

1 **Evaluating the response of $\delta^{13}\text{C}$ in *Haloxylon ammodendron*, a**
2 **dominant C₄ species in Asian desert ecosystem, to water and nitrogen**
3 **addition as well as the availability of its $\delta^{13}\text{C}$ as the indicator of water**
4 **use-efficiency**

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23 **Abstract**

24 Variations in precipitation and atmospheric N deposition affect water and N
25 availability in desert, and thus may have significant effects on desert ecosystems.
26 *Haloxylon ammodendron* is a dominant plant in Asian desert, and addressing its
27 physiological acclimatization to the changes in precipitation and N deposition can
28 provide an insight into how desert plants adapt extreme environment by physiological
29 adjustment. Carbon isotope ratio ($\delta^{13}\text{C}$) in plants has been suggested as a sensitive
30 long-term indicator of physiological acclimatization. Therefore, this study evaluated
31 the effect of precipitation change and increasing atmospheric N deposition on $\delta^{13}\text{C}$ of
32 *H. ammodendron*. Furthermore, *H. ammodendron* is a C_4 plant, whether its $\delta^{13}\text{C}$ can
33 indicate water use-efficiency (WUE) has not been addressed. In the present study, we
34 designed a field experiment with a completely randomized factorial combination of N
35 and water, and measured $\delta^{13}\text{C}$ and gas exchange of *H. ammodendron*, then we
36 calculated the degree of bundle-sheath leakiness (ϕ) and WUE of the assimilating
37 branches of *H. ammodendron*. $\delta^{13}\text{C}$ and ϕ remained stable under N and water supply,
38 while N addition, water addition and their interaction affected gas exchange and WUE
39 in *H. ammodendron*. In addition, $\delta^{13}\text{C}$ had no correlation with WUE. These results
40 were associated with the irrelevance between $\delta^{13}\text{C}$ and the ratio of intercellular to
41 ambient CO_2 concentration (c_i/c_a), which might be caused by a special value (0.37) of
42 the degree of bundle-sheath leakiness (ϕ) or a lower activity of carbonic anhydrase
43 (CA) of *H. ammodendron*. In conclusion, $\delta^{13}\text{C}$ of *H. ammodendron* is not sensitive to
44 global change in precipitation and atmospheric N deposition, and cannot be used for

45 indicating its WUE.

46

47 **1 Introduction**

48 Recently, global precipitation pattern has changed significantly (Frank et al., 2015;
49 Knapp et al., 2015), and atmospheric N deposition has continued to rise (Galloway et
50 al., 2004; Liu et al., 2013; Song et al., 2017). Previous researchers have suggested that
51 arid ecosystems are most sensitive to climate change (Reynolds et al., 2007; Huang et
52 al., 2016), while global change in precipitation and atmospheric N deposition has an
53 important impact on water and N availability in desert (Huang et al., 2018). Thus,
54 these changes may have significant effects on desert ecosystems. *Haloxylon*
55 *ammodendron* is a dominant species in desert regions, especially in Asia. Studying the
56 physiological responses of *H. ammodendron* to global change can provide an insight
57 into how desert plants adapt extreme environment by physiological adjustment.
58 Carbon isotope ratio ($\delta^{13}\text{C}$) in plants depends on the ratio of intercellular to ambient
59 CO_2 concentration (c_i/c_a), which reflects the balance between inward CO_2 diffusion
60 rate, regulated by stomatal conductance (g_s), and CO_2 assimilating rate (A) (Farquhar
61 and Richards, 1984), and has been suggested as a sensitive long-term indicator of
62 physiological acclimatization (Battipaglia et al., 2013; Cernusak et al., 2013; Tranan
63 and Schubertt, 2016; Wang and Feng, 2012). Therefore, investigating the variations in
64 $\delta^{13}\text{C}$ of *H. ammodendron* under water and nitrogen addition can enhance
65 understanding of physiological responses of desert plants to future changes in
66 precipitation and atmospheric N deposition.

67 A large quantity of works has been devoted to the relationships between C₃ plant
68 $\delta^{13}\text{C}$ and water availability or precipitation (e.g., Diefendorf et al., 2010; Kohn, 2010;
69 Liu et al., 2005; Ma et al., 2012; Serret et al., 2018; Stewart et al., 1995; Wang et al.,
70 2005, 2008) and nitrogen availability (e.g. Cernusak et al., 2007; Li et al., 2016;
71 Sparks and Ehleringer, 1997; Yao et al., 2011; Zhang et al., 2015). However, a
72 relatively small amount of research has focused on the responses of C₄ plant $\delta^{13}\text{C}$ to
73 water availability or precipitation (Ellsworth et al., 2017; Liu et al., 2005; Rao et al.,
74 2017; Wang et al., 2006) and nitrogen availability (Ma et al., 2016; Schmidt et al.,
75 1993). For C₄ plants, $\delta^{13}\text{C}$ is controlled by both the c_i/c_a ratio and the degree of
76 bundle-sheath leakiness (ϕ), the proportion of CO₂ produced within bundle sheath
77 cells from C₄ acids that leaks back to mesophyll cells (Ellsworth and Cousins, 2016;
78 Ellsworth et al., 2017; Farquhar, 1983). Thus, the responses of C₄ plant $\delta^{13}\text{C}$ to water
79 and N availability are also affected by ϕ . Genetic factors control ϕ values, which
80 causes the interspecific differences in $\delta^{13}\text{C}$, even the responses of plant $\delta^{13}\text{C}$ to water
81 and N availability (Gresset et al., 2014). On the other hand, enzymatic activity of
82 carbonic anhydrase (CA) may influence $\delta^{13}\text{C}$ in C₄ plants (Cousins et al., 2006). CA is
83 an enzyme that catalyzes the hydration of CO₂ in mesophyll cells to form bicarbonate
84 (HCO₃⁻). Previous studies showed that CA activity in most C₄ plants is usually low,
85 just sufficient to support photosynthesis (Cousins et al., 2006; Gillon and Yakir, 2000,
86 2001; Hatch and Burnell, 1990). *H. ammodendron* is a typical C₄ plant. How its $\delta^{13}\text{C}$
87 responds to water and N availability has never been addressed.

88 Foliar $\delta^{13}\text{C}$ in C₃ plants has been considered as a useful indicator of intrinsic water

89 use-efficiency (WUE) (Farquhar, 1983). However, although some studies have
90 suggested that $\delta^{13}\text{C}$ of C_4 plants could also indicate its WUE (Henderson et al., 1992;
91 Wang et al., 2005; Cernusak et al., 2013; Ellsworth and Cousins, 2016), this statement
92 is still controversial. The relationship between $\delta^{13}\text{C}$ and WUE is based on the links
93 between c_i/c_a ratio and $\delta^{13}\text{C}$ and between c_i/c_a ratio and WUE (Ehleringer and Cerling,
94 1995). For C_3 plants, $\delta^{13}\text{C}$ always decreases with an increase in c_i/c_a ratio; but for C_4
95 plants, the correlation between $\delta^{13}\text{C}$ and c_i/c_a ratio depends on ϕ value (Cernusak et al.,
96 2013) and CA activity (Cousins et al., 2006). As mentioned above, ϕ value is under
97 genetic control, and the CA activity changes across species (Cousins et al., 2006;
98 Gillon and Yakir, 2000, 2001; Hatch and Burnell, 1990), thus, the correlation between
99 $\delta^{13}\text{C}$ and c_i/c_a ratio, as well as the relationship between WUE and $\delta^{13}\text{C}$, shows
100 interspecific difference. Whether $\delta^{13}\text{C}$ of *H. ammodendron* indicates WUE has never
101 been evaluated.

102 In this study, we designed an experiment with multiple water and nitrogen supply in
103 the southern Gurbantunggut Desert in Xinjiang Uygur Autonomous Region, China.
104 We measured the $\delta^{13}\text{C}$, gas exchange and WUE of the assimilating branches of *H.*
105 *ammodendron*. We had two objectives. One objective was to evaluate the response of
106 the dominant plant of Asian desert to future changes in precipitation and atmospheric
107 N deposition by revealing the effects of water and N supply on $\delta^{13}\text{C}$ of *H.*
108 *ammodendron*. The other was to explore the availability of $\delta^{13}\text{C}$ as the indicator of
109 water use-efficiency in *H. ammodendron*.

110

111 **2 Materials and methods**

112 **2.1 Definitions and Basic Equations**

113 Stable carbon isotopic ratio ($\delta^{13}\text{C}$) of natural materials is expressed as:

$$114 \quad \delta^{13}\text{C}(\text{‰}) = \left[\frac{(^{13}\text{C}/^{12}\text{C})_{\text{sample}}}{(^{13}\text{C}/^{12}\text{C})_{\text{standard}}} - 1 \right] \times 1000 \quad (1)$$

115 where $(^{13}\text{C}/^{12}\text{C})_{\text{sample}}$ and $(^{13}\text{C}/^{12}\text{C})_{\text{standard}}$ are the $^{13}\text{C}/^{12}\text{C}$ ratio of the sample and of Pee
116 Dee Belemnite standard (PDB), respectively. Farquhar (1983) proposed the pattern of
117 carbon isotopic discrimination (Δ) in C_4 plant:

$$118 \quad \Delta = \frac{\delta^{13}\text{C}_{\text{air}} - \delta^{13}\text{C}_{\text{plant}}}{1 + \delta^{13}\text{C}_{\text{plant}}/1000} \approx \delta^{13}\text{C}_{\text{air}} - \delta^{13}\text{C}_{\text{plant}} = a + [b_4 + \varphi (b - s) - a] \frac{c_i}{c_a} \quad (2)$$

119 where $\delta^{13}\text{C}_{\text{plant}}$ and $\delta^{13}\text{C}_{\text{air}}$ are the $\delta^{13}\text{C}$ values of plants and CO_2 in the ambient air,
120 respectively. The parameter a ($= 4.4\text{‰}$, Craig, 1954) is the carbon isotopic
121 fractionation in the diffusion of CO_2 into internal leaves; b_4 ($= -5.9\text{‰}$, O'Leary, 1984)
122 is the combined carbon isotopic fractionations occurring in the processes of gaseous
123 CO_2 dissolution, hydration/dehydration reactions of CO_2 and HCO_3^- in mesophyll
124 cells, and HCO_3^- carboxylation by PEP (phosphoenolpyruvate) carboxylase; s ($= 1.8\text{‰}$,
125 O'Leary, 1984) is the carbon isotopic fractionation during diffusion of CO_2 out of the
126 bundle-sheath cells, and b ($= 27\text{‰}$, Farquhar and Richards, 1984) is the carbon
127 isotopic fractionation of CO_2 carboxylation by RuBP (ribulose-1,5-bisphosphate)
128 carboxylase. The variable φ is the proportion of CO_2 producing within bundle sheath
129 cells from C_4 acids that leaks back to mesophyll cells, and c_i/c_a is the ratio of
130 intercellular to ambient CO_2 concentration. Eq. (2) can be transformed into the
131 following format:

$$132 \quad \delta^{13}\text{C}_{\text{plant}} = -[b_4 + \varphi(b - s) - a] \frac{c_i}{c_a} + \delta^{13}\text{C}_{\text{air}} - a \quad (3)$$

133 According to Eq. (3), if the coefficient $[b_4 + \varphi (b - s) - a]$ is greater than 0, $\delta^{13}\text{C}$
134 decreases with increasing c_i/c_a ; if this coefficient is lower than 0, $\delta^{13}\text{C}$ increases with
135 increasing c_i/c_a .

136 Water use-efficiency (WUE) is defined as the amount of assimilated carbon dioxide
137 by plants under the consumption of per unit water. There are two characteristics of
138 WUE, instantaneous WUE (ins-WUE) and intrinsic WUE (int-WUE), respectively.
139 Ins-WUE can be calculated by:

$$140 \quad \text{ins-WUE} = A/E = (c_a - c_i)/1.6v = c_a(1 - c_i/c_a)/1.6v \quad (4)$$

141 where A is photosynthetic rate, E is transpiration rate and v is calculated
142 by:

$$143 \quad v = (e_i - e_a)/p \quad (5)$$

144 where e_i and e_a are the water vapor pressure inside and outside the leaves, p is the
145 atmospheric pressure.

146 The definition of int-WUE is:

$$147 \quad \text{int-WUE} = A/g_s = (c_a - c_i)/1.6 = c_a(1 - c_i/c_a)/1.6 \quad (6)$$

148 where g_s is stomatal conductance.

149 **2.2 Study site and species**

150 This experiment was conducted at the Fukang Station of Desert Ecology, Chinese
151 Academy of Sciences, on the southern edge of the Gurbantunggut Desert (44°26' N,
152 87°54' E) in northwestern China. The altitude of the study site is 436.8 m above
153 average sea level (a.s.l.). It is a typical continental arid, temperate climate, with a hot
154 summer and cold winter in the area. The mean annual temperature is 7.1 °C and the

155 mean annual precipitation is 215.6 mm, with a potential evaporation of about 2000
156 mm. The mean annual temperature and the annual rainfall amount in the sampling
157 year are 10.23 °C and 122.7 mm (Cui, 2018). The soil type is grey desert soils
158 (Chinese classification) with aeolian sands on the surface (0-100 cm). The
159 percentages of clay (< 0.005 mm), silt (0.005-0.063 mm), fine sand (0.063-0.25 mm)
160 and medium sand (0.25-0.5 mm) range from 1.63-1.76%, 13.79-14.15%,
161 55.91-56.21% and 20.65-23.23%, respectively (Chen et al., 2007). The soil is highly
162 alkaline (pH = 9.55 ± 0.14) with low fertility. The vegetation is dominated by
163 *Haloxylon ammodendron* and *Haloxylon persicum* with about 30% coverage. Herbs
164 include ephemerals, annuals and small perennials, with a cover of ca. 40% (Fan et al.,
165 2013). Although the coverage of the two *Haloxylon* species is a little lower than that
166 of herbs, the biomass of the former is much larger than that of the latter, because
167 *Haloxylon* plants are shrubs with an average height of 1.5 m whereas the latter are
168 very low herbaceous plants. Biological soil crusts are distributed widely on the soil
169 between the herbs and *Haloxylon*, with almost 40% coverage (Zhang et al., 2007).

170 The present study focused on *Haloxylon ammodendron* because it is the dominant
171 species in Asian desert. *H. ammodendron* is a species of Chenopodiaceae, which is a
172 xerophytic and halophytic woody plant (Cui et al., 2017). The leaves of *H.*
173 *ammodendron* have been completely degraded due to the extreme drought, and the
174 assimilation branches, which are the glossy green branches (Fig. S1), perform the
175 same functions as the leaves. Due to its drought tolerance, *H. ammodendron* is widely
176 distributed in desert areas.

177 **2.3 Experimental design**

178 A field experiment with a completely randomized factorial combination of water and
179 nitrogen has been conducted from 2014 to 2017. We designed two water addition
180 levels (0, 60 mm·yr⁻¹; W0, W1) based on the prediction that precipitation will
181 increase by 30% in northern China in the next 30 years (Liu et al., 2010), and three
182 levels of N addition (0, 30, 60 kg N·ha⁻¹·yr⁻¹; N0, N1 and N2), because N deposition
183 has reached 35.4 kg N·ha⁻¹·yr⁻¹ in the nearby city, Urumqi (Cui et al., 2017) and will
184 double by 2050 relative to the early 1990s (Galloway et al., 2008). Therefore, there
185 were six treatments (W0N0, W0N1, W0N2, W1N0, W1N1, W1N2) in this
186 experiment. Four replicates of each treatment were set, making a total of 24 plots with
187 a size of 10 m × 10 m. A small sub-plot with a size of 1.5 m × 1.5 m was set in each
188 plot. A well-grown *H. ammodendron* was enclosed in the center of the sub-plot. The
189 average height and coverage of an individual *H. ammodendron* were 1.5 m and 1.9 m²,
190 respectively, and did not vary significantly across the plots. The type of nitrogen used
191 in the present study is NH₄NO₃. To simulate natural water and N inputs, the
192 treatments were applied in equal amounts, twelve times, once a week in April, July
193 and September, as 5 mm·m⁻² of water and 2.5 or 5 kg N·ha⁻¹ each week (Cui et al.,
194 2017). Usually, water addition was with a sprinkler kettle, irrigating over the canopy
195 of *H. ammodendron*.

196 **2.4 Measurements of gas exchange and WUE**

197 As mentioned above, the main assimilating organ of *H. ammodendron* is the
198 assimilation branches. Thus, we conducted gas exchange measurements on the

199 assimilation branches of the *H. ammodendron* grown in the sub-plots by LI-6400
200 portable photosynthesis system. The measurements were conducted on 27-29, June
201 2016, which is the main growing season of *H. ammodendron*. It may be the most
202 appropriate to take measurements during this period, and the results of the
203 measurements are therefore more representative. Previous studies have also usually
204 conducted this measurement during the growing season (Nyongesah and Wang, 2013;
205 Cui, 2018; Gong et al., 2019). The ins-WUE was calculated based on these measured
206 gas exchange traits by the Eq. (4), and int-WUE by Eq. (6). At each plot, the top
207 assimilating branches of a mature individual was selected randomly for the
208 measurement of gas exchange, which includes photosynthetic rate (A), stomatal
209 conductance (g_s), transpiration rate (E), the ambient CO_2 concentration (c_a) and the
210 intercellular CO_2 concentration (c_i). Before the measurement of gas exchange, it takes
211 about 5s to stabilize after the assimilating branches was inserted in the cuvette. We
212 repeated 10 times on the same assimilating branches for each measurement. Our
213 measurements were carried out under the conditions of a standard $450 \text{ mmol}\cdot\text{mol}^{-1}$
214 CO_2 concentration at a flow rate of $500 \text{ mmol}\cdot\text{s}^{-1}$ above saturation in photo flux
215 density of $1600 \text{ mmol}\cdot\text{m}^{-2}\cdot\text{s}^{-1}$. The temperature of the measuring assimilating
216 branches varied from $29.5 \text{ }^\circ\text{C}$ to $30.5 \text{ }^\circ\text{C}$ during the entire period of gas exchange
217 measurements.

218 **2.5 Samples collection**

219 Sample collection was conducted in 20, July, during the addition of water and
220 nitrogen. Considering that there is a considerable difference in $\delta^{13}\text{C}$ between buds,

221 young and matured leaves, so we collected the mature assimilating branches of *H.*
222 *ammodendron* for the $\delta^{13}\text{C}$ measurements. All *H. ammodendron* individuals grown in
223 plots (10 m \times 10 m) were sampled. Eight pieces of the mature assimilating branches
224 (15-20 cm long) were collected from each individual; two pieces of assimilating
225 branches were collected at each of the four cardinal directions from the positions of
226 full irradiance. All assimilating branches from the same plot were combined into one
227 sample. After the samples were collected, they were immediately divided into two
228 parts randomly and taken back to the laboratory at Fukang Station. The first part was
229 used to determine the chlorophyll content. The second part was immediately
230 inactivated in a 105 °C oven in the laboratory at Fukang Station, and then brought
231 back to Beijing in a ziplock bag. The time interval between sample collection and
232 inactivation is very short. After inactivation, the carbon exchange of the assimilating
233 branches stop, so the isotope composition of the samples will not change anymore.
234 All plant samples of the second part were air-dried immediately in the laboratory in
235 Beijing. Then the samples were ground into a fine powder using a steel ball mixer
236 mill MM200 (Retsch GmbH, Haan, Germany) for the measurements of $\delta^{13}\text{C}$ and N
237 contents.

238 **2.6 Measurements of plant $\delta^{13}\text{C}$, plant N and chlorophyll contents**

239 The $\delta^{13}\text{C}$ and N measurements were performed on a Delta^{Plus} XP mass spectrometer
240 (Thermo Scientific, Bremen, Germany) coupled with an automated elemental
241 analyzer (Flash EA1112, CE Instruments, Wigan, UK) in a continuous flow mode, at
242 the Stable Isotope Laboratory of the College of Resources and Environmental

243 Sciences, China Agricultural University. The carbon isotopic ratios were reported in
 244 the delta notation relative to the V-PDB standard. For this measurement, we obtained
 245 standard deviations low than 0.15‰ for $\delta^{13}\text{C}$ among replicate measurements of the
 246 same sample. And standard deviations for the N measurements were 0.1%.

247 The chlorophyll contents of all samples were determined immediately when the
 248 sample were taken back in the laboratory at Fukang Station. The samples were first
 249 extracted by 95% ethyl alcohol (0.5 g sample to 25 mL ethyl alcohol), and then the
 250 absorbancy was measured under the wave length of 665 and 649 nm by the
 251 spectrophotometer. The content of chlorophyll a, b was calculated by the follow
 252 equations:

$$253 \quad \text{Chlorophyll a (mg/L)} = 13.95 \times \text{OD}_{665} - 6.88 \times \text{OD}_{649} \quad (7)$$

$$254 \quad \text{Chlorophyll b (mg/L)} = 24.96 \times \text{OD}_{649} - 7.32 \times \text{OD}_{665} \quad (8)$$

255 where OD₆₆₅ and OD₆₄₉ are the absorbancy under the wave length of 665 and
 256 649nm, respectively.

257 **2.7 Calculation of the degree of bundle-sheath leakiness**

258 The degree of bundle-sheath leakiness (ϕ) was calculated by the transformation of Eq.
 259 (2):

$$260 \quad \phi = \left(\frac{(\delta^{13}\text{C}_{\text{air}} - \delta^{13}\text{C}_{\text{plant}}) / (1 + \delta^{13}\text{C}_{\text{plant}}/1000) - a}{c_i/c_a} + a - b_4 \right) / (b - s) \quad (9)$$

261 In this equation, parameters a, b₄, b and s are constant, while $\delta^{13}\text{C}_{\text{plant}}$ and c_i/c_a are
 262 the measured values of our samples. We did not measure the $\delta^{13}\text{C}_{\text{air}}$ at our study site,
 263 so we had to use an approximation of the $\delta^{13}\text{C}_{\text{air}}$ to do this ϕ calculation. The
 264 approximated value we used is -9.77‰, which has been measured at Donglingshan

265 Mountain, Beijing, north China in September 2019. The two sites should have similar
266 $\delta^{13}\text{C}_{\text{air}}$ because the two sites are located in countryside with less human activities and
267 have a similar distance from the nearest city. The straight line distances between
268 Donglingshan Mountain and the city center of Beijing, and between our study site and
269 Urumqi city are about 90 km. In addition, since the $\delta^{13}\text{C}_{\text{air}}$ has large diurnal and
270 seasonal variations, we used the published range of $\delta^{13}\text{C}_{\text{air}}$ from May to July in
271 Shangdianzi, China (Data comes from Global Monitoring Laboratory, Earth System
272 Research Laboratories, <https://www.esrl.noaa.gov/gmd>) to calculate the minimum and
273 maximum $\delta^{13}\text{C}_{\text{air}}$ of the time period, which is the main growing season for *H.*
274 *ammოდendron*. Finally, the $\delta^{13}\text{C}_{\text{air}}$ used in the calculation ranged from -10.52‰ to
275 -9.01‰ with an average of -9.77‰.

276 **2.8 Statistical analysis**

277 Statistical analyses were conducted using SPSS software (SPSS for Windows, Version
278 20.0, Chicago, IL, United States). One-way analysis of variance (ANOVA) and
279 two-way analysis of variance (ANOVA) were used to compare the difference of $\delta^{13}\text{C}$
280 and other physiological traits between each treatment. Pearson analysis was used to
281 determine the correlation among $\delta^{13}\text{C}$, WUE and c_i/c_a in *H. ammოდendron*.

282

283 **3 Results**

284 **3.1 Plant $\delta^{13}\text{C}$ under water and nitrogen addition**

285 The $\delta^{13}\text{C}$ of the assimilating branches of *H. ammოდendron* in the six treatments
286 W0N0, W0N1, W0N2, W1N0, W1N1, W1N2 was -14.18 ± 0.19 ‰, -14.71 ± 0.35 ‰,

287 $-14.45 \pm 0.18 \text{ ‰}$, $-14.67 \pm 0.40 \text{ ‰}$, $-14.65 \pm 0.38 \text{ ‰}$, $-14.34 \pm 0.29 \text{ ‰}$, respectively.

288 One-way ANOVA analyses showed no significant variation in $\delta^{13}\text{C}$ across treatments
289 ($p = 0.79$, Fig. 1). Two-way ANOVA analyses suggested that $\delta^{13}\text{C}$ was not affected by
290 water addition ($p = 0.68$), N addition ($p = 0.61$) and their interaction ($p = 0.56$, Table
291 1).

292 Fig.1

293 Table 1

294 **3.2 Gas exchange and WUE under water and nitrogen addition**

295 Photosynthetic rate (A), stomatal conductance (g_s), transpiration rate (E) and c_i/c_a
296 ranged from $12.11 \mu\text{mol CO}_2 \text{ m}^{-2} \text{ s}^{-1}$ to $39.35 \mu\text{mol CO}_2 \text{ m}^{-2} \text{ s}^{-1}$, from $0.09 \text{ mol H}_2\text{O}$
297 $\text{m}^{-2} \text{ s}^{-1}$ to $0.31 \text{ mol H}_2\text{O m}^{-2} \text{ s}^{-1}$, from $2.87 \text{ mmol H}_2\text{O m}^{-2} \text{ s}^{-1}$ to $8.49 \text{ mmol H}_2\text{O m}^{-2} \text{ s}^{-1}$
298 and 0.11 to 0.57, respectively. One-way ANOVA analyses showed significant changes
299 in leaf gas exchange across the six treatments ($p < 0.01$ for A, g_s , E and c_i/c_a , Fig. 2).
300 Two-way ANOVA analyses suggested that water addition had exerted effect on c_i/c_a (p
301 < 0.01), that N additions influenced A ($p < 0.01$) and c_i/c_a ($p = 0.009$), and that the
302 interaction between water and N supply played a role in g_s ($p < 0.001$), E ($p < 0.001$)
303 and c_i/c_a ($p < 0.001$, Table 1).

304 Fig. 2

305 Instantaneous WUE (ins-WUE) and intrinsic WUE (int-WUE) ranged from 3.09
306 $\mu\text{mol CO}_2 / \text{mmol H}_2\text{O}$ to $8.49 \mu\text{mol CO}_2 / \text{mmol H}_2\text{O}$ and from $93.64 \mu\text{mol CO}_2 / \text{mol}$
307 H_2O to $208.47 \mu\text{mol CO}_2 / \text{mmol H}_2\text{O}$, respectively. One-way ANOVA analyses
308 showed significant changes in these two indexes across these treatments (both $p <$

309 0.001, Fig. 3). Two-way ANOVA analyses suggested that water addition, N addition
310 and their interaction all have significant effect on these two indexes (all $p < 0.05$,
311 Table 1).

312 Fig. 3

313 **3.3 Correlations among $\delta^{13}\text{C}$, WUE and c_i/c_a ratio**

314 In order to test whether $\delta^{13}\text{C}$ in *H. ammodendron* can indicate WUE, the relationships
315 among $\delta^{13}\text{C}$, ins-WUE, int-WUE and c_i/c_a ratio were revealed in this study. Our
316 results showed no correlation between $\delta^{13}\text{C}$ and ins-WUE ($p = 0.23$, Fig. 4a), between
317 $\delta^{13}\text{C}$ and int-WUE ($p = 0.23$, Fig. 4c), and between $\delta^{13}\text{C}$ and c_i/c_a ratio ($p = 0.18$, Fig.
318 4e). However, there was a negative correlation between ins-WUE and c_i/c_a ratio ($p <$
319 0.001 , Fig. 4b), and between int-WUE and c_i/c_a ratio ($p < 0.001$, Fig. 4d).

320 Fig. 4

321 **3.4 The degree of bundle-sheath leakiness under water and nitrogen addition**

322 The ϕ value calculated from the minimum $\delta^{13}\text{C}_{\text{air}}$ ranged from 0.16 to 0.50 with a
323 mean value of 0.35; the ϕ value calculated from the maximum $\delta^{13}\text{C}_{\text{air}}$ ranged from
324 0.44 to 0.70 with a mean value of 0.55, and the ϕ value calculated from the average
325 $\delta^{13}\text{C}_{\text{air}}$ ranged from 0.32 to 0.59 with a mean value of 0.45. One-way ANOVA
326 analyses showed no significant variation in ϕ calculated from the minimum, average
327 and maximum $\delta^{13}\text{C}_{\text{air}}$ across treatments ($p = 0.60$ for the ϕ calculated from the
328 minimum $\delta^{13}\text{C}_{\text{air}}$, $p = 0.77$ for the ϕ calculated from the average $\delta^{13}\text{C}_{\text{air}}$, and $p = 0.90$
329 for the ϕ calculated from the maximum $\delta^{13}\text{C}_{\text{air}}$, Fig. 5). Two-way ANOVA analyses
330 suggested that ϕ was not affected by water addition ($p = 0.46$ for the ϕ calculated

331 from the minimum $\delta^{13}\text{C}_{\text{air}}$, $p = 0.64$ for the ϕ calculated from the average $\delta^{13}\text{C}_{\text{air}}$, and
332 $p = 0.98$ for the ϕ calculated from the maximum $\delta^{13}\text{C}_{\text{air}}$, N addition ($p = 0.65$ for the
333 ϕ calculated from the minimum $\delta^{13}\text{C}_{\text{air}}$, $p = 0.60$ for the ϕ calculated from the average
334 $\delta^{13}\text{C}_{\text{air}}$, and $p = 0.55$ for the ϕ calculated from the maximum $\delta^{13}\text{C}_{\text{air}}$) and their
335 interaction ($p = 0.30$ for the ϕ calculated from the minimum $\delta^{13}\text{C}_{\text{air}}$, $p = 0.52$ for the ϕ
336 calculated from the average $\delta^{13}\text{C}_{\text{air}}$, and $p = 0.87$ for the ϕ calculated from the
337 maximum $\delta^{13}\text{C}_{\text{air}}$, Table 1).

338 Fig. 5

339

340 **4 Discussion**

341 The $\delta^{13}\text{C}$ of the assimilating branches in *H. ammodendron* did not change across
342 treatments (Fig. 1, Table 1), suggesting that neither water addition nor nitrogen
343 addition influenced the $\delta^{13}\text{C}$ of *H. ammodendron*. Previous studies also reported no
344 significant relationship between $\delta^{13}\text{C}$ of C_4 plant and water availability (Swap et al.,
345 2004; Wang et al., 2008), and between $\delta^{13}\text{C}$ of C_4 plant and nitrogen availability (Yao
346 et al., 2011, Yang et al., 2017).

347 In general, the effects of water availability and nitrogen availability on $\delta^{13}\text{C}$ are
348 dependent on c_i/c_a ratio, which reflects the balance between stomatal conductance (g_s)
349 and photosynthetic rate (A) (Farquhar and Richards, 1984). Stomatal conductance (g_s)
350 usually increases with increasing water availability under water addition. Although
351 two-way ANOVA analyses suggested that water addition had no effect on both A and
352 g_s (Table 1); one-way ANOVA analyses showed that g_s was higher in W1N0 than that

353 in W0N0 (Fig. 2b), indicating that water addition had a positive effect on g_s under
354 ambient N condition. Increasing g_s under water supply will lead to the rise of
355 intercellular CO_2 because of the decrease of diffusional resistance to CO_2 . As the
356 results, c_i/c_a ratio was observed to increase with increasing moisture (Fig. 2d, Table 1).
357 However, $\delta^{13}C$ remained stable under water addition (Fig. 1, Table 1). Thus, c_i/c_a ratio
358 could not explain the observed response of $\delta^{13}C$ to water supply.

359 For most plants in natural ecosystems, nitrogen is the key factor limiting plant
360 growth (Hall et al., 2011). Thus, nitrogen addition usually causes plants to absorb
361 more N. However, extreme drought could prevent plants from absorbing N even
362 under high N supply. In the present experiment, N supply was found to have an effect
363 on N contents in *H. ammodendron*. Relative to the control treatment (W0N0), N
364 contents increased with N supply under low N addition, but kept unchanged under
365 high addition (Table S1, S2). Nitrogen is the main constituent of Rubisco (ribulose -
366 1,5 - biphosphate carboxylase oxygenase) and chlorophyll in plants. Thus,
367 chlorophyll a was found to have the similar pattern as N contents under water and N
368 supply. Chlorophyll a was higher in W0N1 than W0N0, and there was no difference in
369 chlorophyll a between W0N0 and W0N2 (Table S1). Increasing chlorophyll contents
370 in W0N1 should lead to the increase in photosynthetic rate (A). However, different
371 from our prediction, one-way ANOVA analyses suggested that A in W0N1 did not
372 differ from that in W0N0, and that A in W0N2 was lower than that in W0N0 (Fig., 2a).
373 Two-way ANOVA analyses showed that N addition had an influence on A (Table 1).
374 Both the analyses suggested that N supply played a negative role in A. These results

375 might be associated with the extremely high light intensity at the study site. Due to the
376 high light intensity, photosynthetic rate might not be correlated with chlorophyll
377 contents (Gabrielsen, 1948). The negative effect of N supply on A led to the decrease
378 in the consumption of intercellular CO₂. Consequently, c_i/c_a ratio increased with N
379 supply (Fig. 2d, Table 1). Therefore, the variations in c_i/c_a ratio with N addition could
380 not account for the unchanged pattern in $\delta^{13}\text{C}$ under N supply (Fig. 1).

381 The co-application of water and nitrogen had a negative effect on A but no effect on
382 g_s (W0N0 vs. W1N1, W1N2, Fig. 2a, b). The responses of A and g_s to the
383 co-application of water and nitrogen resulted in an increase in c_i/c_a ratio (Fig., 2d).
384 Since $\delta^{13}\text{C}$ remained unchanged under the co-application of water and nitrogen (Fig.
385 1), c_i/c_a ratio could not also explain the observed $\delta^{13}\text{C}$ response to the co-application
386 of water and nitrogen.

387 In summary, the unchanged $\delta^{13}\text{C}$ across treatments was not dependent on the c_i/c_a
388 ratio in *H. ammodendron* (Fig. 4e). The observed $\delta^{13}\text{C}$ stability across treatments
389 might be associated with the ϕ value and carbonic anhydrase (CA) in *H.*
390 *ammodendron*. For C₄ plants, the relationship between carbon isotope discrimination
391 ($\Delta \approx \delta^{13}\text{C}_{\text{air}} - \delta^{13}\text{C}_{\text{plant}}$, see Eq. (2)) and c_i/c_a ratio is controlled by ϕ values (Ellsworth
392 and Cousins, 2016; Ellsworth et al., 2017; Farquhar, 1983; Wang et al., 2008). Some
393 studies suggested that ϕ value was stable for a given species under a wide range of
394 environmental conditions (Henderson et al., 1992; Wang et al., 2008; Cernusak et al.,
395 2013). However, other studies had different conclusions that ϕ value was influenced
396 by irradiation (Bellasio and Griffiths, 2014; Kromdijk et al., 2010; Pengelly et al.,

397 2010; Ubierna et al., 2013), temperature (von Caemmerer et al., 2014), water stress
398 (Fravolini et al., 2002; Gong et al., 2017; Williams et al., 2001; Yang et al., 2017) and
399 nitrogen supply (Fravolini et al., 2002; Meinzer and Zhu, 1998; Yang et al., 2017). In
400 current study, the ϕ value of *H. ammodendron* remained unchanged across six
401 treatments (Fig. 5), and two-way ANOVA analyses suggested that water supply and N
402 supply had no effect on ϕ (Table 1). Therefore, the ϕ value of *H. ammodendron* was
403 insensitive to water and N addition in this study. Even if the ϕ value remains stable,
404 the relationship between Δ and c_i/c_a ratio is also associated with the magnitude of the
405 ϕ value. Cernusak et al. (2013) predicted that when ϕ value is greater than 0.37, the
406 correlation between Δ and c_i/c_a ratio is positive; conversely, when ϕ value is less than
407 0.37, the correlation is negative. In particular, when ϕ value is equal to 0.37, there will
408 be no correlation between them, because the coefficient ($[b_4 + \phi (b - s) - a]$ in Eq. (2))
409 of c_i/c_a ratio equals to 0 (Cernusak et al., 2013). The ϕ value calculated from the
410 average $\delta^{13}C_{air}$ ranged from 0.32 to 0.59 with a mean value of 0.45 in present study.
411 Thus, the correlation between Δ and c_i/c_a in *H. ammodendron* should be positive
412 based on the prediction by Cernusak et al. (2013). Δ always changes in the opposite
413 direction to $\delta^{13}C_{plant}$ changes according to Eq. (2), thus, a negative relationship
414 between $\delta^{13}C_{plant}$ and c_i/c_a is expected. In fact, this study observed no correlation
415 between $\delta^{13}C$ and c_i/c_a in *H. ammodendron* (Fig. 4e); this indicates that ϕ was not the
416 driver of the observed $\delta^{13}C$ pattern in *H. ammodendron*. However, the measured $\delta^{13}C$
417 represents the fixed carbon isotope composition throughout the assimilation branch
418 formation period, which usually spans at least several weeks. And the measured c_i/c_a

419 is an instant indicator. As a result, there were some uncertainties in the calculation of
420 ϕ value using Eq. (2) based on the measured $\delta^{13}\text{C}$ and c_i/c_a . In addition, the mean ϕ
421 value calculated from the minimum and maximum $\delta^{13}\text{C}_{\text{air}}$ were 0.35 and 0.55,
422 respectively, suggesting that the ϕ value of *H. ammodendron* might be close to 0.37,
423 which led to the observed insensitive response of $\delta^{13}\text{C}$ to water and N addition.

424 The enzymatic activity of CA may be another mechanism behind the unchanged
425 $\delta^{13}\text{C}$ across treatments. Cousins et al. (2006) suggested that enzymatic activity of CA
426 affects carbon isotope discrimination in most C_4 plants because CA can result in the
427 parameter b_4 changes (see Eq. (2)). But in traditional view, the parameter b_4 was a
428 constant. However, it is only true when the ratio of PEP carboxylation rate to the CO_2
429 hydration rate (V_p/V_h) is equal to zero, which is caused by a high CA activity. If
430 V_p/V_h is not zero, b_4 will change and be controlled by V_p/V_h (Cousins et al., 2006).
431 Previous studies reported that CA activity was low in most C_4 plants (Cousins et al.,
432 2006; Gillon and Yakir, 2000, 2001; Hatch and Burnell, 1990). Thus, CA activity in *H.*
433 *ammodendron* might also be low, leading to the change in b_4 with V_p/V_h , and thus
434 $\delta^{13}\text{C}$. Cousins et al. (2006) added V_p/V_h into the discrimination pattern of C_4 plants
435 and predicted that at a given ϕ value, when the V_p/V_h is 0 or 1, the correlation
436 between Δ and c_i/c_a ratio is negative or positive, respectively. Since CA activity is low
437 in most C_4 plants, and the V_p/V_h always ranges from 0 to 1, we speculate that no
438 correlation between Δ and c_i/c_a ratio may also occur when the V_p/V_h is a certain value
439 between 0 and 1. The irrelevance of Δ and c_i/c_a ratio also means that $\delta^{13}\text{C}_{\text{plant}}$ is not
440 related to c_i/c_a ratio due to the negative correlation between Δ and $\delta^{13}\text{C}_{\text{plant}}$ according

441 to Eq. (2). Thus, the uncorrelated pattern between $\delta^{13}\text{C}$ and c_i/c_a ratio in *H.*
442 *ammodendron* might be related to this specific V_p/V_h value due to low CA activity.

443 In addition, the unchanged $\delta^{13}\text{C}$ across treatments may also be controlled by the
444 water sources of *H. ammodendron*. Previous study has found that the root of *H.*
445 *ammodendron* can be inserted into the soil layer deeper than 3 m (Sheng et al., 2004),
446 which made it easy to uptake groundwater. Therefore, *H. ammodendron* may be less
447 sensitive to water addition. However, a study conducted in the same region has found
448 that the shallow soil water (0-40 cm) and groundwater are two important water
449 sources for *H. ammodendron* (Dai et al., 2014), and another study has reported that
450 water addition resulted in an increase of soil water contents in shallow soil layer (Cui,
451 2018). Moreover, gas exchange changed across treatments in the present study (Fig.
452 2). Thus, the utilization of groundwater by *H. ammodendron* may be one of the
453 reasons why its $\delta^{13}\text{C}$ was not sensitive to water and N addition, but it should not be
454 the main reason.

455 Whether foliar $\delta^{13}\text{C}$ of C_4 plants can indicate their WUE is still controversial.
456 Henderson et al. (1992) found that $\delta^{13}\text{C}$ of 10 C_4 species has negative correlation with
457 their WUE. Although this result was just opposite to a positive relationship between
458 $\delta^{13}\text{C}$ and WUE for C_3 plants (Farquhar, 1983; Duquesnay et al., 1998; Feng, 1998), it
459 is a proof that $\delta^{13}\text{C}$ of C_4 plants can indicate their WUE. In the work of Henderson et
460 al. (1992), they found that the ϕ values in 10 C_4 species was around 0.21 over a range
461 of irradiance and leaf temperature. According to the suggestion by Cernusak et al.
462 (2013) that Δ is negatively related to c_i/c_a ratio when ϕ value is less than 0.37, thus,

463 the $\delta^{13}\text{C}$ of 10 C_4 species has a positive correlation with c_i/c_a ratio. In general, under
464 fixed ambient CO_2 concentration, WUE is always negatively correlated with c_i/c_a ratio
465 (see Eq. (4) and Eq. (6)). This is why a negative relationship between $\delta^{13}\text{C}$ and WUE
466 was observed for the 10 C_4 species. The present study showed that ins-WUE and
467 int-WUE both had no correlation with $\delta^{13}\text{C}$ in *H. ammodendron* (Fig. 4a, 4c), which
468 was different from the results published by Henderson et al. (1992). In general, c_i/c_a
469 ratio is the link between WUE and $\delta^{13}\text{C}$. As mentioned above, if the ϕ value equals to
470 0.37 and/or the activity of CA is very low, $\delta^{13}\text{C}$ would not correlate to c_i/c_a ratio, and
471 thus leads to the uncorrelation between $\delta^{13}\text{C}$ and WUE. In addition, the different time
472 scales of $\delta^{13}\text{C}$, ins-WUE and int-WUE may also result in this uncorrelation. As
473 mentioned above, the measured $\delta^{13}\text{C}$ represents the long-term fixed carbon isotope
474 composition (at least several weeks). And the value of ins-WUE and int-WUE were
475 calculated from the gas exchange of a short-term measurement, which lasted only a
476 few minutes. Therefore, this difference in time scale may also drive the uncorrelation
477 between $\delta^{13}\text{C}$ and WUE. Although the defects in measurements could introduce some
478 uncertainty in the observed relationship between $\delta^{13}\text{C}$ and WUE, $\delta^{13}\text{C}$ remained stable
479 under water and nitrogen addition (Fig. 1, Table 1), while the measured ins-WUE and
480 int-WUE was higher in the control treatment (W0N0) than other treatments (Fig. 3),
481 suggesting water and N supply had a significant effect on WUE (Table 1). These
482 results indirectly confirmed that $\delta^{13}\text{C}$ of *H. ammodendron* could not indicate its WUE.

483 The present study has found that $\delta^{13}\text{C}$ of *H. ammodendron* could not be used as a
484 indicator of its WUE. Although this conclusion cannot be analogous to all C_4 plants,

485 the present study has important implications for the understanding of physiological
486 responses of desert plants to future changes in precipitation and atmospheric N
487 deposition. *H. ammodendron* is a dominant species in Asia desert, which has a great
488 effect on the stabilization of sand dunes, the survival and development of understory
489 plants and the structure and function of desert ecosystems (Sheng et al., 2005; Su et
490 al., 2007; Cui et al., 2017). Thus, *H. ammodendron* is widely distributed in desert
491 areas, and the prediction of its drought adaptation is crucial in desert ecosystem.

492

493 **5 Conclusion**

494 Global changes including precipitation and atmospheric N deposition have been
495 proved to have an important influence on ecosystems, especially for the arid
496 ecosystems. The present study showed that water and N addition had little effect on
497 the $\delta^{13}\text{C}$ values and the degree of bundle-sheath leakiness (ϕ) of *H. ammodendron*, but
498 played an important role in the change of its gas exchange and water use efficiency
499 (WUE). In addition, different pattern of instantaneous WUE (ins-WUE), intrinsic
500 WUE (int-WUE) and $\delta^{13}\text{C}$ across treatment and no correlation between instantaneous
501 WUE (ins-WUE) and $\delta^{13}\text{C}$, and between intrinsic WUE (int-WUE) and $\delta^{13}\text{C}$ has been
502 found in this study, suggesting that $\delta^{13}\text{C}$ of *H. ammodendron* could not indicate its
503 WUE. This result was caused by the lack of the correlation between $\delta^{13}\text{C}$ and the ratio
504 of intercellular to ambient CO_2 concentration (c_i/c_a), which might be associated with
505 the degree of bundle-sheath leakiness (ϕ) or the low activity of carbonic anhydrase
506 (CA). Thus, the current experiment implies that the availability of $\delta^{13}\text{C}$ as the

507 indicator of WUE could be not universal for C₄ species.

508

509 **Conflict of interest**

510 The authors declare that they have no conflict of interest.

511

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516

517 **Authors' Contributions**

518 G Wang and J Li designed the experiment and modified the manuscripts. Z Chen
519 designed and executed the experiment and wrote the manuscripts. X Liu designed the
520 experiment. X Cui executed the experiment. Y Han executed the experiment.

521

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530

531 **Data availability**

532 The datasets analyzed in this manuscript are not publicly available. Requests to access
533 the datasets should be directed to gawang@cau.edu.cn.

534

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742 **Table 1 The p values of all measured and calculated indexes in plants under two-way ANOVA**
 743 **analysis of water (W) and nitrogen (N) additions**

	W	N	W*N
$\delta^{13}\text{C}$	0.678	0.607	0.563
Photosynthetic rate (A)	0.331	0.008**	0.183
Stomatal conductance (g_s)	0.533	0.871	<0.001***
Transpiration rate (E)	0.622	0.883	<0.001***
c_i/c_a	0.004**	0.009**	<0.001***
ins-WUE	0.002**	<0.001***	<0.001***
int-WUE	0.004**	0.018*	<0.001***
φ_{\min}	0.463	0.645	0.295
φ_{ave}	0.644	0.600	0.521
φ_{\max}	0.982	0.549	0.866

744 Note. φ_{\min} , φ_{ave} and φ_{\max} represent the φ values calculated from the minimum, average and maximum $\delta^{13}\text{C}_{\text{air}}$. *, **, ***

745 *** indicates a significant influence.

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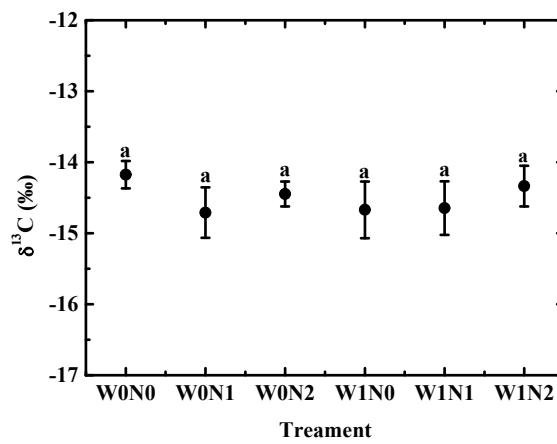
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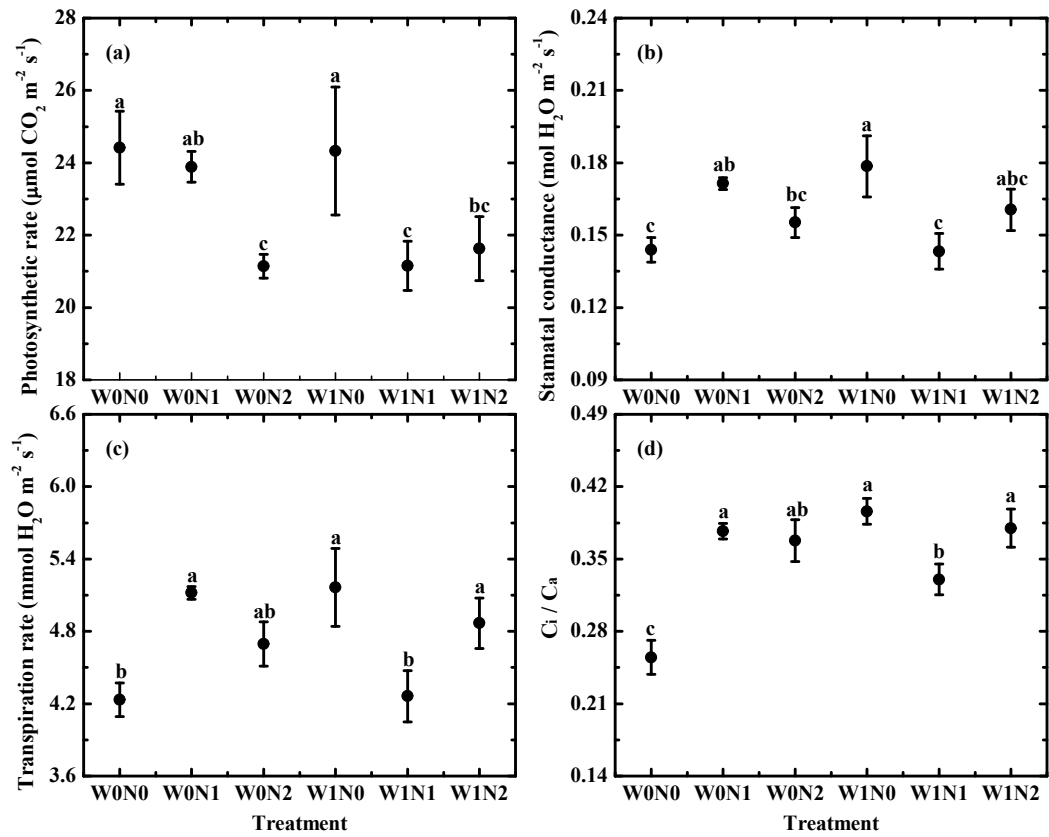
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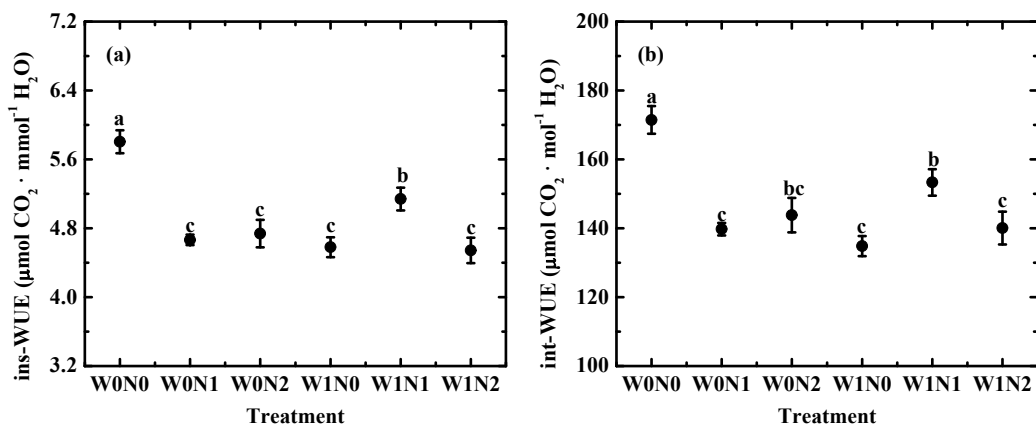
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762 **Fig. 1** The $\delta^{13}\text{C}$ of assimilating branches of *Haloxylon ammodendron* under water (W) and

763 **nitrogen (N) additions.** The spot represents the mean value of four replicates with error bars
 764 denoting the standard error (SE).
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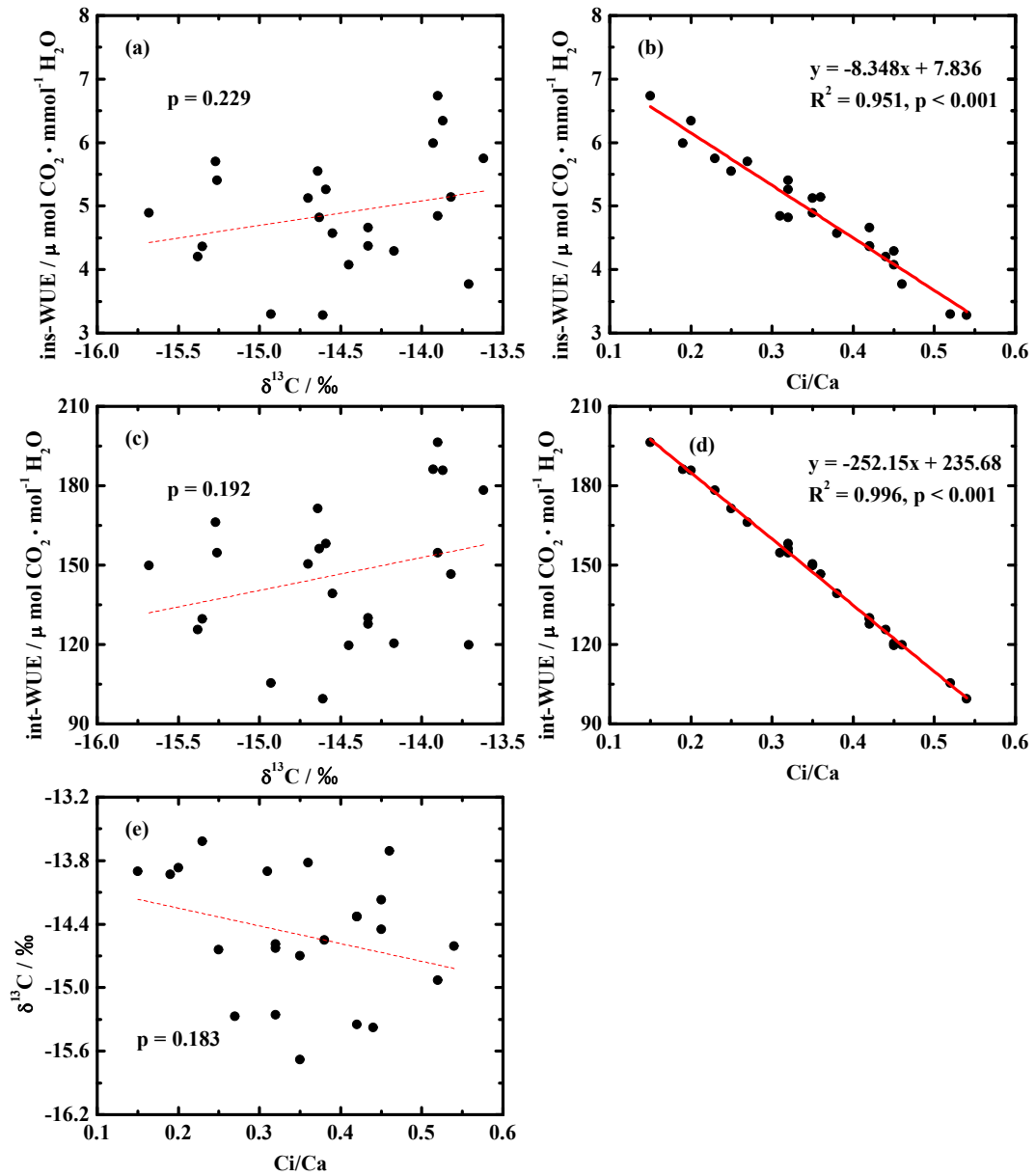
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 768 **Fig. 2 Variations in photosynthetic rate (a), stomatal conductance (b), water use-efficiency (c)**
 769 **and c_i/c_a (d) across water (W) and nitrogen (N) additions.** The spot represents the mean value
 770 of four replicates with error bars denoting the standard error (SE).
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 774 **Fig. 3 Variations in ins-WUE (a) and int-WUE (b) across water (W) and nitrogen (N)**
 775 **additions.** The spot represents the mean value of four replicates with error bars denoting the
 776 standard error (SE).

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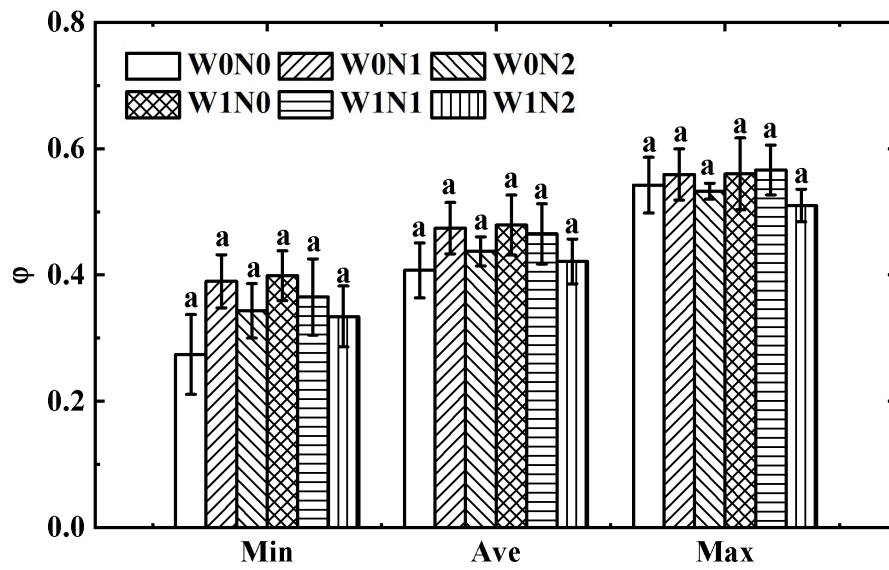
780 **Fig. 4** Correlations of ins-WUE vs. $\delta^{13}\text{C}$ (a), ins-WUE vs. c_i/c_a (b), int-WUE vs. $\delta^{13}\text{C}$ (c),

781 int-WUE vs. c_i/c_a (d), and $\delta^{13}\text{C}$ vs. c_i/c_a (e) of assimilating branches of *Haloxylon*

782 *ammodendron*

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786 **Fig. 5** Variations in ϕ calculated by the from the minimum (Min), average (Ave) and
 787 maximum (Max) values of $\delta^{13}\text{C}_{\text{air}}$ across water (W) and nitrogen (N) additions. The box
 788 represents the mean value of four replicates with error bars denoting the standard error (SE).

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