta_i - \Delta_{obs}) - \frac{eR_D + f\Gamma^*}{C_a}}. \quad (16)$$

229 In the calculation of g_m , terms of ^{respiration} ~~respiratory~~ and ^{photorespiration} ~~photorespiratory~~ could be ignored and e and f are
 230 assumed to be zero ~~or cancelled~~ in the calculation of g_m . ^{can}

231 Then Eqn. (16) can be rewritten as

$$g_m = \frac{(b-e_s-a_l) \frac{P_n}{C_a}}{\Delta_l - \Delta_{obs}} \quad (17)$$

Therefore, the contribution of post- carboxylation fractionation can be estimated by

$$\text{Contribution of post - carboxylation fractionation} = \frac{(\text{Total }^{13}\text{C fractionation} - \text{fractionation from mesophyll conductance})}{\text{Total }^{13}\text{C fractionation}} \times 100\%. \quad (18)$$

3 Results

3.1 Foliar gas exchange measurements

When SWC increased between the treatments, P_n , g_s and T_r in *P. orientalis* and *Q. variabilis* peaked at 70–80% of FC and for 100% of FC (Fig. 2). The C_i in *P. orientalis* rose as SWC increased. It peaked at 60–70% of FC and declined thereafter with increased SWC in *Q. variabilis*. The carbon uptake and C_i were significantly improved by elevated $[\text{CO}_2]$ at all SWC for the two species ($p < 0.05$). Greater increases of P_n in *P. orientalis* were found at 50–70% of FC from C_{400} to C_{800} , which was at 35–45% of FC in *Q. variabilis*. As water stress was reduced (at 70–80% and 100% of FC), reduction of g_s in *P. orientalis* was more pronounced with elevated $[\text{CO}_2]$ at a given SWC ($p < 0.01$). Nevertheless, g_s in *Q. variabilis* for C_{400} , C_{500} , and C_{600} was significantly higher than that for C_{800} at 50–80% of FC ($p < 0.01$). Coordinated with g_s , T_r of the two species for C_{400} and C_{500} was significantly higher than that for C_{600} and C_{800} , except at 35–60% of FC ($p < 0.01$, Figs. 2g and 2h). P_n , g_s , C_i and T_r in *Q. variabilis* was significantly greater than the corresponding values in *P. orientalis* ($p < 0.01$, Fig. 2).

3.2 $\delta^{13}\text{C}$ of water-soluble compounds in leaves

After observations of photosynthetic traits in leaves of the two species, the same leaves were immediately frozen and WSCs were extracted for all orthogonal treatments. The carbon isotope composition of WSCs ($\delta^{13}\text{C}_{\text{WSC}}$) of both species increased as SWC increased (Figs. 3a and 3b, $p < 0.01$). The mean $\delta^{13}\text{C}_{\text{WSC}}$ of *P. orientalis* and *Q. variabilis* ranged from $-27.44 \pm 0.155\text{‰}$ to $-26.71 \pm 0.133\text{‰}$, and from $-27.96 \pm 0.129\text{‰}$ to $-26.49 \pm 0.236\text{‰}$, respectively. The photosynthetic capacity varied with increased SWC and the mean $\delta^{13}\text{C}_{\text{WSC}}$ of the two species, reaching their respective maxima at 70–80% of FC. With gradual enrichment of $[\text{CO}_2]$, mean $\delta^{13}\text{C}_{\text{WSC}}$ in both species declined when $[\text{CO}_2]$ exceeded 600 ppm ($p < 0.01$). Except for C_{400} at 50–100% of FC, the $\delta^{13}\text{C}_{\text{WSC}}$ in *P. orientalis* was significantly higher than that in *Q. variabilis* for most $[\text{CO}_2] \times \text{SWC}$ treatments ($p < 0.01$, Fig. 3).

3.3 Estimations of WUE_{ge} and WUE_{cp}

Figure 4a shows that increments of WUE_{ge} in *P. orientalis* under severe drought (i.e., 35–45% of FC) were highest for most $[\text{CO}_2]$, ranging from 90.7% to 564.65%. The WUE_{ge} in *P. orientalis* decreased as SWC increased and increased as $[\text{CO}_2]$ increased. Differing from variation in WUE_{ge} in *P. orientalis* with moistened soil, WUE_{ge} in *Q. variabilis* increased slightly at 100% of FC for C_{600} or C_{800} (Fig. 4b). The maximum WUE_{ge} occurred at 35–45% of FC for C_{800} among all orthogonal treatments associated with both species. Elevated $[\text{CO}_2]$ enhanced the WUE_{ge} in *Q. variabilis* at any SWC, except at 60–80% of FC. Thirty-two saplings of *P. orientalis* had greater WUE_{ge} than did *Q. variabilis* for the same $[\text{CO}_2] \times \text{SWC}$ treatments ($p < 0.05$).

As illustrated in Fig. 5a, WUE_{cp} in *P. orientalis* for C_{600} or C_{800} increased as water stress was alleviated beyond 50–60% of FC, as well as that for C_{400} or C_{500} , while SWC exceeded 60–70% of FC. *Q. variabilis* showed variable WUE_{cp} with increasing SWC (Fig. 5b). Except for C_{400} , WUE_{cp} in *Q. variabilis*

271 decreased abruptly at 50–60% of FC, and then increased as SWC increased for C₅₀₀, C₆₀₀, and C₈₀₀. In
272 contrast to the results for WUE_{ge}, WUE_{cp} in *Q. variabilis* was more pronounced than in *P. orientalis*
273 among all orthogonal treatments.

274 3.4 ¹³C fractionation from the site of carboxylation to cytoplasm before sugar transportation

275 We evaluated the total ¹³C fractionation from the site of carboxylation to the cytoplasm by gas-
276 exchange measurements and WSCs in leaves (Table 2), which can help track the path of ¹³C fractionation
277 in leaves. Comparing $\delta^{13}C_{WSC}$ with $\delta^{13}C_{model}$ from Eqns. (4, 7–9), the total ¹³C fractionation in *P. orientalis*
278 ranged from 0.0328 to 0.0472‰, which was less than that in *Q. variabilis* (0.0384 to 0.0466‰). The
279 total fractionation in *P. orientalis* was magnified with increasing SWC especially when SWC reached
280 35–80% of FC from C₄₀₀ to C₈₀₀ (increased by 21.30–42.04%). The total fractionation for C₄₀₀ and C₅₀₀
281 were amplified as SWC increased until 50–60% of FC in *Q. variabilis*, whereas they were increased at
282 50–80% of FC and decreased at 100% of FC for C₆₀₀ and C₈₀₀. Elevated [CO₂] enhanced the mean total
283 fractionation in *P. orientalis*, while fractionation in *Q. variabilis* declined sharply from C₆₀₀ to C₈₀₀. Total
284 ¹³C fractionation, with increased SWC, in *P. orientalis* increased more rapidly than it did in *Q. variabilis*.

285 3.5 *g_m* imposed on the interaction of CO₂ concentration and water stress

286 A comparison between online leaf $\delta^{13}C_{WSC}$ and the values of gas-exchange measurements is given to
287 estimate the *g_m* over all treatments in Fig. 6 [Eqns. (10–17)]. A significant increasing trend occurred in
288 *g_m* with decreasing water stress in *P. orientalis*, ranging from 0.0091–0.0690 mol·CO₂·m⁻²·s⁻¹ (*p* < 0.05),
289 which reached a maximum at 100% of FC under a given [CO₂]. Increases in *g_m* in *Q. variabilis* with
290 increasing SWC were not significant, except those under C₄₀₀. With increasing [CO₂], *g_m* in the two
291 species increased at different rates. With *P. orientalis* under C₄₀₀, *g_m* increased gradually and reached a
292 maximum under C₈₀₀ at 35–60% and 100% of FC (*p* < 0.05). However, that was maximized under C₆₀₀
293 (*p* < 0.05) and reduced under C₈₀₀ at 60–80% of FC. The maximum increment in *g_m* (8.2–58.4%) occurred
294 at C₈₀₀ at all SWC for *Q. variabilis*. The *g_m* in *Q. variabilis* was clearly greater than that in *P. orientalis*
295 under the same treatments, conditions.

296 3.6 Contribution of post-carboxylation fractionation

297 We evaluated the difference between Δ_i and Δ_{obs} in ¹³C fractionation derived from mesophyll
298 conductance. The post-photosynthetic fractionation after carboxylation can be calculated by subtracting
299 *g_m*-sourced fractionation from the total ¹³C fractionation (Table 2). The *g_m*-sourced fractionation provided
300 a smaller contribution to the total ¹³C fractionation than did post-carboxylation fractionation irrespective
301 of treatment (Table 2). The *g_m*-sourced fractionation in the two species illustrated different variations
302 with increasing SWC, which declined at 50–80% of FC and increased at 100% of FC in *P. orientalis*;
303 yet, in *Q. variabilis*, it increased with water stress alleviation at 50–80% of FC and then decreased at
304 100% of FC. Nevertheless, in the two species post-carboxylation fractionation in leaves all increased as
305 SWC increased. The *g_m*-sourced fractionation in *P. orientalis* and *Q. variabilis* reached their peaks under
306 C₆₀₀ and C₈₀₀, respectively. Post-carboxylation fractionation was magnified with increases in [CO₂] in
307 *P. orientalis*, and reached a maximum under C₆₀₀ and then declined under C₈₀₀.

308 3.7 Relationship between *g_s*, *g_m* and total ¹³C fractionation

309 Total ¹³C fractionation may be correlated with resistances associated with stomata and mesophyll cells.
310 We performed linear regressions between *g_s*/*g_m* and total ¹³C fractionation in *P. orientalis* and *Q.*
311 *variabilis* (Fig. 7 and 8). The total ¹³C fractionation was correlated to *g_s* (*p* < 0.01). The positive linear
312 relationships between *g_m* and total ¹³C fractionation (*p* < 0.01) indicated that the variation of [CO₂]

313 through the chloroplast was correlated with carbon discrimination following leaf photosynthesis.

314 4 Discussion

315 4.1 Photosynthetic traits

316 The exchange of CO₂ and water vapor via stomata can be modulated by the soil/leaf water potential
317 (Robredo et al., 2010). Saplings of *P. orientalis* reached maximum P_n and g_s at 70–80% of FC irrespective
318 of [CO₂] treatments. As SWC exceeded this soil water threshold, elevated CO₂ caused a greater reduction
319 in g_s , as was previously reported for barley and wheat (Wall et al., 2011). The decrease in g_s responding
320 to elevated [CO₂], could be mitigated with ^{increases in} increased SWC. The C_i in *Q. variabilis* peaked at 60–70% of
321 FC and then declined as soil moisture increased (Wall et al., 2006; Wall et al., 2011). This may be because
322 stomata tend to maintain a constant C_i or C_i/C_a when ambient [CO₂] is increased, which would determine
323 the amount of CO₂ used directly in the chloroplast (Yu et al., 2010). This result could be explained as
324 stomatal limitation (Farquhar and Sharkey, 1982; Xu, 1997). However, C_i in *P. orientalis* increased
325 considerably, while SWC exceeded 70–80% of FC, as found by Mielke et al. (2000). One possible
326 contributing factor is plants close their stomata to reduce water loss during organic matter synthesis
327 simultaneously decreasing the availability of CO₂ and generating respiration of organic matter (Robredo
328 et al., 2007). Another possible explanation is that the limited root volume of potted plants may be unable
329 to absorb sufficient water to support the full growth of shoots (Leakey et al., 2009; Wall et al., 2011). In
330 the present study, increasing [CO₂] may cause nonstomatal limitation when SWC exceeds a soil moisture
331 threshold of 70–80% of FC. The accumulation of nonstructural carbohydrates in leaf tissue may induce
332 mesophyll-based and/or biochemical-based transient inhibition of photosynthetic capacity (Farquhar and
333 Sharkey, 1982). Xu and Zhou (2011) developed a five-level SWC gradient to examine the effect of water
334 on the physiology of a perennial, *Leymus chinensis*, and demonstrated that there was a clear maximum
335 in SWC, below which the plant could adjust to changing environmental conditions. Micanda-Apodaca
336 et al. (2014) also concluded that in suitable water conditions, elevated CO₂ levels augmented CO₂
337 assimilation in herbaceous plants.

338 The P_n of the two woody plant species increased with elevated [CO₂] similar to results seen with other
339 C₃ woody plants (Kgope et al., 2010). Increasing [CO₂] alleviated severe drought and the need for heavy
340 irrigation, suggesting that photosynthetic inhibition produced by a lack or excess of water may be
341 mediated by increased [CO₂] (Robredo et al., 2007; Robredo et al., 2010) and ameliorate the effects of
342 drought stress by reducing plant transpiration (Kirkham, 2016; Kadam et al., 2014; Micanda-Apodaca et
343 al., 2014; Tausz-Posch et al., 2013).

344 4.2 Differences between WUE_{ge} and WUE_{cp}

345 Increases in WUE_{ge} in *P. orientalis* and *Q. variabilis* that resulted from the combination of P_n increase
346 and g_s decrease were followed by a reduction in T_r (Figs. 2a, 2g, 2b and 2h). This result was also
347 demonstrated by Ainsworth and McGrath (2010). Comparing P_n and T_r in the two species, a lower WUE_{ge}
348 in *Q. variabilis* was obtained due to its ^{different} physiological and morphological traits, such as larger leaf area,
349 rapid growth, and higher stomatal conductance than that in *P. orientalis* (Adiredjo et al., 2014). Medlyn
350 et al. (2001) reported that stomatal conductance of broadleaved species is more sensitive to elevated
351 [CO₂] than conifer species. There is no agreement on the patterns of iWUE at the leaf level, related to
352 SWC (Yang et al., 2010). The WUE_{ge} in *P. orientalis* and *Q. variabilis* were enhanced with soil drying,
353 as presented by Parker and Pallardy (1991), DeLucia and Heckathorn (1989), Reich et al. (1989), and
354 Leakey (2009).

355 Böggelein et al. (2012) confirmed that WUE_{cp} was more consistent with daily mean WUE_{ge} than with
356 WUE_{phloem} (calculated with the $\delta^{13}C$ of phloem). The WUE_{cp} of the two species demonstrated similar
357 variations to those in $\delta^{13}C_{WSC}$, which differed from those of WUE_{ge} . Pons et al. (2009) noted that Δ of
358 leaf soluble sugar is coupled with environmental dynamics over a period ranging from a few hours to
359 1–2 days. The WUE_{cp} of our materials responded to $[CO_2] \times SWC$ treatments over a number of
360 cultivated days, whereas WUE_{ge} was characterized as the instantaneous physiological change in plants
361 to new conditions. Consequently, WUE_{cp} and WUE_{ge} had different degrees of variations in response to
362 different treatments.

cultivation -X
-X

363 4.3 Influence of mesophyll conductance on the fractionation after carboxylation

364 Mesophyll conductance, g_m , has been identified to coordinate with environmental factors more rapidly
365 than stomatal conductance (Galmés et al., 2007; Tazoe et al., 2011; Flexas et al., 2007). During our 7-
366 day cultivations, g_m increased and WUE_{ge} decreased with increasing SWC. It has been documented that
367 g_m can improve WUE under drought pretreatment (Han et al., 2016). However, the mechanism by which
368 g_m responds to the fluctuation of $[CO_2]$ is unclear. Terashima et al. (2006) demonstrated that CO_2
369 permeable aquaporin, located in the plasma membrane and inner envelope of chloroplasts, could regulate
370 the change in g_m . In our study, g_m is species-specific to the $[CO_2]$ gradient. The g_m in *P. orientalis*
371 significantly decreased by 9.0%–44.4% from C_{600} to C_{800} at 60–80% of FC; these are similar to the results
372 of Flexas et al. (2007). A larger g_m in *Q. variabilis* under C_{800} was observed compared to *P. orientalis*.

373 Furthermore, g_m contributed to the total ^{13}C fractionation that followed carboxylation, while
374 photosynthate had not been transported to the sapling twigs. The ^{13}C fractionation of CO_2 from the air
375 surrounding the leaf to sub-stomatal cavities may be simply explained by stomatal resistance, which also
376 contains the fractionation derived from mesophyll conductance between sub-stomatic cavities and the
377 site of carboxylation in the chloroplast that cannot be neglected and should be elucidated (Pons et al.,
378 2009; Cano et al., 2014). In estimating the post-carboxylation fractionation, g_m -sourced fractionation
379 must be subtracted from the total ^{13}C fractionation (the difference between $\delta^{13}C_{WSC}$ and $\delta^{13}C_{model}$), which
380 is closely associated with g_m (Fig. 8, $p = 0.01$). Variations in g_m -sourced fractionation are coordinated
381 with those in g_m with changing environmental conditions (Table 2).

382 4.4 Post-carboxylation fractionation generated before photosynthate moves out of leaves

383 Photosynthesis, a biochemical and physiological process (Badeck et al., 2005), is characterized by
384 discrimination in ^{13}C , which leaves an isotopic signature in the photosynthetic apparatus. Farquhar et al.
385 (1989) reviewed the carbon-fractionation in leaves and covered the significant aspects of photosynthetic
386 carbon isotope discrimination. The post-carboxylation/photosynthetic fractionation associated with the
387 metabolic pathways of non-structural carbohydrates (NSC; defined here as soluble sugars + starch)
388 within leaves, and fractionation during translocation, storage, and remobilization prior to tree ring
389 formation is unclear (Epron et al., 2012; Gessler et al., 2014; Rinne et al., 2016). The synthesis of sucrose
390 and starch before transportation to twigs falls within the domain of post-carboxylation fractionation
391 generated in leaves. Hence, we hypothesized that ^{13}C fractionation might exist. When we completed the
392 leaf gas-exchange measurements, leaf samples were collected immediately to determine the $\delta^{13}C_{WSC}$.
393 Presumably, ^{13}C fractionation generated in the synthetic processes of sucrose and starch was contained
394 within the ^{13}C fractionation from the site of carboxylation to cytoplasm before sugar transportation.
395 Comparing $\delta^{13}C_{WSC}$ with $\delta^{13}C_{obs}$, the total ^{13}C fractionation in *P. orientalis* ranged from 0.0328 to
396 0.0472%, which was somewhat less than that in *Q. variabilis* (from 0.0384 to 0.0466%). Post-
397 carboxylation fractionation contributed 75.30–98.9% to total ^{13}C fractionation, determined by subtracting

max
^

398 the fractionation in g_m from total ^{13}C fractionation. Gessler et al. (2004) reviewed the environmental
399 components of variation in photosynthetic carbon isotope discrimination in terrestrial plants. Total ^{13}C
400 fractionation in *P. orientalis* was enhanced by the increase in SWC, consistent with that in *Q. variabilis*,
401 except at 100% of FC. The ^{13}C isotope signature in *P. orientalis* was depleted with elevated $[\text{CO}_2]$. Yet,
402 ^{13}C -depletion was weakened in *Q. variabilis* for C_{600} and C_{800} . Linear regressions between g_s and total
403 ^{13}C fractionation indicated that the post-carboxylation fractionation in leaves depends on the variation of
404 g_s and that stomata aperture was correlated with environmental change.

405 5 Conclusions

406 Through orthogonal treatments of **four $[\text{CO}_2]$ \times five SWC**, WUE_{cp} calculated by $\delta^{13}\text{C}_{\text{WSC}}$ and WUE_{ge}
407 derived from simultaneous leaf gas-exchange, were estimated to differentiate the $\delta^{13}\text{C}$ signal variation
408 before leaf-level translocation of primary assimilates. The influence of g_m on ^{13}C fractionation between
409 the sites of carboxylation and ambient air is important. It requires consideration when testing the
410 hypothesis that the post-carboxylation contributes to the ^{13}C fractionation from the site of carboxylation
411 to cytoplasm before sugar transport. In response to the interactive effects of $[\text{CO}_2]$ and SWC, WUE_{ge} in
412 the two tree species both decreased with increasing SWC, and increased with elevated $[\text{CO}_2]$ at **35–80%**
413 of FC. We concluded that relative soil drying, coupled with elevated $[\text{CO}_2]$, can improve WUE_{ge} by
414 strengthening photosynthetic capacity and reducing transpiration. WUE_{ge} in *P. orientalis* was
415 significantly greater than that in *Q. variabilis*, while the opposite was the case for WUE_{cp} . The g_m and
416 post-carboxylation both contributed to the total ^{13}C fractionation. Rising $[\text{CO}_2]$ and/or moistening soil
417 generated increasing disparities between $\delta^{13}\text{C}_{\text{WSC}}$ and $\delta^{13}\text{C}_{\text{model}}$ in *P. orientalis*; nevertheless, the
418 differences between $\delta^{13}\text{C}_{\text{WSC}}$ and $\delta^{13}\text{C}_{\text{model}}$ in *Q. variabilis* increased when $[\text{CO}_2]$ was less than 600 ppm
419 and/or water stress was alleviated. Total ^{13}C fractionation in the leaf was linearly dependent on g_s . With
420 respect to carbon isotope fractionation in post-carboxylation and transportation processes, we note that
421 ^{13}C fractionation derived from the synthesis of sucrose and starch is likely influenced by environmental
422 changes. A clear description of the magnitude and environmental dependence of post-carboxylation
423 fractionation is worth considering.

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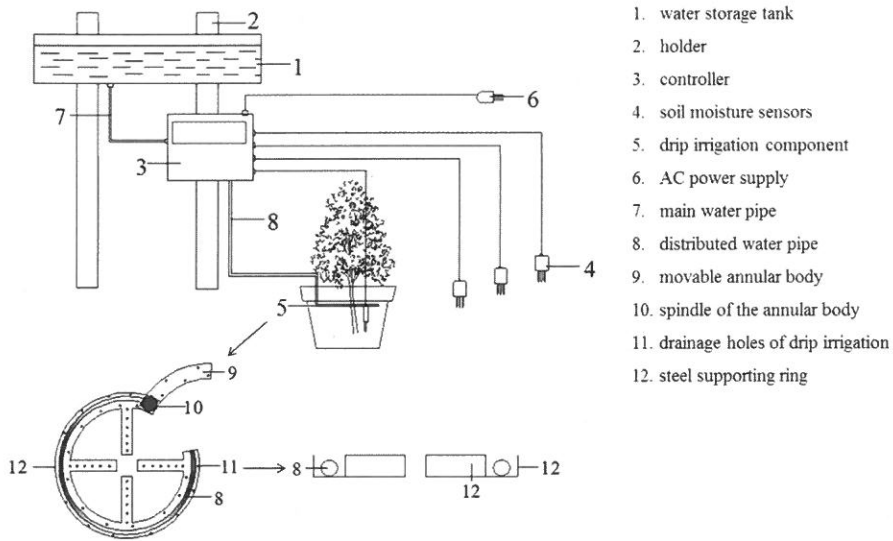
618 **Author contributions**

619 N. Zhao and Y. He collected field samples, and performed the experiments. N. Zhao analyzed the data
620 and wrote the paper. P. Meng commented on the theory and study design. X. Yu revised and edited the
621 manuscript.

622

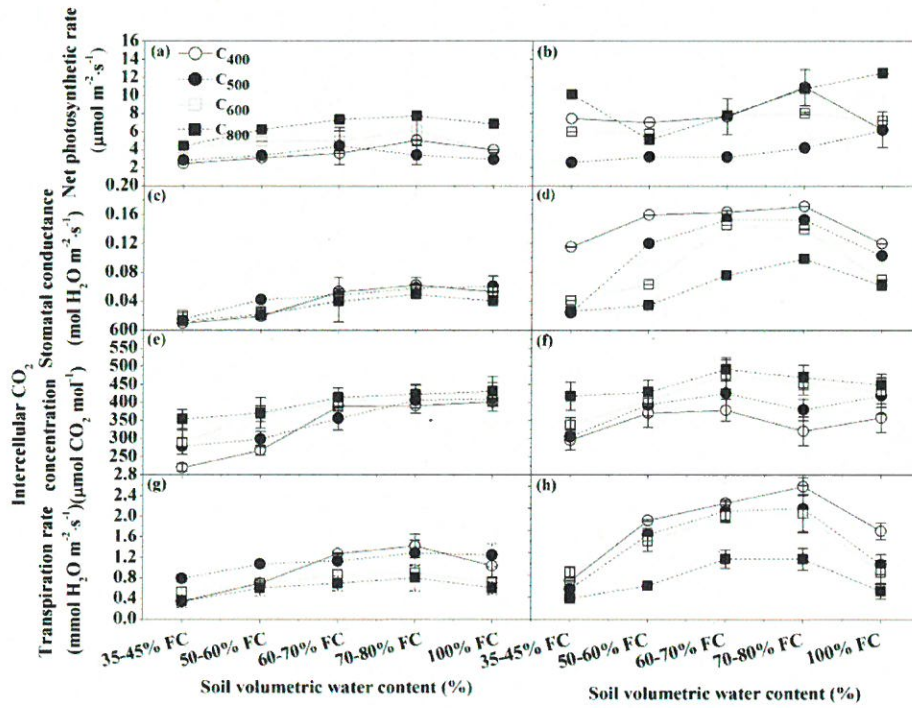
623 *Acknowledgements.* Financial support for this project was provided by the National Natural Science
624 Foundation of China (grant No. 41430747) and the Beijing Municipal Education Commission (CEFF-
625 PXM2017_014207_000043). We thank Beibei Zhou and Yuanhai Lou for collection of materials and
626 management of saplings. We are grateful to anonymous reviewers for constructive suggestions regarding
627 this manuscript. Due to space limitations we cited selected references involving this study topic and
628 apologize for authors whose work was not cited.

Figure

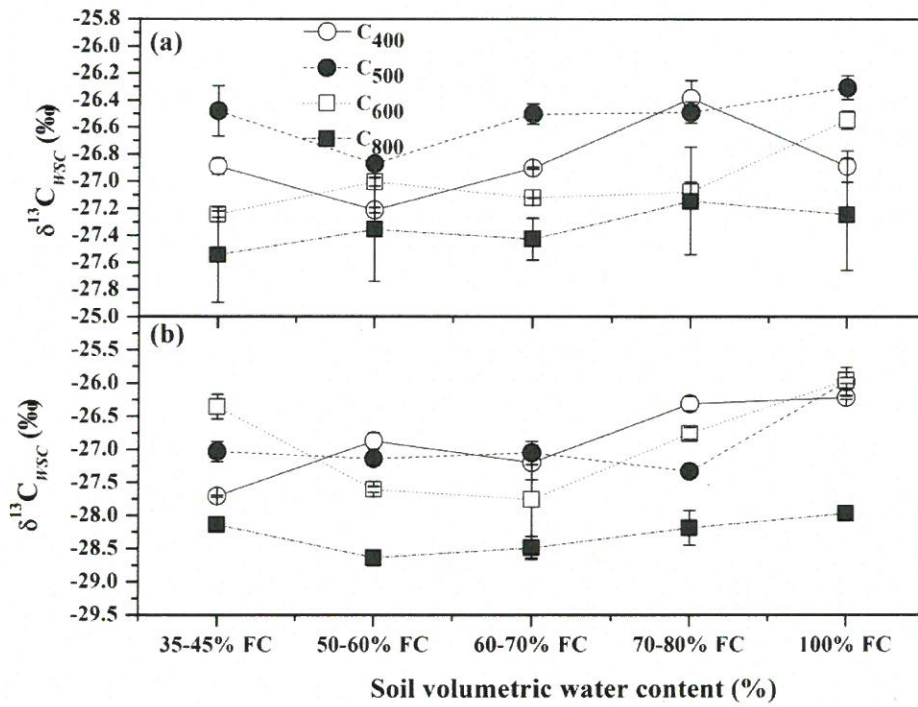


- 1. water storage tank
- 2. holder
- 3. controller
- 4. soil moisture sensors
- 5. drip irrigation component
- 6. AC power supply
- 7. main water pipe
- 8. distributed water pipe
- 9. movable annular body
- 10. spindle of the annular body
- 11. drainage holes of drip irrigation
- 12. steel supporting ring

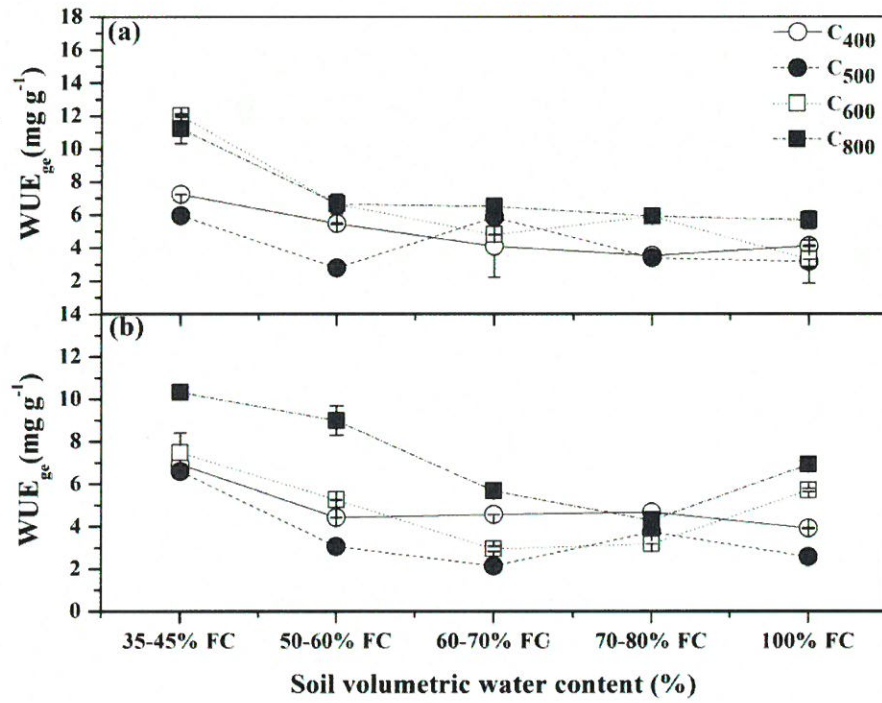
630 **Figure 1.** Diagram of the automatic drip irrigation device used in this study; numbers indicate the
 631 individual parts of the irrigation device (No. 1–12). The lower-left corner of this figure presents the
 632 detailed schematic for the drip irrigation component (No. 8–12).



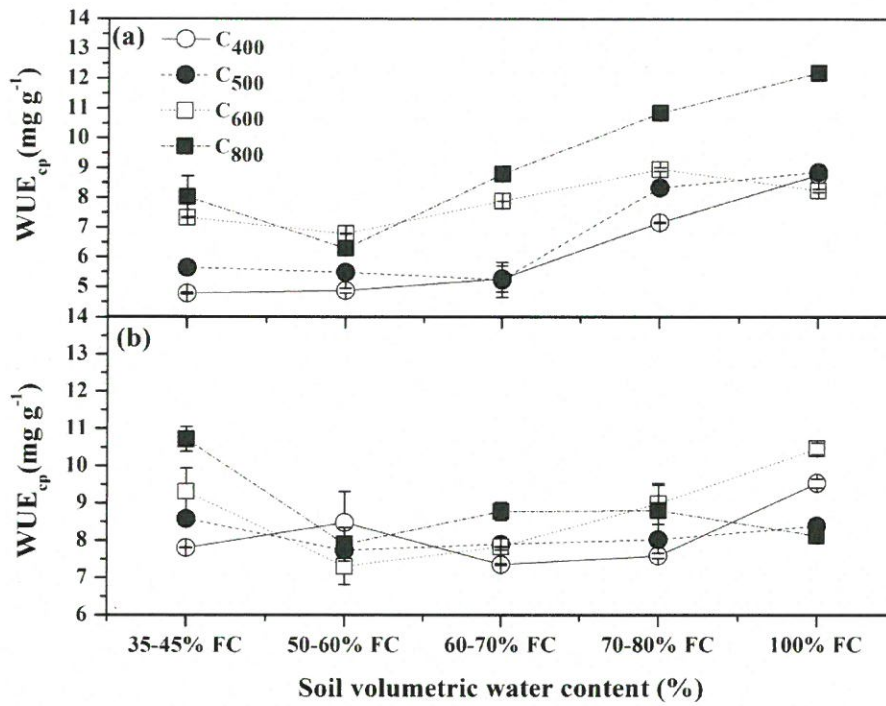
633 **Figure 2.** Net photosynthetic rates (P_n , $\mu\text{mol m}^{-2} \text{s}^{-1}$, a and b), stomatal conductance (g_s , $\text{mol H}_2\text{O m}^{-2} \text{s}^{-1}$,
 634 c and d), intercellular CO_2 concentration (C_i , $\mu\text{mol CO}_2 \text{mol}^{-1}$, e and f), and transpiration rates (T_r ,
 635 $\text{mmol H}_2\text{O m}^{-2} \text{s}^{-1}$, g and h) in *P. orientalis* and *Q. variabilis* for four CO_2 concentration \times five soil
 636 volumetric water content treatments. Means \pm SDs, $n=32$.



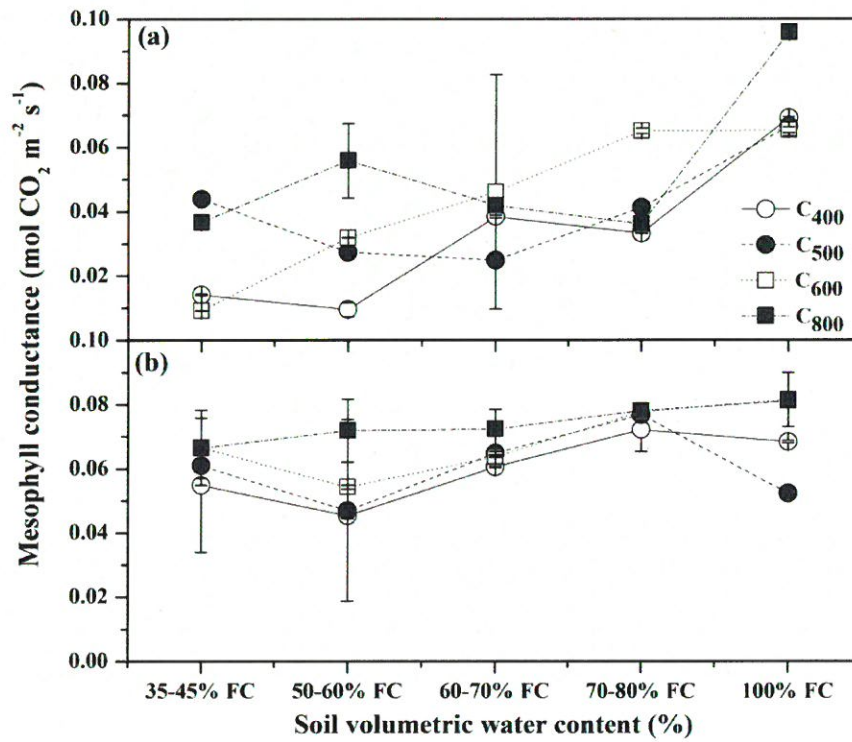
637 **Figure 3.** Carbon isotope composition of water-soluble compounds ($\delta^{13}C_{wsc}$) extracted from leaves of *P.*
 638 *orientalis* (a) and *Q. variabilis* (b) for four CO₂ concentration × five soil volumetric water content
 639 treatments. Means ± SDs, n= 32.



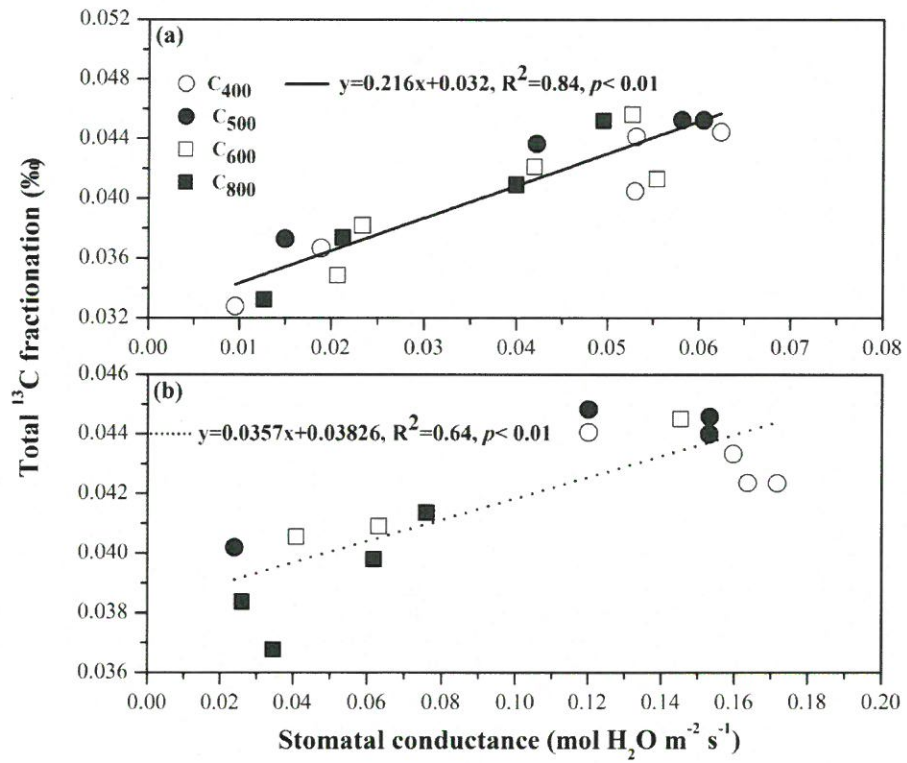
640 **Figure 4.** Instantaneous water use efficiency through gas exchange measurements (WUE_{ge}) for leaves
 641 from *P. orientalis* (a) and *Q. variabilis* (b) for four CO₂ concentration × five soil volumetric water content
 642 treatments. Means ± SDs, n= 32.



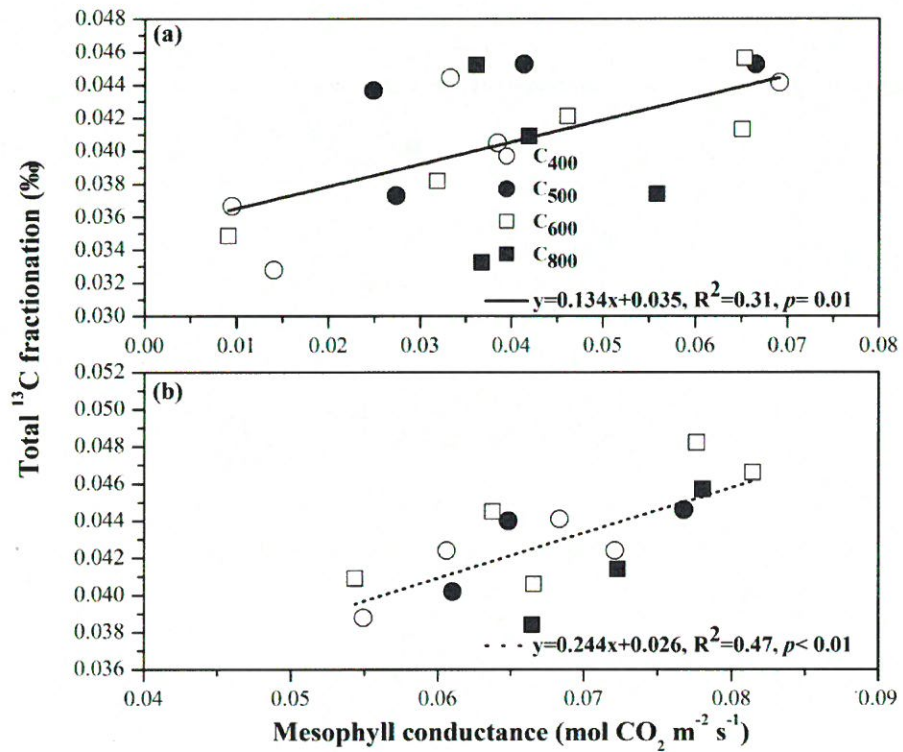
643 **Figure 5.** Instantaneous water use efficiency estimated by $\delta^{13}\text{C}$ of water-soluble compounds (WUE_{cp})
 644 from leaves of *P. orientalis* (a) and *Q. variabilis* (b) for four CO₂ concentration \times five soil volumetric
 645 water content treatments. Means \pm SDs, n= 32.



646 **Figure 6.** Mesophyll conductance in *P. orientalis* (a) and *Q. variabilis* (b) for four CO₂ concentration ×
 647 five soil volumetric water content treatments. Means ± SDs, n= 32.



648 **Figure 7.** Regressions between stomatal conductance and total ^{13}C fractionation in *P. orientalis* (a) and
 649 *Q. variabilis* (b) for four CO_2 concentration \times five soil volumetric water content treatments ($p < 0.01$, $n =$
 650 32).



651 **Figure 8.** Regressions between mesophyll conductance and total ^{13}C fractionation in *P. orientalis* (a)
 652 and *Q. variabilis* (b) for four CO_2 concentration \times five soil volumetric water content treatments ($p \leq$
 653 0.01, $n=32$).

Table 2. Carbon-13 isotope fractionation in *P. orientalis* and *Q. variabilis* under four CO₂ concentration × five soil volumetric water content treatments.

Species	SWC (of FC)	CO ₂ concentration (ppm)														
		¹³ C fractionation (%)					¹³ C fractionation (%)									
		400	500	600	800	fractionation	400	500	600	800	fractionation	400	500	600	800	
<i>P. orientalis</i>	35–45%	0.0328	0.0373	0.0349	0.0332		0.0081	0.0030	0.0034	0.0072		0.0247	0.0343	0.0315	0.0260	
	50–60%	0.0367	0.0437	0.0382	0.0374		0.0018	0.0058	0.0094	0.0004		0.0349	0.0379	0.0288	0.0370	
	60–70%	0.0405	0.0366	0.0421	0.0409		0.0018	0.0050	0.0026	0.0007		0.0387	0.0316	0.0395	0.0402	
	70–80%	0.0444	0.0453	0.0413	0.0452		0.0044	0.0052	0.0103	0.0013		0.0400	0.0401	0.0310	0.0439	
	100%	0.0441	0.0453	0.0456	0.0472	Mesophyll	0.0057	0.0040	0.0025	0.0039	Post-	0.0384	0.0413	0.0431	0.0433	
		Total ¹³ C fractionation (%)					conductance					photosynthesis				
<i>Q. variabilis</i>	35–45%	0.0388	0.0402	0.0406	0.0384		0.0007	0.0025	0.0006	0.0091		0.0381	0.0377	0.0400	0.0293	
	50–60%	0.0433	0.0448	0.0409	0.0368		0.0061	0.0084	0.0023	0.0018		0.0372	0.0364	0.0386	0.0350	
	60–70%	0.0424	0.0440	0.0445	0.0414		0.0066	0.0086	0.0078	0.0041		0.0358	0.0354	0.0367	0.0373	
	70–80%	0.0424	0.0446	0.0482	0.0457		0.0034	0.0016	0.0074	0.0028		0.0390	0.0430	0.0408	0.0429	
	100%	0.0441	0.0466	0.0466	0.0398		0.0027	0.0076	0.0022	0.0125		0.0414	0.0390	0.0444	0.0273	

Table

655 **Table 1.** Orthogonal treatments applied to *P. orientalis* and *Q. variabilis*.

<i>P. orientalis</i>	Repeats (cultivated period)	B ₁	B ₂	B ₃	B ₄	B ₅
A ₁	R ₁ :June 2–9	A ₁ B ₁ R ₁	A ₁ B ₂ R ₁	A ₁ B ₃ R ₁	A ₁ B ₄ R ₁	A ₁ B ₅ R ₁
	R ₂ :June 12–19	A ₁ B ₁ R ₂	A ₁ B ₂ R ₂	A ₁ B ₃ R ₂	A ₁ B ₄ R ₂	A ₁ B ₅ R ₂
A ₂	R ₁ :July 11–18	A ₂ B ₁ R ₁	A ₂ B ₂ R ₁	A ₂ B ₃ R ₁	A ₂ B ₄ R ₁	A ₂ B ₅ R ₁
	R ₂ :July 22–29	A ₂ B ₁ R ₂	A ₂ B ₂ R ₂	A ₂ B ₃ R ₂	A ₂ B ₄ R ₂	A ₂ B ₅ R ₂
A ₃	R ₁ :June 2–9	A ₃ B ₁ R ₁	A ₃ B ₂ R ₁	A ₃ B ₃ R ₁	A ₃ B ₄ R ₁	A ₃ B ₅ R ₁
	R ₂ :June 12–19	A ₃ B ₁ R ₂	A ₃ B ₂ R ₂	A ₃ B ₃ R ₂	A ₃ B ₄ R ₂	A ₃ B ₅ R ₂
A ₄	R ₁ :July 11–18	A ₄ B ₁ R ₁	A ₄ B ₂ R ₁	A ₄ B ₃ R ₁	A ₄ B ₄ R ₁	A ₄ B ₅ R ₁
	R ₂ :July 22–29	A ₄ B ₁ R ₂	A ₄ B ₂ R ₂	A ₄ B ₃ R ₂	A ₄ B ₄ R ₂	A ₄ B ₅ R ₂
<i>Q. variabilis</i>	Repeats (cultivated period)	B ₁	B ₂	B ₃	B ₄	B ₅
A ₁	P ₁ :June 21–28	A ₁ B ₁ P ₁	A ₁ B ₂ P ₁	A ₁ B ₃ P ₁	A ₁ B ₄ P ₁	A ₁ B ₅ P ₁
	P ₂ :July 2–9	A ₁ B ₁ P ₂	A ₁ B ₂ P ₂	A ₁ B ₃ P ₂	A ₁ B ₄ P ₂	A ₁ B ₅ P ₂
A ₂	P ₁ :August 4–11	A ₂ B ₁ P ₁	A ₂ B ₂ P ₁	A ₂ B ₃ P ₁	A ₂ B ₄ P ₁	A ₂ B ₅ P ₁
	P ₂ :August 15–22	A ₂ B ₁ P ₂	A ₂ B ₂ P ₂	A ₂ B ₃ P ₂	A ₂ B ₄ P ₂	A ₂ B ₅ P ₂
A ₃	P ₁ :June 21–28	A ₃ B ₁ P ₁	A ₃ B ₂ P ₁	A ₃ B ₃ P ₁	A ₃ B ₄ P ₁	A ₃ B ₅ P ₁
	P ₂ :July 2–9	A ₃ B ₁ P ₂	A ₃ B ₂ P ₂	A ₃ B ₃ P ₂	A ₃ B ₄ P ₂	A ₃ B ₅ P ₂
A ₄	P ₁ :August 4–11	A ₄ B ₁ P ₁	A ₄ B ₂ P ₁	A ₄ B ₃ P ₁	A ₄ B ₄ P ₁	A ₄ B ₅ P ₁
	P ₂ :August 15–22	A ₄ B ₁ P ₂	A ₄ B ₂ P ₂	A ₄ B ₃ P ₂	A ₄ B ₄ P ₂	A ₄ B ₅ P ₂