- 1 Exogenous DCPTA ameliorates the soil drought effect on nitrogen metabolism in
- 2 maize during the pre-female inflorescence emergence stage
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## 29 Highlights:

This is the first article that explores the effects of DCPTA on nitrogen metabolism and the first article that explores the effects of DCPTA on crops under soil drought

- 32 conditions.
- 33 Abstract

34• 2-(3,4-Dichlorophenoxy) triethylamine (DCPTA) regulates many aspects of plant 35 development; however, its effects on soil drought tolerance are unknown. We pre-treated maize (Zea mays L.) by foliar application of DCPTA and subsequently 36 37 exposed the plants to soil drought and rewatering conditions during the pre-female 38 inflorescence emergence stage. Exogenous DCPTA significantly alleviated drought-induced decreases in maize yield, shoot and root relative growth rate (RGR), 39 40 leaf relative water content (RLWC), net photosynthetic rate (Pn), stomatal conductance (Gs) and transpiration rate (Tr), nitrate (NO<sub>3</sub><sup>-</sup>), nitrite (NO<sub>2</sub><sup>-</sup>), and soluble 41 protein contents, and nitrate reductase (NR), nitrite reductase (NiR), isocitrate 42 43 dehydrogenase (ICDH), alanine aminotransferase (AlaAT) and aspartate 44 aminotransferase (AspAT) activities; increases in the intercellular CO<sub>2</sub> concentration 45 (Ci), the ammonium  $(NH_4^+)$  and free amino acid contents, and the glutamate 46 dehydrogenase (GDH) and protease activities. Simultaneously, exogenous DCPTA 47 improved the spatial and temporal distribution of roots and increased the root hydraulic conductivity (Lp), flow rate of root-bleeding sap and  $NO_3^-$  delivery rates. 48 49 Moreover, Exogenous DCPTA protected the chloroplast structure from drought injury. Taken together, our results suggest that exogenous DCPTA mitigates the repressive 50 effects of drought on N metabolism and subsequently enhances drought tolerance 51 52 during the pre-female inflorescence emergence stage of maize.

53 **Key words:** DCPTA; maize (*Zea mays* L.); nitrogen metabolism; pre-female 54 inflorescence emergence stage; soil drought; yield

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# 57 Introduction

Crops are frequently exposed to drought during the growth period because of 58 limited and erratic rainfall patterns due to global climate change, which leads to 59 60 restrictions on agricultural productivity worldwide (Trenberth et al., 2014). Maize (Zea mays L.), an essential component of global food security, is widely cultivated 61 around the word. The majority of the cultivated area of maize is almost wholly 62 rain-fed and experiences sporadic drought and rewetting cycles (Nuccio et al., 2015). 63 However, maize is considered to be a drought-sensitive crop and loses approximately 64 1/4 potential yield annually due to drought (Ziyomo and Bernardo, 2013). By 2050, 65 the world population will reach 9 billion people, resulting in a high demand for maize 66 (projected to double); furthermore, at that time, drought will severely restrict crop 67 68 growth for more than 50% of the cultivated land (Schiermeier, 2014).

To stabilize and increase global crop production to satisfy the demand of the 69 globally burgeoning population, it is imperative to design agronomic research to 70 improve maize performance under drought stress (Zhao et al., 2018). The application 71 of plant growth regulators has been considered an effective way to enhance crop 72 drought resistance (Ali et al., 2017). Multiple investigations have indicated that a 73 tertiary amine bioregulator known as 2-(3,4-dichlorophenoxy) triethylamine (DCPTA) 74 75 regulates many aspects of plant development; for example, DCPTA promotes plant growth (Keithly et al., 1990a), enlarges chloroplast volume (Keithly et al., 1990b), 76 enhances photosynthetic enzyme activity (Wan and Mendoza, 1992), accelerates  $CO_2$ 77 fixation (Gausman et al., 1985), and stimulates carotenoid biosynthesis (Benedict et 78 al., 1985). As far as we know, very few studies of DCPTA have focused on crops, and 79 the effect of DCPTA on crops exposed to soil drought are still unclear. 80

Nitrogen (N) metabolism is a fundamental process in determining the growth and productivity of plants (Kusano *et al.*, 2011). After being taken up by root systems, nitrate (NO<sub>3</sub><sup>-</sup>) is converted to nitrite (NO<sub>2</sub><sup>-</sup>) by nitrate reductase (NR), the first step of N uptake and utilization. Subsequently, nitrite (NO<sub>2</sub><sup>-</sup>) is converted to NH<sub>4</sub><sup>+</sup> by nitrite reductase (NiR) with reduction-ferredoxin (Fd<sub>red</sub>) as an electron donor (Rajasekhar *et* 

al., 2010). Afterward, the ammonium  $(NH_4^+)$ , derived from NO<sub>3</sub><sup>-</sup> reduction, 86 photorespiration and/or other metabolic processes is assimilated into glutamine by the 87 glutamine synthase/glutamine oxoglutarate aminotransferase (GS/GOGAT) cycle or 88 89 the alternative glutamate dehydrogenase (GDH) pathway with 2-oxoglutarate (2-OG) and reducing equivalents provided by photosynthesis (Chardon et al., 2012). 90 Subsequently, glutamate serving as a donor of the amino group is used for the 91 synthesis of other amino acids, which are used for the synthesis of various organic 92 molecules such as chlorophyll, proteins and nucleic acids. The reactions are catalysed 93 by aminotransferases such as alanine aminotransferase (AlaAT) and aspartate 94 aminotransferase (AspAT) (Slattery et al., 2017). 95

Drought disrupts N metabolism mainly via inhibiting the uptake and/or long-distance transportation of  $NO_3^-$  (Nacer *et al.*, 2013), altering the activities of enzymes involved in N metabolism (Robredo *et al.*, 2011), inhibiting amino acid synthesis, and promoting protein hydrolysis (Fresneau *et al.*, 2007). At present, the study of plant growth regulators mainly concentrates on the improvement of photosynthesis and antioxidant systems, and there have been only a limited number of publications related to N metabolism.

Our previous hydroponic trial found that exogenous DCPTA drastically promoted growth under non-stress conditions and mitigated the PEG-simulated drought-induced growth inhibition of maize at the seedling stage (Xie *et al.*, 2017). The present study was conducted to explore whether DCPTA can alleviate soil drought injuries to maize and whether the effects are associated with the modulation of nitrogen metabolism.

#### 109 Materials and Methods

110 Plant material, growth conditions, design and sampling

Seeds of the maize cultivar ZhengDan 958 and DCPTA were obtained from the
Henan Academy of Agricultural Sciences in China and the China Zhengzhou
Zhengshi Chemical Limited Company, respectively.

114 These experiments were performed in 2016 and 2017 at the Experimental Station

of Northeast Agricultural University, Harbin (126°73'E, 45°73'N), Heilongjiang 115 province, China. The research field area has a temperate continental monsoon climate. 116 The rainfall and mean temperature data during the study period (2016 and 2017, 117 118 May–October) are listed in Fig. 1. Pits (inner length, 10 m; width, 7 m; and height, 1.2 m) in the field were used as experiment containers (Fig. 2). Plastic sheets were used to 119 cover the inner sides of the pits, and a rain-proof shed was used to ensure the crops 120 were solely dependent on soil moisture and irrigation over the course of the 121 experiment to maintain the soil water conditions. The soil used was Chernozem and 122 was sieved (pore size, 1 cm) and diluted with vermiculite (particle diameter, 4-8 mm; 123 soil to vermiculite, v/v, 2:1). Before planting, soil chemical analysis was conducted 124 125 according to Cottenie et al. (1982), and the results are presented in Table 1. 126 Fertilization was carried out by adding ammonium nitrate (33.5% N), calcium superphosphate (15.5%  $P_2O_5$ ), and potassium sulfate (48%  $K_2O$ ) at the rates of 8.0, 127 8.0 and 20 kg pit<sup>-1</sup>, respectively, before planting. No fertilizer was applied after 128 129 planting. All containers were watered to 85% before planting. The seeds were manually sown on 2nd May 2016 and 4th May 2017 and were harvested on 7th 130 October 2016 and 3rd October 2017, respectively. Three seeds were sowed per hole to 131 ensure germination, and only the healthiest seedling within 20 days was kept at each 132 133 site. Each container consisted of 10 rows, and the plant-to-plant and row-to-row distances were 20 cm and 65 cm, respectively. In addition, the ground around the 134 containers was manually sown with the same plant-to-plant and row-to-row distances. 135 The control of plant diseases and insect pests was conducted by managers. 136

The maize at the nine-leaf stage (during the pre-female inflorescence emergencestage) were treated as follows:

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(1) plants were irrigated continuously and sprayed with either 10 mL water (well-watered) or DCPTA (well-watered+DCPTA) per plant;

(2) irrigation was stopped to form the drought conditions and sprayed with 10
mL of either water (drought) or DCPTA (drought +DCPTA) per plant; plants were
rehydrated after 20 days of drought treatment.

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The concentrations of DCPTA (25 mg/L) were based on the results of previous screening experiments, and Tween-20 (0.03%) was added as a surfactant to the solution for spraying. Each treatment had five replicates, and experiments were performed in a completely randomized design. The dynamic changes in soil water contents during the experimental stage are exhibited in Fig. 3.

Random plants from each treatment were sampled on days 0, 5, 10, 15, 20, 25 149 and 30. For leaf sampling, the middle part of the  $9^{th}$  leaf (numbered basipetally) was 150 sampled for analysis of leaf gas exchange, and the same part of the leaf was stored at 151 -80°C after immersion in liquid nitrogen for 30 min for determination of physiological 152 153 parameters. For root sampling, a hand-held soil auger (inner diameter of 20 cm) was 154 used to obtain soil cores from 0 to100 cm depth of the soil profile at 10 cm increments. 155 The soil cores were soaked in a plastic container overnight, and roots were stirred and sieved through a mesh (400 holes  $cm^{-2}$ ). The soil cores were the carefully washed by 156 157 swirling water through the cores. The soil material and old dead roots debris were 158 manually separated from the live roots.

159 **Plant measurement and analysis** 

#### 160 **Relative growth rate (RGR) and plant productivity**

The shoots and roots of maize were oven dried at  $105^{\circ}$ C for 45 min and then held at 80°C for 48 h; the shoot and root dry weights plant<sup>-1</sup> were determined soon afterwards. The RGR was determined as follows: RGR (fresh weight) = [ln (final dry weight) – ln (initial dry weight)]/(duration of treatment days) (Kingsbury *et al.*, 1984). A leaf area metre (Li-COR 3100; Li-COR, Lincoln, NE, USA) was used to estimate the leaf area; number of grains plant<sup>-1</sup> (GN) and grain yield plant<sup>-1</sup> (GY) were recorded at the maize physiological maturity stage.

## 168 Leaf relative water content of (RLWC) and soil water content (SWC)

169 The RLWC was determined on fresh leaf disks  $(2 \times 2 \text{ cm})$  from the middle part of

the 8th leaves (numbered basipetally). After they were weighed (FW), the disks were

immersed in distilled water at 25 °C overnight to obtain the turgid weight (TW). The

172 leaves were dried at 80°C for 48 h and then weighed a third time (DW). RLWC was

173 calculated as follows:

174 
$$RLWC$$
 (%) = [(FW-DW)/(TW-DW)] × 100.

SWC was determined in the soil from the internal area of each container. After
being weighed (FW), the soil portion was dried at 85°C for 96 h and then weighed
(DW). SWC was calculated as follows:

178 SWC (%) = 
$$[(FW-DW)/DW] \times 100.$$

## 179 Gas exchange

The photosynthetic rate (Pn), transpiration rate (Tr), stomatal conductance (Gs), and intercellular CO<sub>2</sub> concentration (Ci) values were determined with a portable photosynthesis system (LI-6400XT; LI-COR Biosciences, Lincoln, NE, USA) between at 13:00~14:00 h. The 6-cm<sup>2</sup> leaf chamber was used, and the photo flux density was 1000 lmol m<sup>-2</sup>s<sup>-1</sup>.

# 185 Transmission electron microscopy of chloroplasts

Observations were performed according to the description of Hu *et al.* (2014), and the chloroplast ultrastructure was observed under a H-7650 transmission electron microscope (manufacture: Hitachi, Japan).

# Root morphological traits, root hydraulic conductivity, and the collection of root-bleeding sap

191 Roots from each soil core were scanned using a digital scanner (Epson V700, 192 Indonesia). The root images were analysed using the WinRHIZO Image Analysis 193 system (Version 2013e) (Regent Instruments Inc., Canada). The root length density 194 (RLD, cm root cm<sup>-3</sup> soil) and root square area density (RSD, cm<sup>2</sup> root cm<sup>-3</sup> soil) were 195 calculated according to the method described by Mosaddeghi *et al.* (2009).

The hydrostatic root hydraulic conductivity (Lp) was measured with a
Scholander pressure chamber according to the method described by López-Pérez *et al*.
(2007).

199 The plants were cut by scissors at 10-12 cm above the soil surface at 18:00-19:00. Centrifuge tubes (inner diameter 40 mm) with cotton were placed on the upper end of 200 201 the stalks, and the stalk joints and centrifuge tubes were wrapped by plastic film to keep impurities and insects out (Fig. 4). The bleeding sap was collected for 12 h; then, 202 the cotton was extracted from each centrifuge tube and placed into a glass syringe 203 (100 ml), and the root-bleeding sap was squeezed out for volume measurement. The 204  $NO_3^{-}$  content in the root-bleeding sap was determined by AA3 Continuous Flow 205 206 Analytical System (Seal, Germany) according to Guan et al. (2014) The flow rate of the root-bleeding sap and the  $NO_3^-$  delivery rate were expressed as ml  $h^{-1}$  root<sup>-1</sup> and 207  $\mu$ g h<sup>-1</sup> root<sup>-1</sup>, respectively. 208

# 209 Foliar NO<sub>3</sub>, NO<sub>2</sub> and NH<sub>4</sub><sup>+</sup> contents

The foliar NO<sub>3</sub><sup>-</sup> content determination by the reduction of NO<sub>3</sub><sup>-</sup> to NO<sub>2</sub><sup>-</sup> followed 210 the salicylic acid methods of Cataldo et al. (1975), the absorbance was monitored at 211 410 nm. The foliar NO<sub>2</sub><sup>-</sup> content was determined using the method described by Barro 212 et al. (1991). The  $NO_2^-$  content was calculated according to the standard curve 213 obtained by known concentrations of KNO<sub>3</sub>. The NH<sub>4</sub><sup>+</sup> content was determined by 214 measuring the absorbance changes at 620 nm, as described by Weber (2007). The 215 216 NH4<sup>+</sup> content was calculated according to the standard curve obtained by known 217 concentrations of  $(NH_4)_2SO_4$ .

# 218 Enzyme activities involved in nitrogen metabolism

The activities of foliar NR and NiR were measured based on the conversion of NO<sub>3</sub><sup>-</sup> to NO<sub>2</sub><sup>-</sup> and NO<sub>3</sub><sup>-</sup> to NH<sub>4</sub><sup>+</sup>, following the methods of Barro *et al.* (1991), and Ida and Morita (1973), respectively. One NR and NiR unit was expressed as the enzyme amount required for the conversion of 1 mmol of NO<sub>3</sub><sup>-</sup> to NO<sub>2</sub><sup>-</sup> per hour and the

enzyme amount required for the conversion of 1 mmol of  $NO_3^-$  to  $NH_4^+$  min<sup>-1</sup>, 223 respectively. GS activity was measured according to the methods of O'Neal and Joy 224 (1973). One unit of GS was expressed as the enzyme amount required to catalyse the 225 226 formation of 1 mmol of glutamylmonohydroxamate per min. The results were expressed as µmol NADH used per minute per milligram of Pro. The GOGAT activity 227 was measured based on the conversion of 2-ketoglutarate to glutamate, following the 228 methods of Groat and Vance (1981). One unit of GOGAT was expressed as the 229 number of enzymes catalysing the oxidation of 1 mmol of NADH per min. The 230 deaminating GDH activity (NAD-GDH) and aminating GDH activity (NADH-GDH) 231 were determined by recording the reduction of NAD and the oxidation of NADH, 232 233 respectively, as described by Groat and Vance (1981). One unit of GDH was 234 calculated in units of mmol of NADH oxidized/NAD reduced per minute. Isocitrate 235 dehydrogenase (ICDH) was assayed according to the method of Lòpez-Millàn et al. (2000). The activity was expressed as U per minute per milligram of protein. 236

## 237 Activity of AlaAT and AspAT

The AlaAT and AspAT activities were determined according to the methods of Jia *et al.* (2015). Enzyme activity was expressed as  $\mu$ mol g<sup>-1</sup>min<sup>-1</sup>.

#### 240 Contents of free amino acids and soluble proteins and protease activity

241 The free amino acid contents were assayed with the ninhydrin reagent method 242 according to Yemm and Cocking (1955), followed by absorbance readings at 570 nm 243 using glycine as the standard. The soluble protein contents were determined using the Coomassie Brilliant Blue G-250 reagent following the description of Bradford (1976), 244 245 followed by absorbance readings at 595 nm using bovine serum albumin as the 246 standard. Protease activity was determined by the casein digestion assay described by 247 Drapeau (1974). By this method, one unit is the number of enzymes required to release acid-soluble fragments equivalent to 0.001 A280 per minute at 37°C and pH 248 7.8. 249

#### 250 Statistical analysis

The data were analysed using SPSS 17.0 and all the values are presented as the mean  $\pm$  SE. The means were separated using the least significant difference (LSD) test at the 5% probability level.

254 **Results** 

255 **Yield** 

Drought stress significantly inhibited the maize yield (Fig. 5). Compared with 256 257 the well-watered treatment, in the drought treatment, the grain number decreased by 258 29.23% in 2016 and 33.24% in 2017, and the grain yield decreased by 34.06% in 259 2016 and by 38.22% in 2017. However, the decrease in maize yield was partially 260 recovered by DCPTA. Compared with the well-watered treatment, in the 261 drought+DCPTA treatment, the grain number decreased by 14.01% in 2016 and by 262 16.55% in 2017, and the grain yield decreased by 17.98% in 2016 and by 20.54% in 263 2017. Moreover, the application of DCPTA improved maize yield under well-watered 264 conditions. Compared with the well-watered treatment, in the DCPTA treatment, the 265 grain number increased by 5.97% in 2016 and by 6.50% in 2017, and the grain yield 266 increased by 7.31% in 2016 and by 8.02% in 2017.

#### **Relative growth rate (RGR) of shoot and root**

The shoot growth rate was inhibited during the drought period, the root growth 268 rate was promoted over days 0-10 and subsequently decreased, and the growth rates 269 270 of shoots and roots recovered during rehydration (Fig. 6 and 7). In the drought 271 treatment compared with the control in 2016, the RGR of shoots and roots decreased by 52.26% and by 48.59%, respectively, between day 15 and 20 and by 39.76% and 272 273 37.70%, respectively, between day 25 and 30 in 2016. In the drought treatment 274 compared with the control in 2017, the RGR of shoots and roots decreased by 65.18% 275 and 66.52%, respectively, between day 15 and 20 and by 51.97% and 43.74%, 276 respectively, between day 25 and 30 in 2017. However, the decrease in growth was

partially recovered by DCPTA. In the drought+DCPTA treatment compared with the 277 control in 2016, the RGR of shoots and roots decreased by 29.96% and 26.31%, 278 respectively, between day 15 and 20, and by 10.11% and 11.49%, respectively, 279 280 between day 25 and 30. In the drought+DCPTA treatment compared with the control in 2017, the RGR of shoots and roots decreased by 34.20% and 42.23%, respectively, 281 between day 15 and 20, and by 12.41% and 14.75%, respectively, between day 25 and 282 30. Moreover, the application of DCPTA improved maize growth under well-watered 283 conditions. The shoot RGR difference between the well-watered+DCPTA treatment 284 and well-watered treatment was significant at days 16-20 and 26-30 in 2016 and days 285 11-30 in 2017, respectively. Similarly, the root RGR difference between the DCPTA 286 287 treatment and well-watered treatment was significant at days 6-10 and 16-25 in 2016 288 and days 11-30 in 2017, respectively.

#### **Root length density (RLD) and root surface area density (RSD)**

290 Drought dramatically inhibited RLD and RSD in the 0-60 cm soil profile in 291 2016 and 2017 (Fig. 8). At day 20, there was a significant difference in RLD for the 292 drought treatment and drought+DCPTA treatment in 20-40 cm in 2016 and in 20-50 293 cm in 2017, respectively. On day 30, the RLD declines were partially reversed by 294 rehydration. On day 30, there was a significant difference in RLD for the drought 295 treatment and drought+DCPTA treatment in 0-50 cm in both 2016 and 2017. On day 30, there was a significant (P > 0.05) difference in RLD for the well-watered 296 297 treatment and well-watered+DCPTA treatment in 0-20 cm in both 2016 and 2017. On 298 day 20, there was a significant difference in RLD between the drought treatment and 299 drought+DCPTA treatment in 10-40 cm in 2016 and 10-50 cm in 2017. On day 30, 300 the declines in RLD were partially reversed by rehydration. At day 30, there was a 301 significant (P > 0.05) difference in RLD between the drought treatment and drought+DCPTA treatment in 0-40 cm in 2016 and 0-50 cm in 2017. On day 30, 302 303 there was a significant difference between RLD for the well-watered treatment and well-watered+DCPTA treatment in 0-20 cm in 2016 and in 0-30 cm in 2017. 304

# Root hydraulic conductivity, flow rate of root-bleeding sap and NO<sub>3</sub><sup>-</sup> delivery rates

The root hydraulic conductivity, flow rate of root-bleeding sap and NO<sub>3</sub><sup>-</sup> 307 concentrations in the root-bleeding sap declined continuously during the drought 308 309 period and recovered during rehydration (Fig. 9). In the drought treatment compared 310 with the control, the root hydraulic conductivity, flow rate of root-bleeding sap and 311  $NO_3^-$  delivery rates decreased by 34.21%, 75.69% and 76.58%, respectively, on day 312 20 and by 19.34%, 35.96% and 57.37%, respectively, on day 30 in 2016; these values 313 decreased by 47.01%, 78.80% and 61.34%, respectively, on day 20 and by 29.03%, 314 33.79%, and 50.47%, respectively, on day 30 in 2017. However, the DCPTA 315 application partially reversed the decline in root hydraulic conductivity, root-bleeding sap flow and  $NO_3$  delivery rates caused by drought and resulted in a faster recovery 316 317 after rehydration. In the drought+DCPTA treatment compared with the control, the 318 root hydraulic conductivity, root-bleeding sap flow and NO<sub>3</sub><sup>-</sup> delivery rates decreased 319 by 17.91%, 46.73% and 41.90%, respectively, on day 20 and by 9.87%, 18.80% and 320 23.51%, respectively, on day 30 in 2016; these values decreased by 26.60%, 47.66% 321 and 37.06%, respectively, on day 20 and by 16.94%, 14.68% and 25.23%, respectively, on day 30. 322

Under well-watered conditions, DCPTA significantly increased root hydraulic conductivity on the 15<sup>th</sup>, 20<sup>th</sup>, 25<sup>th</sup> and 30<sup>th</sup> days in 2016 and on the 20<sup>th</sup> and 25<sup>th</sup> days in 2017; DCPTA significantly increased the flow rate of root-bleeding sap on the 10<sup>th</sup>, 15<sup>th</sup>, 20<sup>th</sup> and 25<sup>th</sup> days in both 2016 and 2017; DCPTA significantly increased the NO<sub>3</sub><sup>-</sup> concentration in root-bleeding sap on the 10<sup>th</sup>, 15<sup>th</sup>, 20<sup>th</sup>, 25<sup>th</sup> and 30<sup>th</sup> days in 2016 and on the 10<sup>th</sup>, 15<sup>th</sup>, 20<sup>th</sup> and 25<sup>th</sup> days in 2017.

# 329 Leaf water status (RLWC)

The RLWC declined continuously over the drought period and recovered during rehydration (Fig. 10). In the drought treatment compared with the control, RLWC decreased by 30.88% on day 20 and by 13.08% on day 30 in 2016 and

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decreased by 37.48% on day 20 and by 21.89% on day 30 in 2017. However, the DCPTA application partially reversed the decline in RLWC caused by drought and resulted in a faster recovery of the foliar RLWC contents after rehydration. In the drought+DCPTA treatment compared with the control, the RLWC decreased by 11.75% on day 20 and by 3.84% on day 30 in 2016 and decreased by 21.15% on day 20 and by 10.73% on day 30 in 2017. Under well-watered conditions, DCPTA application had no significant effect on the RLWC.

## 340 Gas exchange parameters

341 Pn, Gs and Tr declined continuously over the drought period and recovered during rehydration (Fig. 11). In the drought treatment compared with the control, Pn, 342 343 Gs and Tr decreased by 46.20%, 68.97% and 51.35%, respectively, on day 20 and by 344 35.06%, 46.30% and 38.17%, respectively, on day 30 in 2016 and decreased by 345 56.00%, 71.10% and 62.10%, respectively, on day 20 and by 45.89%, 44.02% and 346 43.79%, respectively, on day 30 in 2017. However, the DCPTA application partially 347 reversed the decline in Pn, Gs and Tr caused by drought and resulted in a faster recovery of Pn, Gs and Tr rehydration. In the drought+DCPTA treatment compared 348 349 with the control, Pn, Gs and Tr decreased by 23.41%, 37.39% and 20.95%, 350 respectively, on day 20 and by 8.77%, 20.89% and 10.61%, respectively, on day 30 in 351 2016 and decreased by 35.31%, 32.03% and 34.11%, respectively, on day 20 and by 14.16%, 15.33% and 8.62%, respectively, on day 30. Under well-watered conditions, 352 DCPTA significantly increased Pn on the 10<sup>th</sup>, 15<sup>th</sup>, 25<sup>th</sup> and 30<sup>th</sup> days in 2016 and on 353 the 15<sup>th</sup>, 20<sup>th</sup> and 25<sup>th</sup> days in 2017. In addition, under well-watered conditions, 354 DCPTA significantly increased Gs on the 15<sup>th</sup>, 20<sup>th</sup>, 25<sup>th</sup> and 30<sup>th</sup> days in 2016 and on 355 the 10<sup>th</sup>, 15<sup>th</sup>, 20<sup>th</sup>, 25<sup>th</sup> and 30<sup>th</sup> days in 2017 and significantly increased Tr on the 10<sup>th</sup>, 356 15<sup>th</sup>, 20<sup>th</sup>, 25<sup>th</sup> and 30<sup>th</sup> days in both 2016 and 2017. 357

In 2016 and 2017, Ci showed the same variation tendency during the drought period. In the drought treatment, Ci declined over days 0-10, subsequently increased over days 10-20, and then decreased after rehydration. In the drought+DCPTA treatment, Ci declined over days 0-15 day and subsequently increased over days
15-30. Under well-watered conditions, DCPTA application had no significant effect
on Ci.

## 364 Chloroplast ultrastructure

Regardless of whether DCPTA was applied, the photosynthetic mesophyll cells 365 of the non-stressed seedlings included a delimited cell wall containing chloroplasts. 366 367 These chloroplasts had intact membranes and a regular arrangement of granal and 368 stromal thylakoids, which were attached to the cell wall and exhibited typical 369 ellipsoidal shapes (Fig. 12). However, in the stressed seedlings, the cell wall structure 370 was incomplete and exhibited indistinct gradation, a lower density, and loose edges. 371 Plasmolysis and degradation were also evident in part of the cell membrane. 372 Moreover, the chloroplasts, which separated from the plasma membrane, were nearly 373 round and swelled asymmetrically, the thylakoids were overly disorganized, and the 374 thylakoid membranes were loose and showed an increased number of plastoglobules. 375 In the PEG-6000+DCPTA treatment, the complete membrane structures of the 376 chloroplasts were present, and the shapes of the chloroplasts changed slightly from 377 elongated ellipses to ellipses close to the cell walls. A well-aligned internal lamellar system and fewer plastoglobules were observed in the leaves of the 378 379 PEG-6000+DCPTA treatment compared with the leaves of the PEG-6000 treatment.

## **ICDH activity**

ICDH activity declined continuously over the drought period and recovered during rehydration (Fig. 13). In the drought treatment compared with the control, ICDH activity decreased by 40.75% on day 20 and by 36.33% on day 30 in 2016 and decreased by 37.77% on day 20 and by 33.07% on day 30 in 2017. However, the DCPTA application partially reversed the decline in ICDH activity caused by drought and resulted in a faster recovery of ICDH activity after rehydration. In the drought+DCPTA treatment compared with the control, ICDH decreased by 24.92% on

day 20 and by 19.94% on day 30 in 2016 and decreased by 19.88% on day 20 and by
13.63% on day 30 in 2017. Under well-watered conditions, the application of DCPTA
significantly increased the foliar ICDH activity on the 10<sup>th</sup>, 15<sup>th</sup>, 20<sup>th</sup> and 25<sup>th</sup> days in
2016 and on the 15<sup>th</sup>, 20<sup>th</sup>, and 25<sup>th</sup> days in 2017.

# 392 Contents of $NO_3^-$ , $NO_2^-$ and $NH_4^+$

393 The foliar  $NO_3^-$  and  $NO_2^-$  contents declined continuously during the drought 394 period and recovered during rehydration (Fig. 14). In the drought treatment compared with the control, the foliar  $NO_3^-$  and  $NO_2^-$  contents decreased by 39.82% and 38.27%, 395 396 respectively, on day 20 and by 33.62% and 35.11%, respectively, on day 30 in 2016 and decreased by 57.97% and 32.27%, respectively, on day 20 and by 42.38% and 397 398 25.21%, respectively, on day 30 in 2017. However, the DCPTA application partially reversed the decline in the foliar NO<sub>3</sub><sup>-</sup> and NO<sub>2</sub><sup>-</sup> contents caused by drought and 399 400 resulted in a faster recovery of the foliar NO<sub>3</sub><sup>-</sup> and NO<sub>2</sub><sup>-</sup> contents after rehydration. In 401 the drought+DCPTA treatment compared with the control, the foliar NO<sub>3</sub><sup>-</sup> and NO<sub>2</sub><sup>-</sup> 402 contents decreased by 25.94% and 23.31%, respectively, on day 20 and by 14.25% 403 and 18.55%, respectively, on day 30 in 2016 and decreased by 33.86% and 18.26%, respectively, on day 20 and by 17.26% and 8.60%, respectively, on day 30 in 2017. In 404 contrast, drought led to a marked elevation in the foliar  $NH_4^+$  content. In the drought 405 treatment compared with the control, the foliar  $NH_4^+$  increased by 52.48% on day 20 406 and by 29.18% on day 30 in 2016 and increased by 98.68% on day 20 and by 72.21% 407 408 on day 30 in 2017. In contrast, the DCPTA application suppressed the increase in the 409 foliar  $NH_4^+$  content induced by drought. In the drought+DCPTA treatment compared with the control, the foliar  $NH_4^+$  increased by 20.83% on day 20 and by 13.33% on 410 411 day 30 in 2016 and increased by 45.37% on day 20 and by 33.29% on day 30 in 2017. 412 Under well-watered conditions, DCPTA significantly increased the foliar NO<sub>3</sub><sup>-</sup> content on the 20<sup>th</sup> and 25<sup>th</sup> days in 2016 and on the 15<sup>th</sup>, 20<sup>th</sup>, 25<sup>th</sup> and 30<sup>th</sup> days in 413 2017. However, the DCPTA application had no significant effect on the foliar 414 contents of  $NO_2^-$  and  $NH_4^+$ . 415

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# 416 Activities of NR and NiR

417 The activities of foliar NR and NiR declined continuously during the drought period and recovered during rehydration (Fig. 15). In the drought treatment compared 418 419 with the control treatment, the activities of foliar NR and NiR decreased by 40.37% 420 and 36.91%, respectively, on day 20 and by 34.42% and 29.82%, respectively, on day 421 30 in 2016 and decreased by 52.80% and 32.46%, respectively, on day 20 and by 422 37.36% and 21.28%, respectively, on day 30 in 2017. However, the DCPTA 423 application partially reversed the declines in the activities of foliar NR and NiR 424 caused by drought and resulted in a faster recovery of the foliar NR and NiR activities 425 after rehydration. In the drought+DCPTA treatment compared with the control, the 426 activities of foliar NR and NiR decreased by 15.54% and 14.21%, respectively, on day 20 and by 10.79% and 7.79%, respectively, on day 30 in 2016 and decreased by 427 428 25.10% and 12.39%, respectively, on day 20 and by 13.55% and 4.96%, respectively, 429 on day 30 in 2017. Under well-watered conditions, DCPTA significantly increased the foliar NR activity on the 10<sup>th</sup>, 15<sup>th</sup> and 25<sup>th</sup> days in 2016 and on the 10<sup>th</sup>, 15<sup>th</sup>, 20<sup>th</sup> 430 and 25<sup>th</sup> days in 2017. Similarly, under well-watered conditions, DCPTA significantly 431 increased the foliar NiR activity on the 10<sup>th</sup> and 25<sup>th</sup> days in 2016 and on the 15<sup>th</sup> and 432 20<sup>th</sup> days in 2017. 433

# 434 Activities of GS, GOGAT and GDH

435 The activities of foliar GS and GOGAT first increased and then decreased during the drought period and recovered during rehydration (Fig. 16). In the drought 436 437 treatment compared with the control, the activities of foliar GS and GOGAT 438 decreased by 40.69% and 60.62%, respectively, on day 20 and by 33.65% and 51.39%, 439 respectively, on day 30 in 2016 and decreased by 47.57% and 66.84%, respectively, 440 on day 20 and by 37.96% and 56.77%, respectively, on day 30 in 2017. However, the 441 DCPTA application partially reversed the decline in the foliar GS and GOGAT activities caused by drought and resulted in a faster recovery of the foliar GS and 442 443 GOGAT activities after rehydration. In the drought+DCPTA treatment compared with

the control, the foliar GS and GOGAT activities decreased by 19.87% and 33.37%, 444 respectively, on day 20 and by 10.99% and 15.17%, respectively, on day 30 in 2016 445 and decreased by 23.56% and 33.02%, respectively, on day 20 and by 14.93% and 446 447 18.20%, respectively, on day 30 in 2017. Likewise, the DCPTA application alone caused an increase in the foliar GS and GOGAT activities. In contrast, drought led to 448 marked increases in the activities of foliar NAD-GDH and NADH-GDH. In the 449 drought treatment compared with the control, the activities of foliar NAD-GDH and 450 NADH-GDH increased by 87.16% and 150.92%, respectively, on day 20 and by 451 84.01% and 134.71%, respectively, on day 30 in 2016 and increased by 103.99% and 452 137.36%, respectively, on day 20 and by 96.13% and 111.20%, respectively, on day 453 454 30 in 2017. However, the DCPTA application partially reversed the increases in the 455 activities of foliar NAD-GDH and NADH-GDH caused by drought and resulted in a 456 faster recovery of the foliar NAD-GDH and NADH-GDH activities after rehydration. 457 In the drought+DCPTA treatment compared with the control, the activities of foliar 458 NAD-GDH and NADH-GDH increased by 49.55% and 92.59%, respectively, on day 20 and by 36.52% and 49.51%, respectively, on day 30 in 2016 and increased by 459 460 46.90% and 80.94%, respectively, on day 20 and by 35.43% and 45.60%, respectively, on day 30 in 2017. 461

Under well-watered conditions, DCPTA significantly increased the foliar GS activity on the 20<sup>th</sup> and 25<sup>th</sup> days in 2016 and on the 10<sup>th</sup> 20<sup>th</sup> and 25<sup>th</sup> days in 2017. Similarly, under well-watered conditions, DCPTA significantly increased the foliar GOGAT activity on day 15 in 2017 and significantly increased the foliar NAD-GDH activity on day 15 in 2016

## 467 Activities of AlaAT and AspAT

The activities of foliar AlaAT and AspAT first increased and then continuously decreased during the drought period and recovered during rehydration (Fig. 17). In the drought treatment compared with the control, the activities of foliar AlaAT and AspAT decreased by 44.18% and 65.51%, respectively, on day 20 and by 36.43% and 38.52%,

respectively, on day 30 in 2016 and decreased by 56.80% and 54.92%, respectively, 472 473 on day 20 and by 41.43% and 41.29%, respectively, on day 30 in 2017. However, the 474 DCPTA application partially reversed the decline in the activities of foliar AlaAT and 475 AspAT caused by drought and resulted in a faster recovery of the foliar AlaAT and 476 AspAT activities after rehydration. In the drought+DCPTA treatment compared with 477 the control, the activities of foliar AlaAT and AspAT decreased by 17.45% and 35.39%, respectively, on day 20 and by 15.84% and 17.17%, respectively, on day 30 478 in 2016 and decreased by 32.49% and 37.82%, respectively, on day 20 and by 17.46% 479 and 18.21%, respectively, on day 30 in 2017. 480

Under well-watered conditions, DCPTA significantly increased the foliar AlaAT activity on the 10<sup>th</sup>, 15<sup>th</sup>, and 20<sup>th</sup> days in 2016 and on the 10<sup>th</sup>, 15<sup>th</sup>, 20<sup>th</sup> and 25<sup>th</sup> days in 2017. Similarly, under well-watered conditions, DCPTA significantly increased the foliar AspAT activity on day 15 in 2016 and on the 5<sup>th</sup>, 10<sup>th</sup>, 15<sup>th</sup> and 25<sup>th</sup> days in 2017.

# 486 **Protease activity and contents of proteins and free amino acids**

The protease activity and free amino acid contents increased continuously 487 during the drought period and decreased during rehydration (Fig. 18). In the drought 488 treatment compared with the control, the protease activity and free amino acid 489 contents increased by 122.48% and 88.92%, respectively, on day 20 and by 55.03% 490 491 and 34.51%, respectively, on day 30 in 2016 and increased by 145.15% and 78.58%, 492 respectively, on day 20 and by 57.08% and 43.29%, respectively, on day 30 in 2017. 493 However, the DCPTA application partially reversed the increases in the protease 494 activity and free amino acid contents caused by drought.

In the drought+DCPTA treatment compared with the control, the protease activity and free amino acid contents increased by 78.47% and 24.77%, respectively, on day 20 and by 16.46% and 6.62%, respectively, on day 30 in 2016 and increased by 65.41% and 30.24%, respectively, on day 20 and 14.50% and 13.57%, respectively, on day 30 in 2017.

In contrast, drought led to a marked decrease in the foliar protein content. In the 500 501 drought treatment compared with the control, the foliar protein content decreased by 35.51% on day 20 and by 18.32% on day 30 in 2016 and decreased by 44.81% on day 502 503 20 and by 22.50% on day 30 in 2017. In contrast, the DCPTA application suppressed 504 the increase in the foliar protein content induced by drought. In the drought+DCPTA 505 treatment compared with the control, the foliar protein content increased by 19.54% on day 20 and by 7.25% on day 30 in 2016 and increased by 25.46% on day 20 and 506 by 10.30% on day 30 in 2017. 507

508 Significant differences between the foliar protein contents of the well-watered 509 treatment and well-watered+DCPTA treatment were observed on day 10 in 2017 and 510 on day 15 in 2017, and significant differences between the free amino acid contents of 511 these treatments were observed on day 10 in 2017.

## 512 Discussion

513 One of the most clear and consistent effects of drought on crops is the inhibition 514 of growth and yield (Sinclair, 2011). In this experiment, exogenous DCPTA partly 515 mitigated the reductions in plant growth and yield induced by drought, as expressed 516 by the stable soot and root RGR, GN and GY (Fig. 5, Fig. 6 and Fig. 7). Moreover, 517 exogenous DCPTA promoted growth and yield under well-watered conditions.

518 N is a necessary macro-nutrient element for plants and a main limiting factor in 519 plant growth and development (Kusano et al., 2011). The pre-female inflorescence 520 emergence stage is the important stage determining maize yield (Talaat et al., 2015). 521 The inhibition of maize growth induced by drought could be partly attributed to nitrogen metabolism during the pre-female inflorescence emergence stage. NO<sub>3</sub><sup>-</sup> is 522 523 the main nitrogen source assimilated by higher plants in agricultural soils (Robredo et al., 2011). Similar to previous reports for tomato (Sánchez-Rodríguez et al., 2011) and 524 525 wheat (Fresneau *et al.*, 2007), drought significantly diminished the  $NO_3^-$  content in 526 maize leaves in both the DCPTA-treated and non-treated leaves (Fig. 14). This decrease may be explained by drought-induced inhibitions in nitrate uptake from the 527

roots and/or nitrate transport. However, the reduction in the non-treated leaves wasgreater than that in the DCPTA-treated leaves in this study.

Maintaining the supply of nutrients and water from the soil to crop depends on 530 531 root morphology, which can be characterized by RLD and RSD (Kamran *et al.*, 2018). Previous studies have reported that DCPTA promotes root development in tomatoes 532 (Keithly et al., 1990a). In this study, there was a significant difference between the 533 RLD and RSD of the well-watered treatment and DCPTA treatment in 0-20 cm and 534 0-30 cm in 2016 and 2017 on day 30 (Fig. 8). This result suggests that DCPTA 535 promoted root development in maize under well-watered conditions. Interestingly, 536 under drought conditions, the DCPTA application significantly increased RLD, RSD 537 538 and the root RGR. These results indicate that the DCPTA application also promoted 539 maize root growth and improved the spatial and temporal distribution of roots, which 540 was beneficial to  $NO_3^-$  uptake under drought conditions.

The xylem sap transports water and nutrients from the roots throughout the plant 541 and depends on transpiration intensity and root pressure. The increased root hydraulic 542 conductivity and flow rate of root-bleeding sap induced by the DCPTA application 543 may, due to the enhanced root pressure, which depend on physiological activity of the 544 545 whole root system (Fig. 9) (Noguchi et al., 2005; Lui et al., 2014). In addition, the 546  $NO_3^-$  delivery rate in the presence of DCPTA was significantly higher than that without DCPTA under drought conditions, which may partly result from the improved 547  $NO_3^-$  absorption and enhanced root pressure induced by DCPTA. The stable RLWC 548 549 in the drought+DCPTA treatment suggests an abundant supply water to the aboveground parts, and balanced transpirational loss and water uptake under drought 550 551 conditions; as a result, the DCPTA-treated plants maintained Gs, which reduced the leaf epidermal resistance and promoted the mass flow of water to the leaf surface and 552 553 the transportation of the  $NO_3^-$  required for N metabolism in leaves (Fig. 10 and Fig. 554 11). Under drought conditions, increases in the foliar  $NO_3^-$  were observed in the 555 DCPTA-treated plants (Fig. 14).

556

Whether stomatal or non-stomatal factors are the main cause of a reduced Pn

may be determined by changes in Gs and Ci (Bethke and Drew, 1992). During the
early period of drought, the change of Ci were accompanied by continuously declined
Gs, then Gs decreases but Ci shows an increase (Fig. 11).

560 Thus, the decrease of the Pn in drought-treated plants was mainly attributed to stomatal limitations firstly, and then, non-stomatal limitations induced by the damage 561 of photochemical mechanism, partly reflected by damaged chloroplast (Fig. 12). 562 However, DCPTA application maintains relatively high Gs, ensuring the availability 563 of  $CO_2$  for the carbon reduction cycle. Simultaneously, the DCPTA application 564 delayed the increase in Ci and protected the chloroplast ultrastructure against 565 drought-induced oxidative damage, which suggests that DCPTA can protect the 566 567 photochemical mechanism and, as a result, ensures a more efficient photosynthesis 568 process after rehydration. Moreover, similar to previous studies on spruce (Keithly et 569 al., 1990a), sugar beets (Keithly et al., 1990b) and guayule (Wan and Mendoza, 1992), DCPTA application also promoted photosynthesis under well-watered conditions. 570

571 In most plants, nitrate reduction occurs in leaves. NO<sub>3</sub>, after being taken up into the leaf cell, is converted to  $NH_4^+$  by two successive steps catalysed by NR and NiR. 572 NR, the rate-limiting enzyme of nitrogen assimilation, is highly sensitive to stress 573 574 (Plett *et al.*, 2016). Similar to previous studies on wheat and barley, the NR activity 575 continuously declined in response to drought (Fig. 15) (Robredo et al., 2011; Fresneau et al., 2007). As a typical nitrate-induced enzyme, NR activity is primarily 576 regulated by the  $NO_3^-$  concentration in the leaves (Chamizo-Ampudia *et al.*, 2017). 577 578 The up-regulation of foliar NR activity in the drought+DCPTA treatment may result from the increase in the foliar  $NO_3^-$  content (Fig. 14). Moreover, the reduction in the 579 foliar NiR activity under drought conditions was significantly reversed by the 580 application of DCPTA, which may be because the DCPTA-stabilized photosynthesis 581 582 resulted in a sufficient supply of Fd<sub>red</sub> (Fig. 11 and Fig. 15), thus promoting the conversion of  $NO_2^-$  to  $NH_4^+$ . The present results indicate that DCPTA pretreatment 583 584 could maintain a high NO<sub>3</sub><sup>-</sup> assimilation ability in maize under drought conditions.

585 Although the foliar NR and NiR activities declined during the drought period, the

foliar  $NH_4^+$  content exhibited an increasing tendency in our experiment (Fig. 14). This increase may be associated with the glycine oxidation in activated photorespiration, which is induced by decreases in *Ci* levels under drought conditions (Wang *et al.*, 2012). The increased *Gs* induced by DCPTA was beneficial to the increase in CO<sub>2</sub> in the cellular spaces of the leaf, implying that photorespiration was partly alleviated (Fig. 11).

In plants cells, excessive levels of  $NH_4^+$  are destructive, and the major  $NH_4^+$ 592 assimilation pathway is the GS/GOGAT cycle in higher plants. When the 593 GS/GOGAT cycle is suppressed and the NH<sub>4</sub><sup>+</sup> content rises continuously under 594 stress,  $NH_4^+$  could serve as a substrate to form glutamate via the reversible amination 595 of 2-OG by GDH catalyse, although the enzyme has a lower affinity for NH<sub>4</sub><sup>+</sup> 596 (Fontaine *et al.*, 2012). In general, drought inhibited  $NH_4^+$  assimilation (Nava *et al.*, 597 2007). During the early period of drought, GDH activity increased sharply, GS 598 activity increased slightly, and GOGAT remained stable (Fig. 16). These results 599 suggest that accelerated  $NH_4^+$  assimilation in maize may be an adaptive mechanism 600 to produce more glutamate and eliminate the accumulation of excess foliar  $NH_4^+$ . 601 Subsequently, the GS and GOGAT activities decreased, which may have resulted 602 from an inadequate supply of energy and 2-OG because of photosynthetic inhibition 603 604 and decreased ICDH activity (Fig. 13). GDH activity decreased with drought, which may be due to the oxidative degradation of GS caused by the overproduction of 605 reactive oxygen species (ROS) induced by drought conditions (Xia et al., 2015). 606

The DCPTA application altered the major  $NH_4^+$  assimilation pathway, 607 maintained the GOGAT/GS cycle and suppressed the GDH pathway, which may 608 have contributed to maintaining the conversion of  $NH_4^+$  to glutamine and the 609 subsequent formation of glutamate from glutamine. This result may occur because 610 611 the photosynthetic stability and ICDH activity induced by the DCPTA application promoted 2-OG synthesis and the reducing power (i.e., NADPH, ATP, or  $Fd_{red}$ ) in 612 plants during the drought period, thus providing the GS/GOGAT cycle with 613 relatively sufficient substrates and energy and favouring the enhancement of foliar 614

GOGAT and GS activities (Du *et al.*, 2016). As a result, with the application of
DCPTA, drought had less of an effect on the activities of GS and GOGAT.

Although DCPTA promoted  $NO_3^-$  assimilation, as expressed by the increased NR and NiR activities (Fig. 15), this treatment compared to the drought treatment caused significant decreases in the  $NH_4^+$  content, which means that exogenous DCPTA resulted in the integration of  $NH_4^+$  into the structure of organic compounds, thereby contributing to the reduction in the  $NH_4^+$  content. Therefore, the DCPTA application effectively modulated the activities of ICDH, GS, GOGAT and GDH and accelerated the conversion of  $NH_4^+$  to glutamate, which is the precursor of other amino acids.

Transamination is a key step in the biosynthesis of various amino acids from 624 625 glutamate, with the availability of C skeletons from the Krebs cycle (Hodges, 2002). 626 In our studies, both the aminotransferases studied, AlaAT and AspAT, showed increased activity in maize during the early drought period (Fig. 17). Such increases 627 in aminotransferases activities under drought conditions might help in the synthesis of 628 increased amounts of amino acids that act as compatible cytoplasmic solutes and 629 protect cell organelles and biomolecules, thus reducing the adverse effects of drought 630 on maize (Munns and Tester, 2008). Subsequently, the AlaAT and AspAT activities 631 decreased, which may be attributable to the weakened GS/NADH-GOGAT pathway 632 633 (Fig. 16) (Gangwar and Singh, 2011). Moreover, stable aminotransferase activities were observed in DCPTA-treated plants. This finding may be associated with 634 increased GS/GOGAT activities, which can generate more glutamate to serve as a 635 substrate for transamination reactions in maize treated with DCPTA under drought 636 conditions. 637

Most soluble proteins are enzymes that participate in various metabolic pathways in plants; therefore, the soluble protein content is considered one of the most important indices reflecting the overall metabolic level in plants. Protein synthesis in plants is very sensitive to abiotic stresses and is positively correlated with stress tolerance (Fresneau *et al.*, 2007). Free amino acids are the building blocks of proteins. Drought increased the free amino acid contents, which may mainly be

attributed to the increased AlaAT and AspAT activities in the early drought period 644 and the subsequent promotion of protein degradation (Fig. 18) (Yang et al., 2013). 645 However, DCPTA-treated seedlings maintained higher soluble protein levels and 646 647 lower free amino acid levels than did non-DCPTA-treated seedlings in response to drought. This result may occur because DCPTA inhibited protein degradation by 648 649 stable protease activities and maintained protein stability, ensuring the series of physiological and biochemical processes that occur normally under stress conditions. 650 Additionally, the DCPTA application increased the amino acid contents under 651 well-watered conditions, which may be attributable to the promoted biosynthesis and 652 accumulation of amino acids, which ultimately improved plant growth and 653 654 development (Talaat and Shawky, 2016).

#### 655 **Conclusions**

The present study suggested that DCPTA treatment increased NO<sub>3</sub><sup>-</sup> uptake and 656 the long-distance transportation of  $NO_3^-$  from the roots to the leaves via the 657 production of excess roots and maintained a stabilized transpiration rate. The 658 increased foliar NO<sub>3</sub><sup>-</sup> content up-regulated NR activity and maintained a high N 659 assimilation ability that was restrained by drought. Exogenous DCPTA effectively 660 661 regulated the ICDH, GS, GOGAT and GDH activities to speed up the conversion of  $NH_4^+$  to Glu, reduced the toxicity of excess  $NH_4^+$  to the plant, and accelerated the 662 663 synthesis of proteins and amino acids. Moreover, DCPTA treatment maintained increased the photosynthetic capacity, supply nitrogen metabolism of energy and 664 carbon skeleton thus alleviating the inhibition of growth by drought in maize. 665

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876	Table 1 Chemical properties of the used soil.										
	Year	pН	HCO <sub>3</sub> <sup>-</sup>	Cl	$SO_4^{2-}$	$\operatorname{Ca_2}^+$	$Mg_2^+$	$Na^+$	$\mathbf{K}^+$	N	Р
			+	(mg	(mg	(mg	(mg	(mg	(mg	(mg	(mg
			CO3 <sup>2-</sup>	kg <sup>-1</sup> )	kg <sup>-1</sup> )	kg <sup>-1</sup> )	kg <sup>-1</sup> )	kg <sup>-1</sup> )	kg <sup>-1</sup> )	kg <sup>-1</sup> )	kg <sup>-1</sup> )
			(mg								
			kg <sup>-1</sup> )								
	2016	7.2	204.3	297.6	463.8	87.4	37.6	4.1	28.9	17.3	3.9
	2017	7.1	201.7	302.5	446.2	90.3	40.9	4.0	31.4	16.1	3.8
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#### 897 Figure legends:

- **Fig. 1** The rainfall (bar) and mean temperature (line) data during the study period (2016 and 2017, May–October).
- Fig. 2 The pits (inner length, 10 m; width, 7 m; and height, 1.2 m) used for this studyand the plastic sheets used to cover inner sides of the pits.
- **Fig. 3** Changes in the soil water content (SWC) in 2016 and 2017. The data represent
- 903 the means of independent measurements with five replicates, and the standard
- deviations are indicated by the vertical error bars. Values with the same letters on the
- bars are not significantly different at P < 0.05 (LSD test).
- Fig. 4 The tools used for the collection of bleeding sap. (A) cotton, (B) centrifuge
  tube, (C) scissors, (D) plastic film, and (E) deionized water.
- **Fig. 5** Changes in the grain number plant<sup>-1</sup> and grain yield plant<sup>-1</sup> (g) of maize in 2016 and 2017. The data represent the means of independent measurements with five replicates, and the standard deviations are indicated by the vertical error bars. Values with the same letters on the bars are not significantly different at P<0.05 (LSD test).
- Fig. 6 Leaf phenotypic features of the maize seedlings after 20 days of treatment with
  drought and/or DCPTA in 2016 and 2017.
- **Fig. 7** Effect of drought and/or DCPTA treatment on the relative growth rate (RGR) of the shoots and roots of the maize in 2016 and 2017. The data represent the means of independent measurements with five replicates, and the standard deviations are indicated by the vertical error bars. Values with the same letters on the bars are not significantly different at P<0.05 (LSD test).
- Fig. 8 Effect of drought and/or DCPTA treatment on root length density (RLD) and
  root surface area density (RSD) for different soil depths on the 20<sup>th</sup> day and 30<sup>th</sup> day
  after treatment in 2016 and 2017.
- **Fig. 9** Effect of drought and/or DCPTA treatment on root hydraulic conductivity, the flow rate of root-bleeding sap and the  $NO_3^-$  delivery rate in 2016 and 2017. The data represent the means of independent measurements with five replicates, and the standard deviations are indicated by the vertical error bars. Values with the same

926 letters on the bars are not significantly different at P<0.05 (LSD test).

**Fig. 10** Effect of drought and/or DCPTA treatment on relative water content of the leaves (RLWC) in 2016 and 2017. The data represent the means of independent measurements with five replicates, and the standard deviations are indicated by the vertical error bars. Values with the same letters on the bars are not significantly different at P<0.05 (LSD test).

**Fig. 11** Effect of drought and/or DCPTA treatment on the gas exchange parameters in the maize leaves in 2016 and 2017. Pn, net photosynthetic rate; Gs, stomatal conductance; Tr, transpiration rate; Ci, intercellular  $CO_2$  concentration. The data represent the means of independent measurements with five replicates, and the standard deviations are indicated by the vertical error bars. Values with the same letters on the bars are not significantly different at P<0.05 (LSD test).

Fig. 12 Ultrastructure of the photosynthetic apparatus in the maize leaves after 20
days of treatment with drought and/or DCPTA in 2016 and 2017. SL, stroma lamella;
GL, grana lamellae; CW, cell wall; and P, plastoglobule. The scale bars for the
photosynthetic apparatus represent 2000 nm.

**Fig. 13** Effects of drought and/or DCPTA treatment on isocitrate dehydrogenase (ICDH) activity in 2016 and 2017. The data represent the means of independent measurements with five replicates, and the standard deviations are indicated by the vertical error bars. Values with the same letters on the bars are not significantly different at P<0.05 (LSD test).

**Fig. 14** Effects of drought and/or DCPTA treatment on the nitrate (NO<sub>3</sub><sup>-</sup>), nitrite (NO<sub>2</sub><sup>-</sup>) and ammonium (NH<sub>4</sub><sup>+</sup>) contents in the leaves of maize in 2016 and 2017. The data represent the means of independent measurements with five replicates, and the standard deviations are indicated by the vertical error bars. Values with the same letters on the bars are not significantly different at P<0.05 (LSD test).

**Fig. 15** Effects of drought and/or DCPTA treatment on nitrate reductase (NR) and nitrite reductase (NiR) activities in the leaves of maize in 2016 and 2017. The data represent the means of independent measurements with five replicates, and the

standard deviations are indicated by the vertical error bars. Values with the same
letters on the bars are not significantly different at P<0.05 (LSD test).</li>

**Fig. 16** Effects of drought and/or DCPTA treatment on the glutamine synthetase (GS), glutamate synthase (GOGAT) and glutamate dehydrogenase (GDH) activities in the leaves of maize in 2016 and 2017. The data represent the means of independent measurements with five replicates, and the standard deviations are indicated by the vertical error bars. Values with the same letters on the bars are not significantly different at P<0.05 (LSD test).

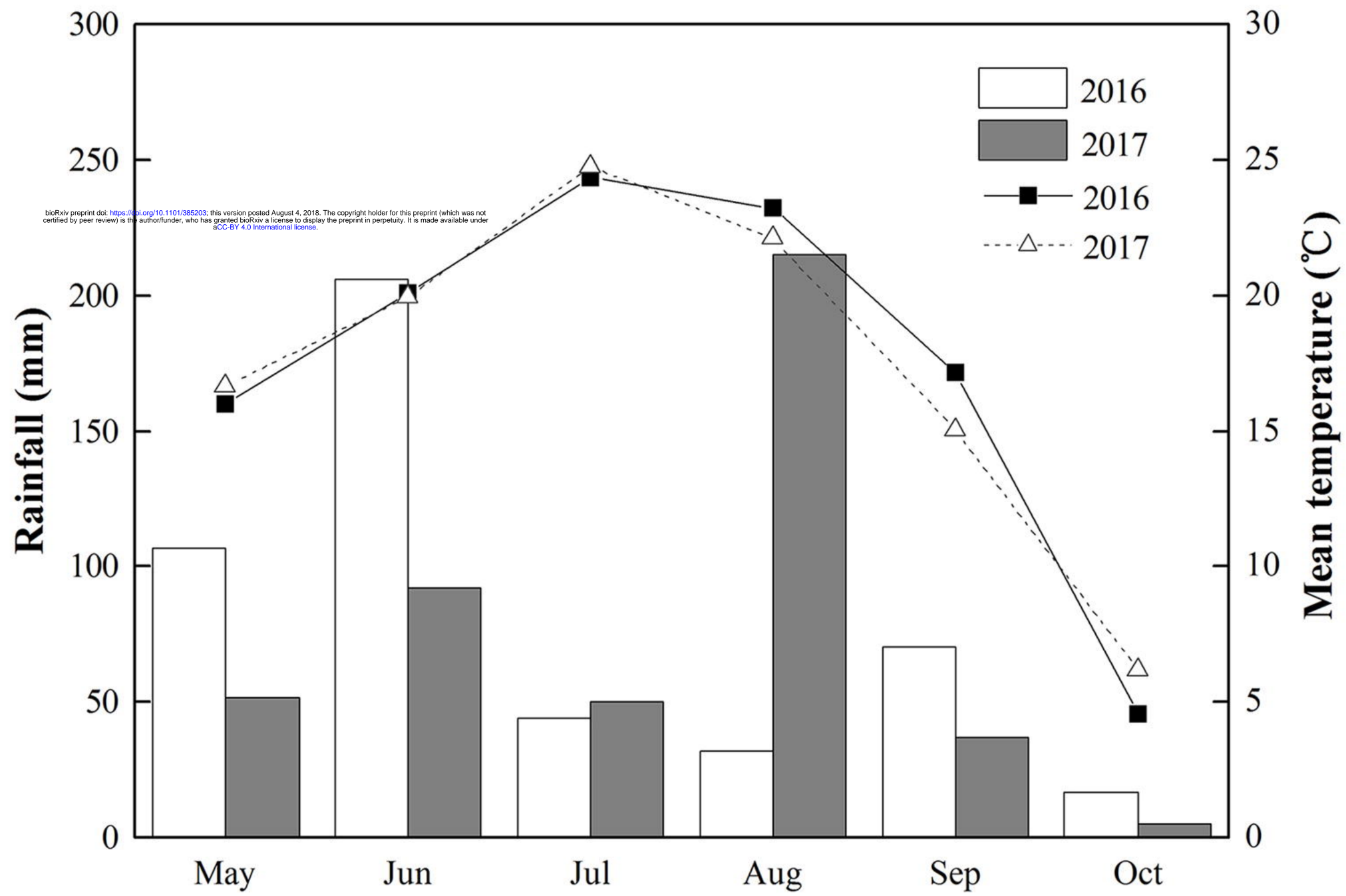
**Fig. 17** Effects of drought and/or DCPTA treatment on the alanine aminotransferase

964 (AlaAT) and aspartate aminotransferase (AspAT) activities in the leaves of maize in

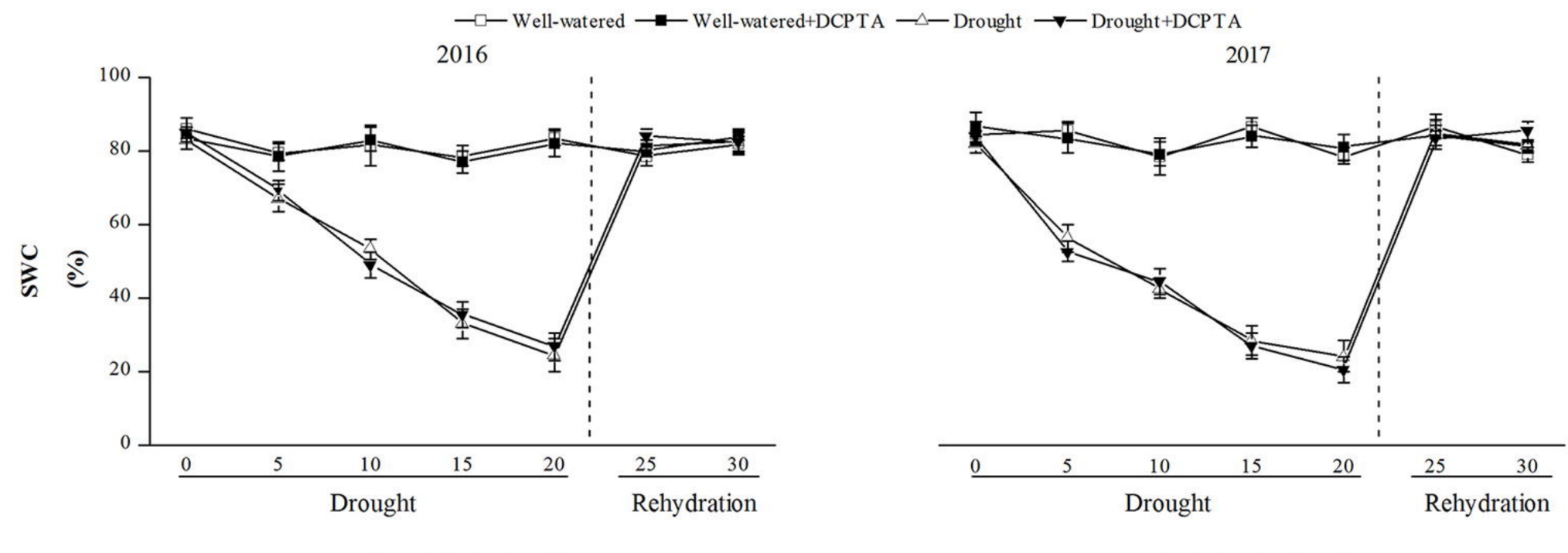
2016 and 2017. The data represent the means of independent measurements with five

replicates, and the standard deviations are indicated by the vertical error bars. Values

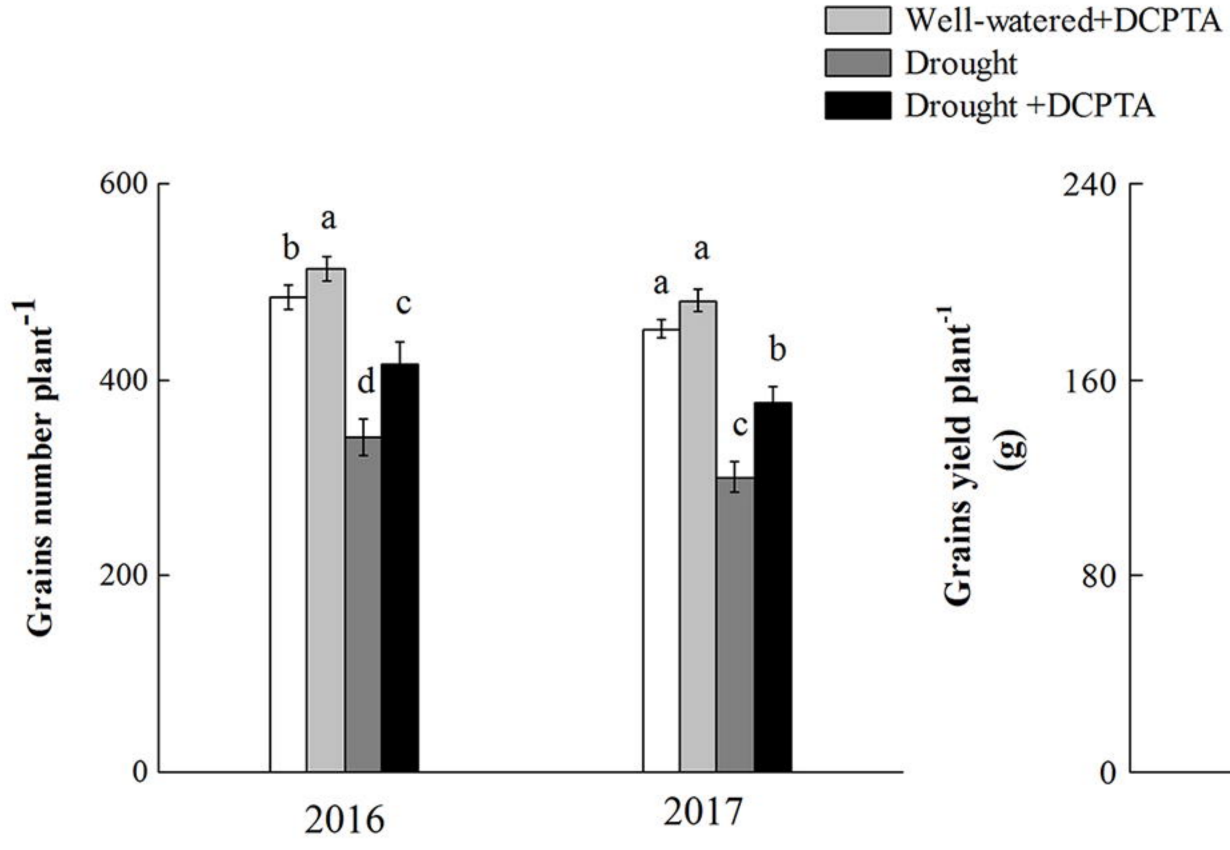
- with the same letters on the bars are not significantly different at P<0.05 (LSD test).
- **Fig. 18** Effects of drought and/or DCPTA treatment on the protease activity and protein and free amino acid contents in the leaves of maize in 2016 and 2017. The data represent the means of independent measurements with five replicates, and the standard deviations are indicated by the vertical error bars. Values with the same letters on the bars are not significantly different at P<0.05 (LSD test).



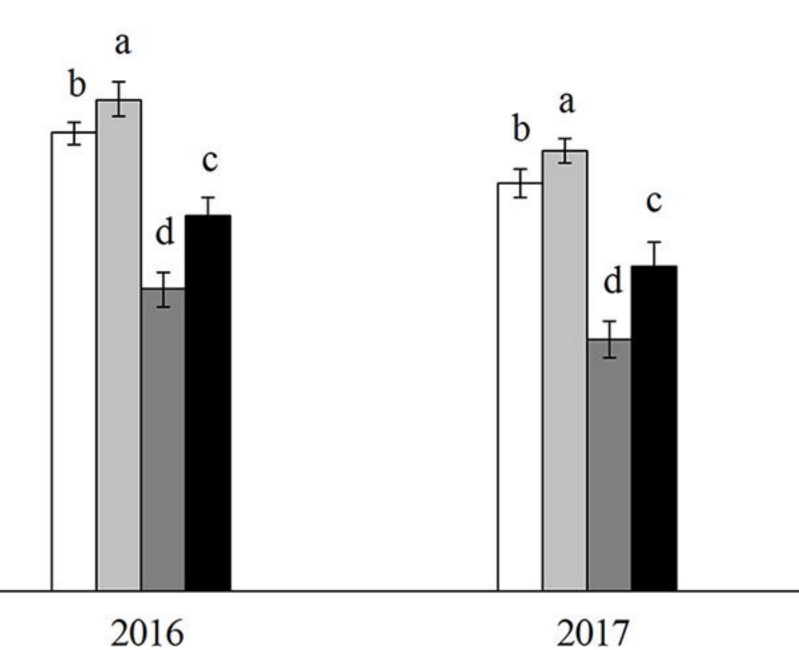








Well-watered



2016



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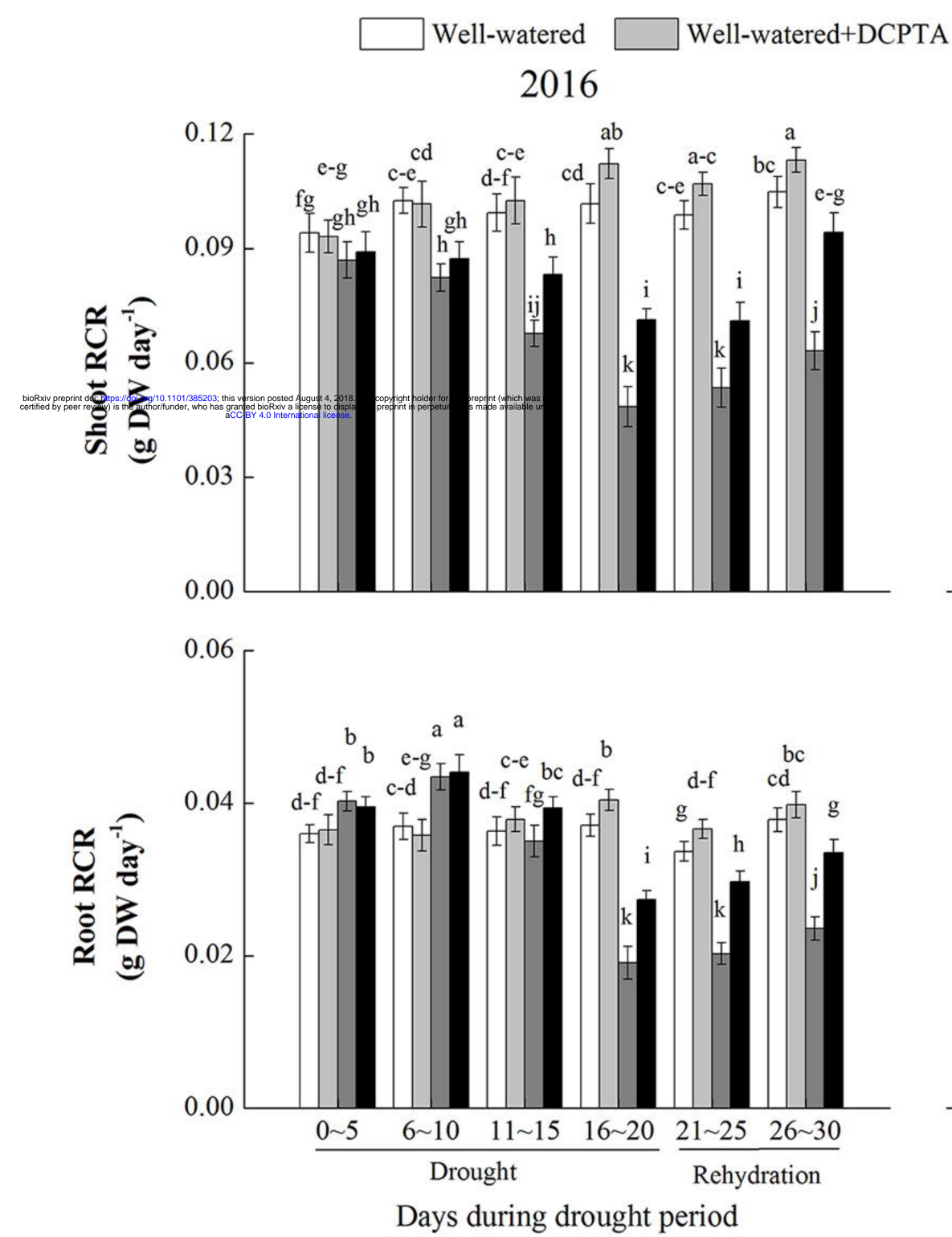
2017

#### Well-watered Well-watered

+DCPTA

#### Drought

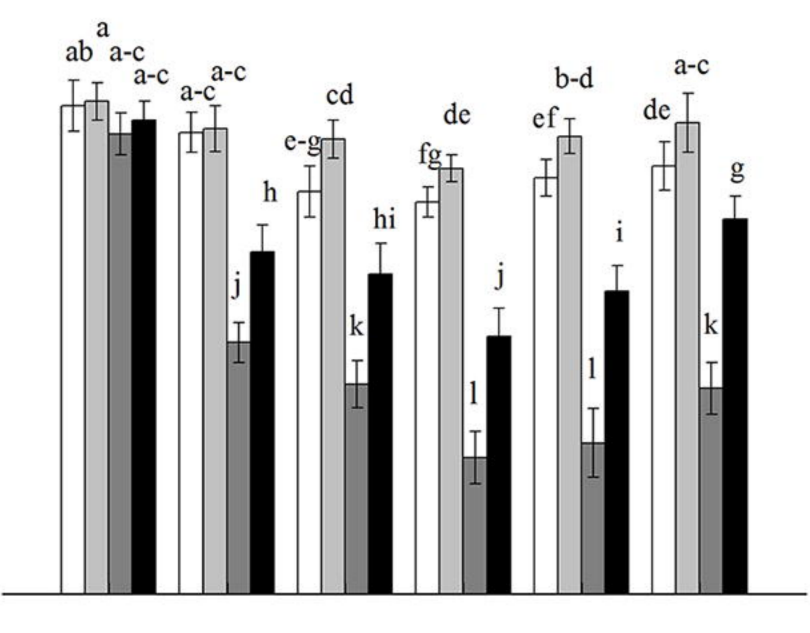
## Drought +DCPTA

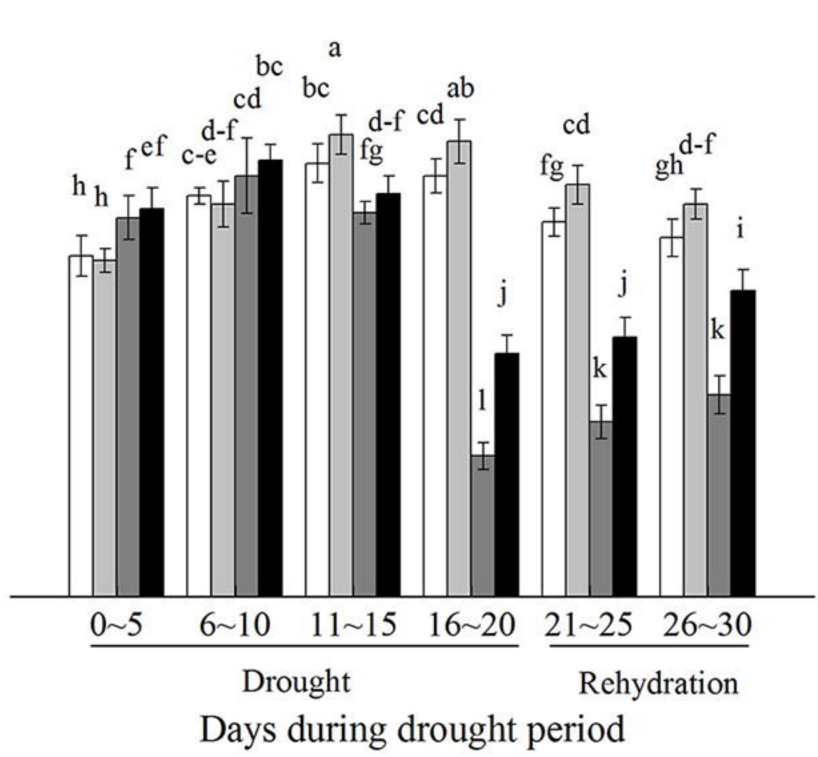


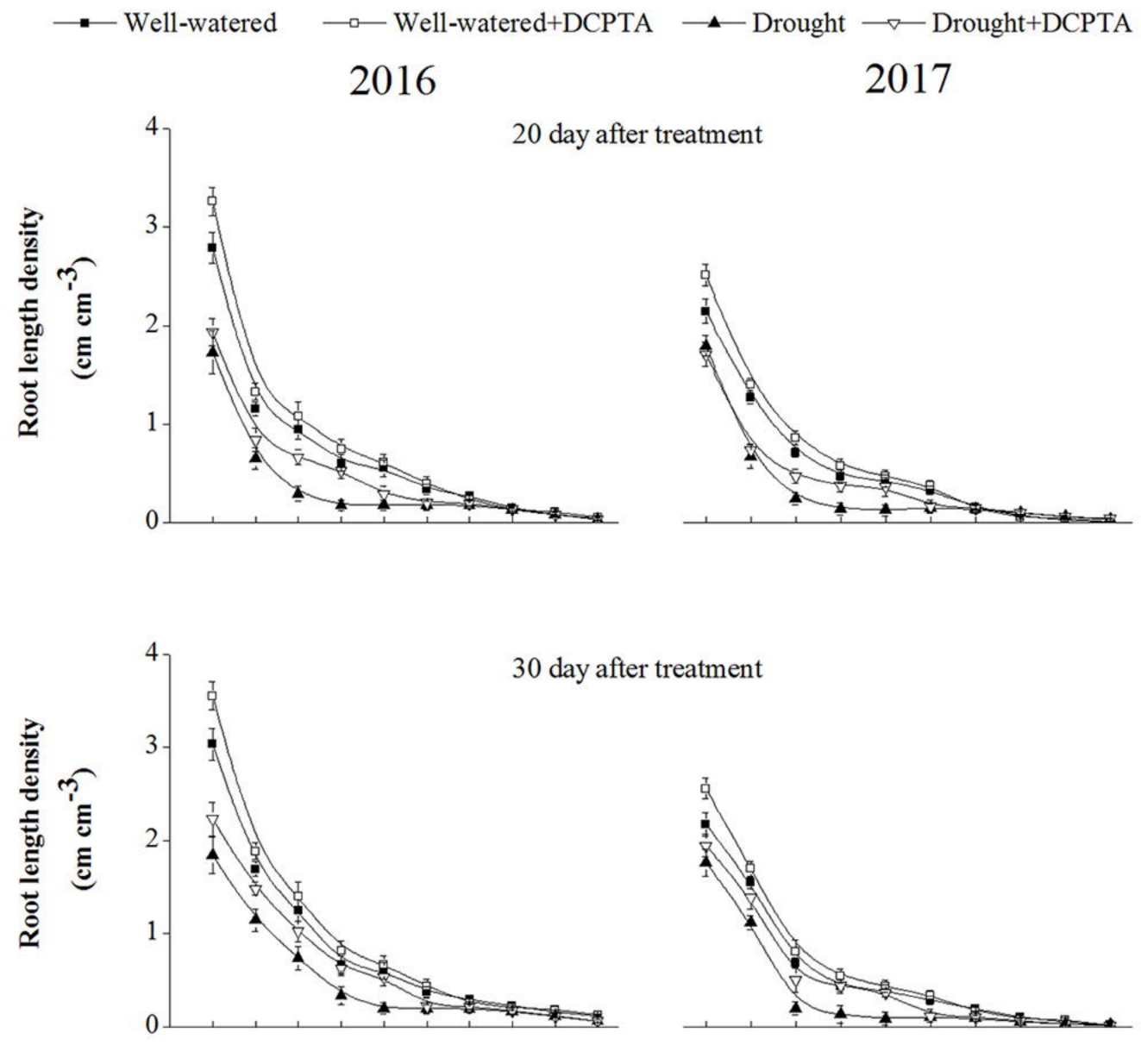


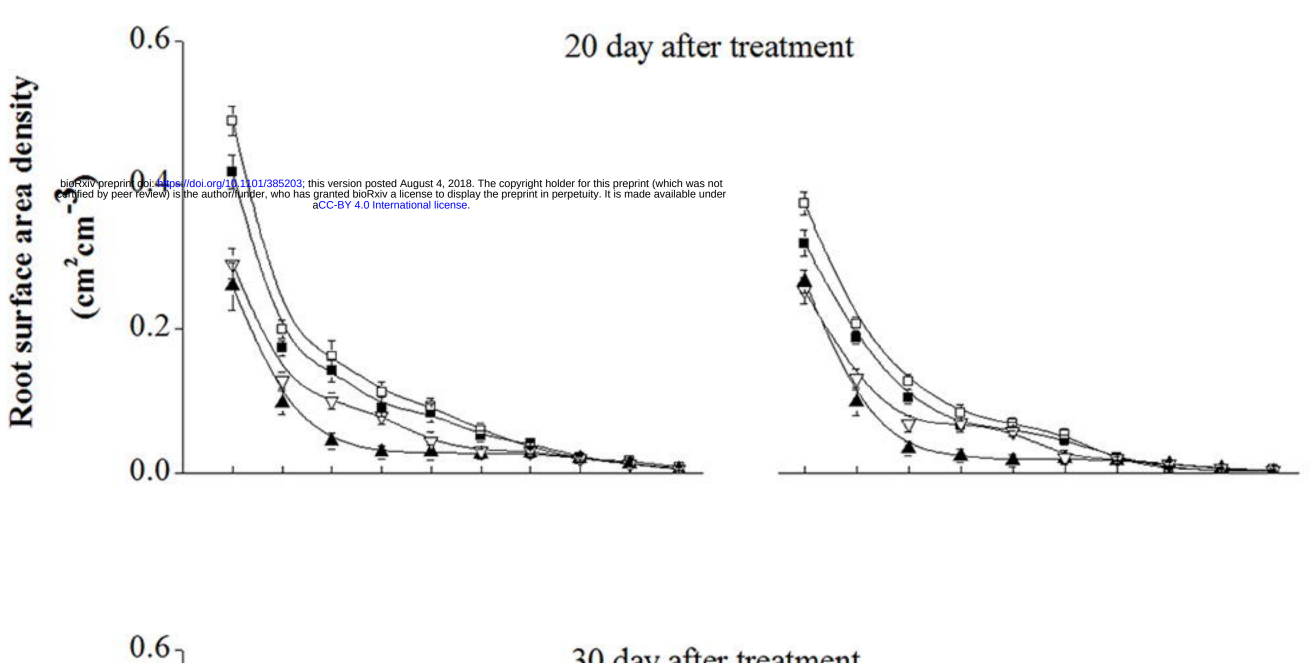
Drought +DCPTA

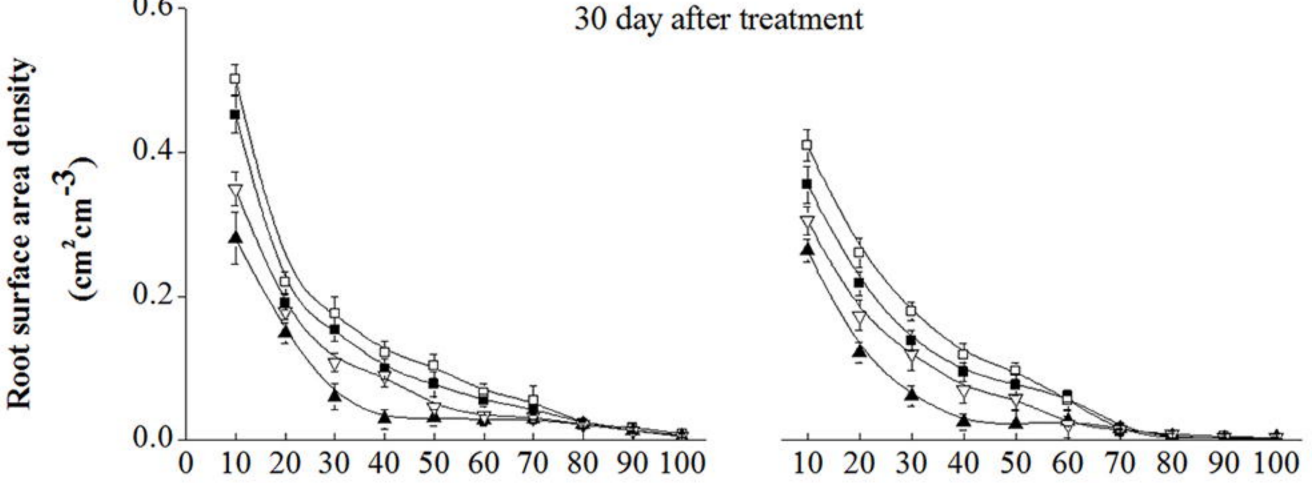
2017

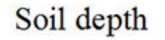




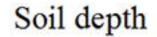




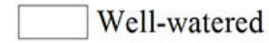




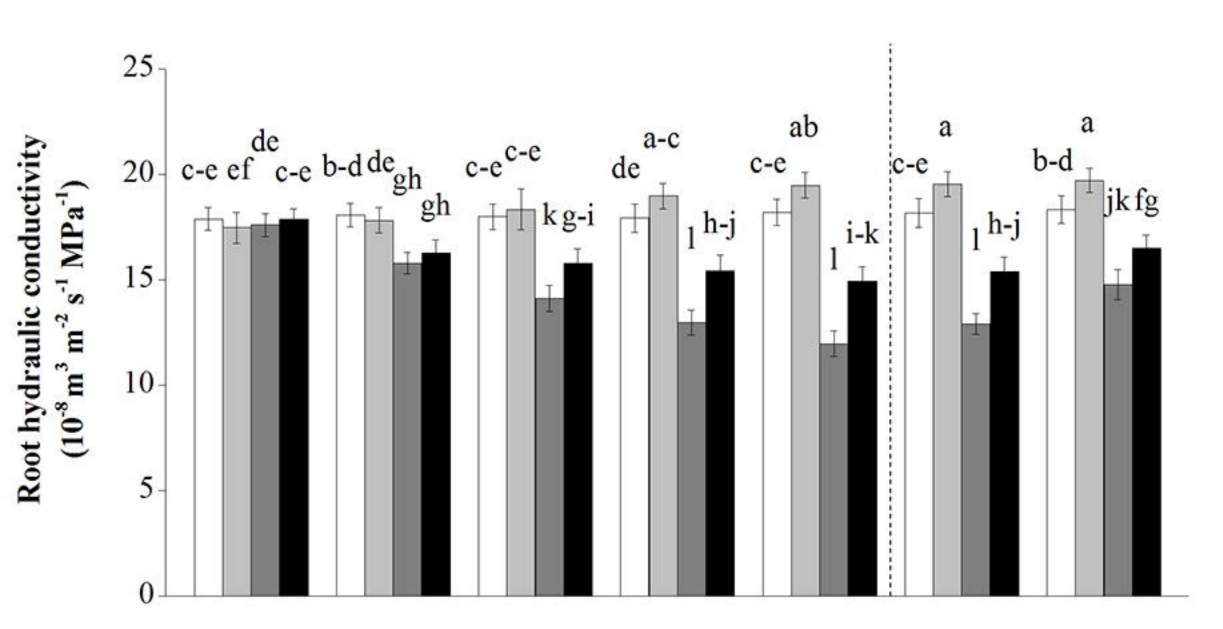


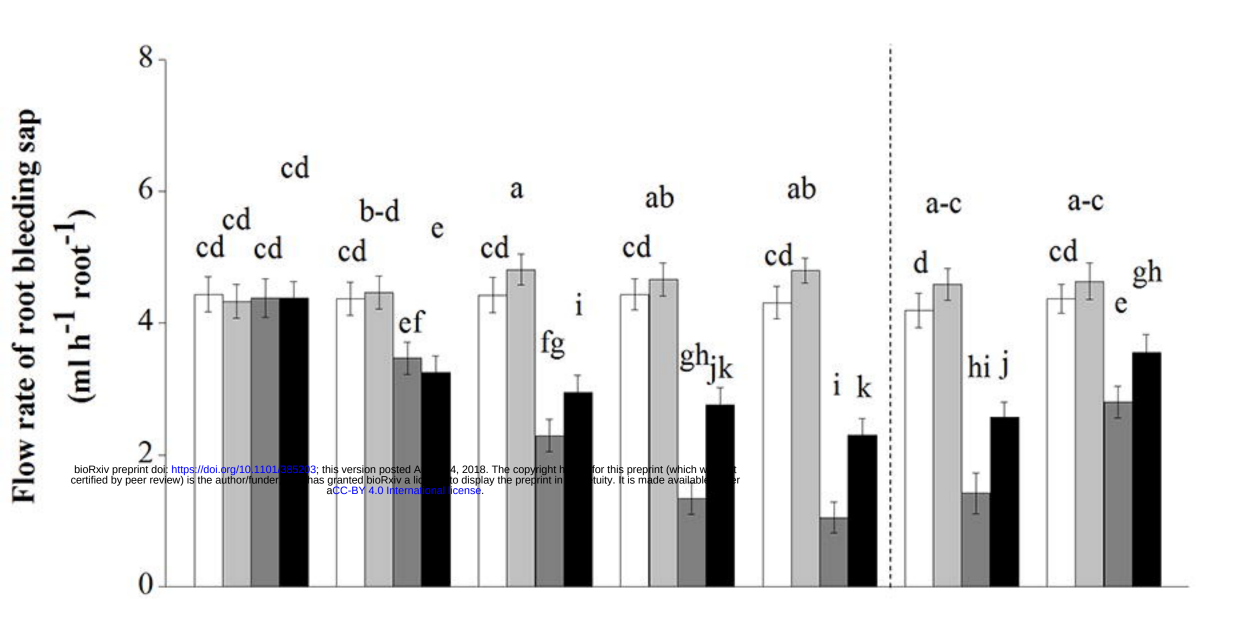


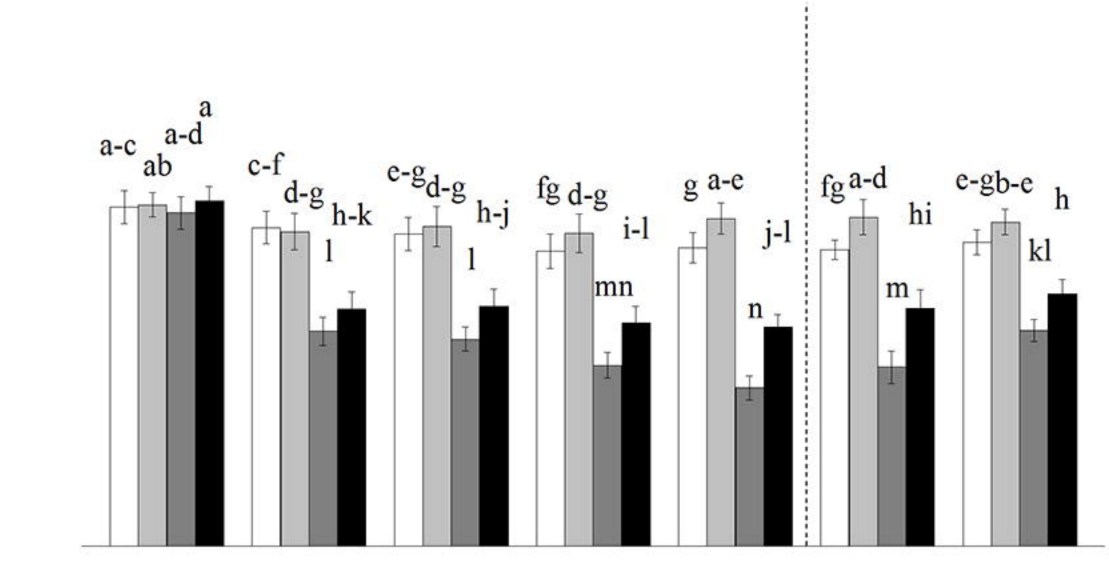


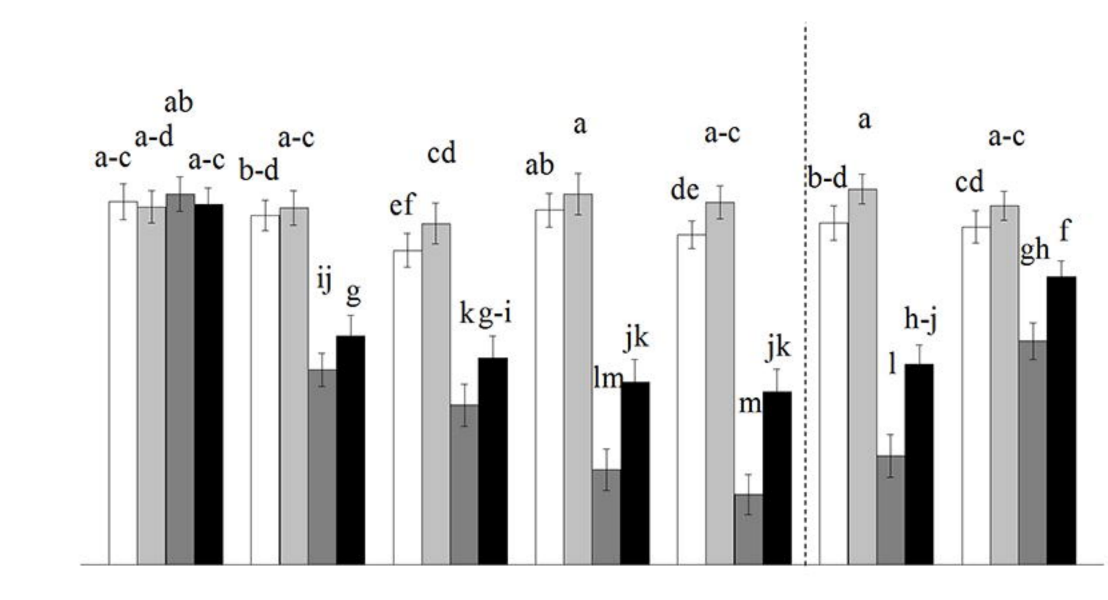


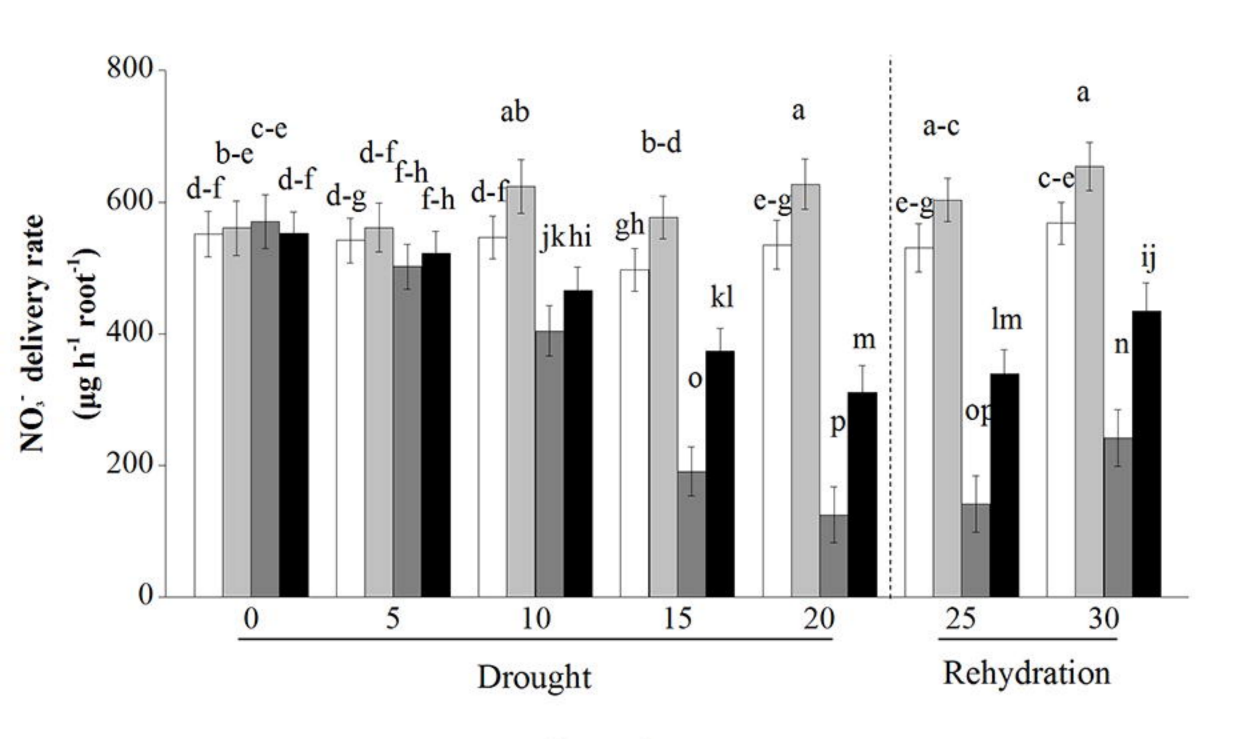


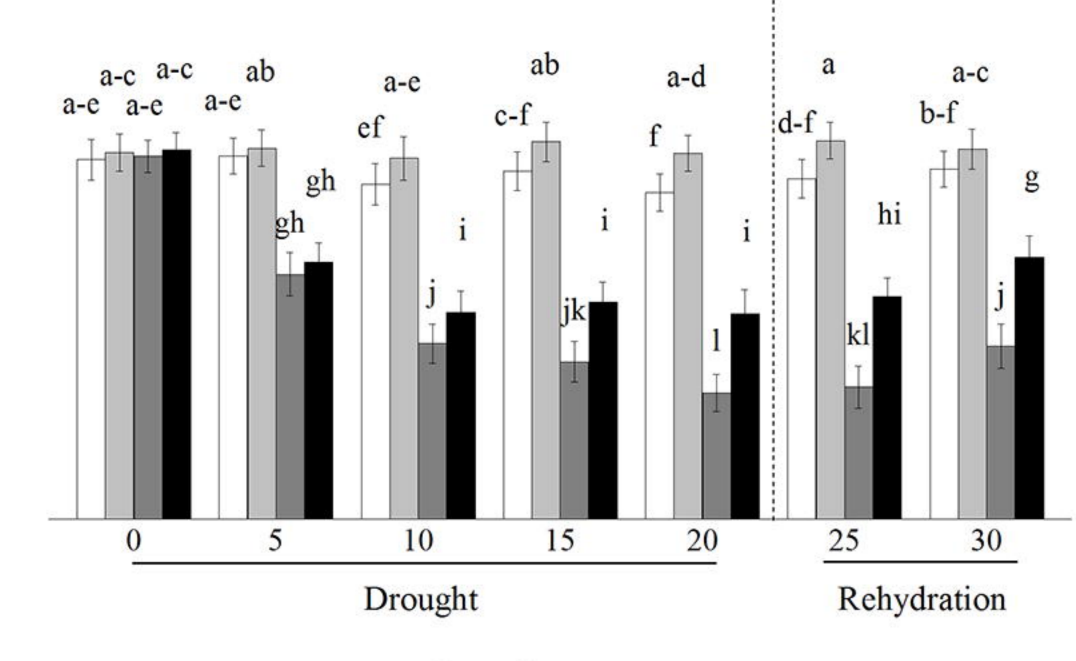




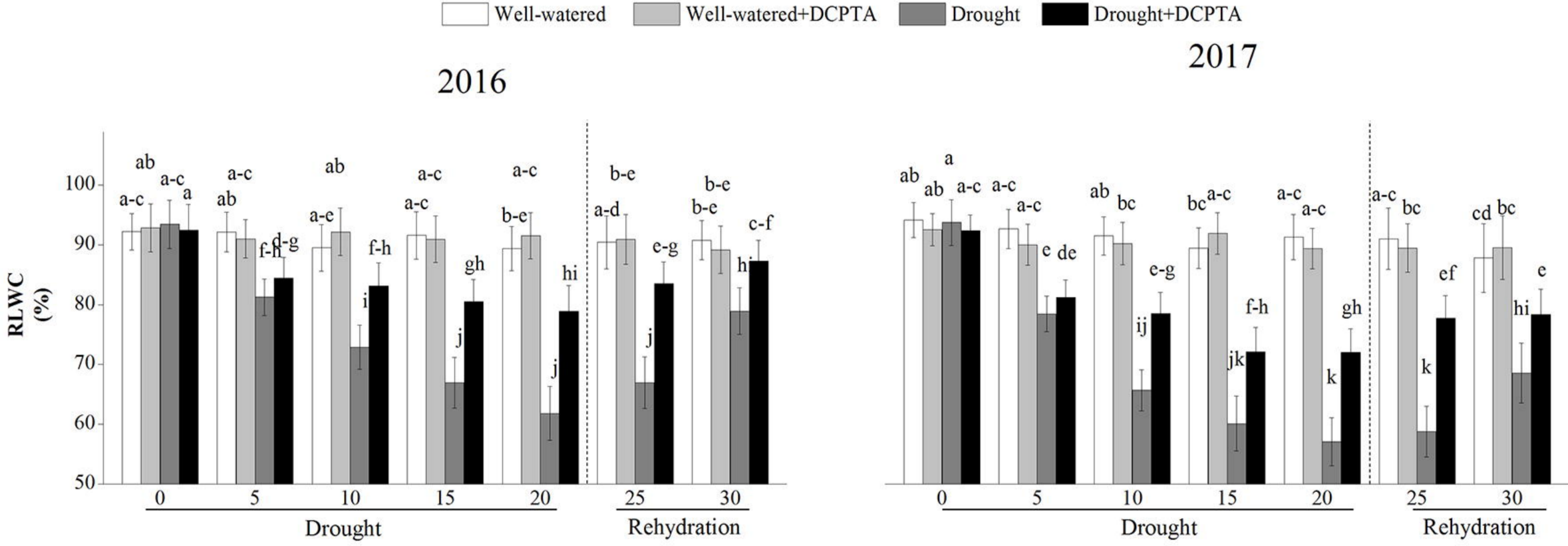


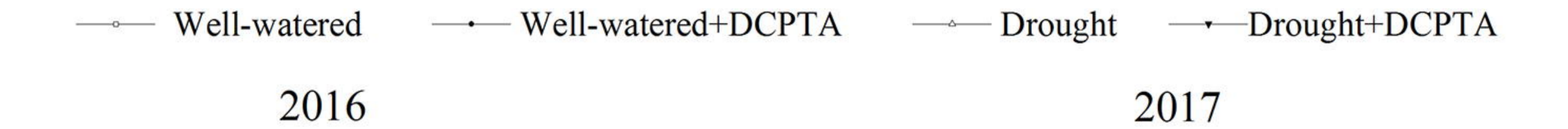


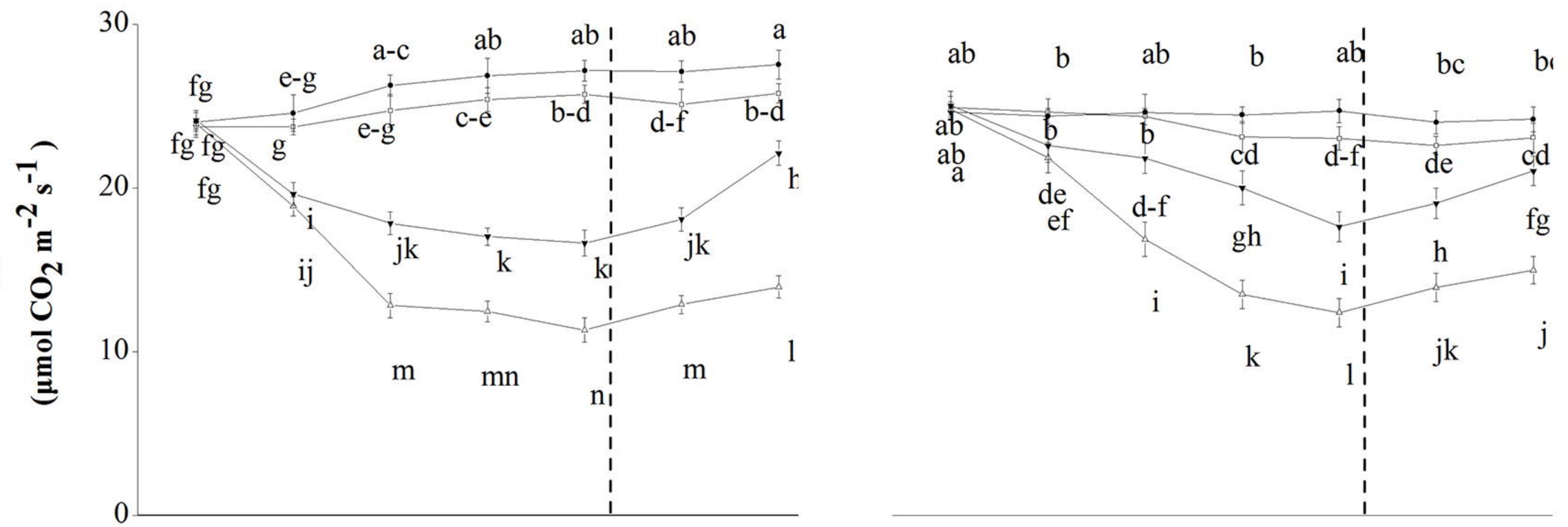


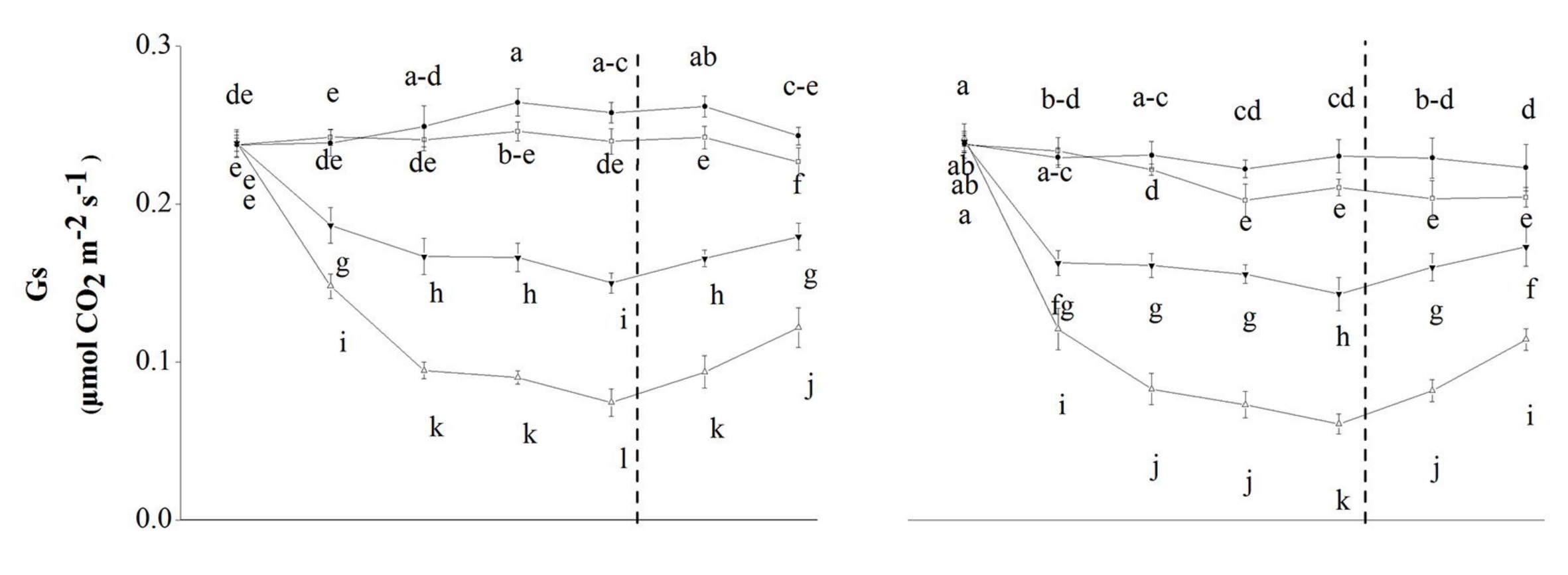


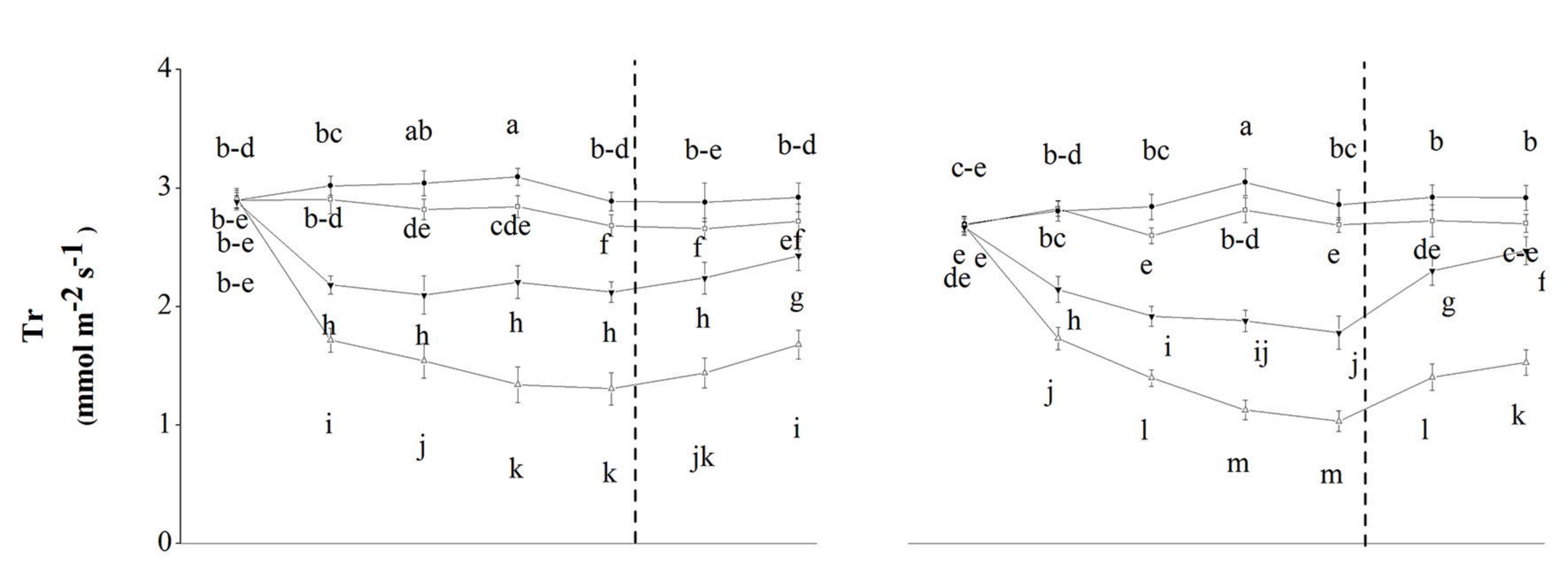
Days after treatment



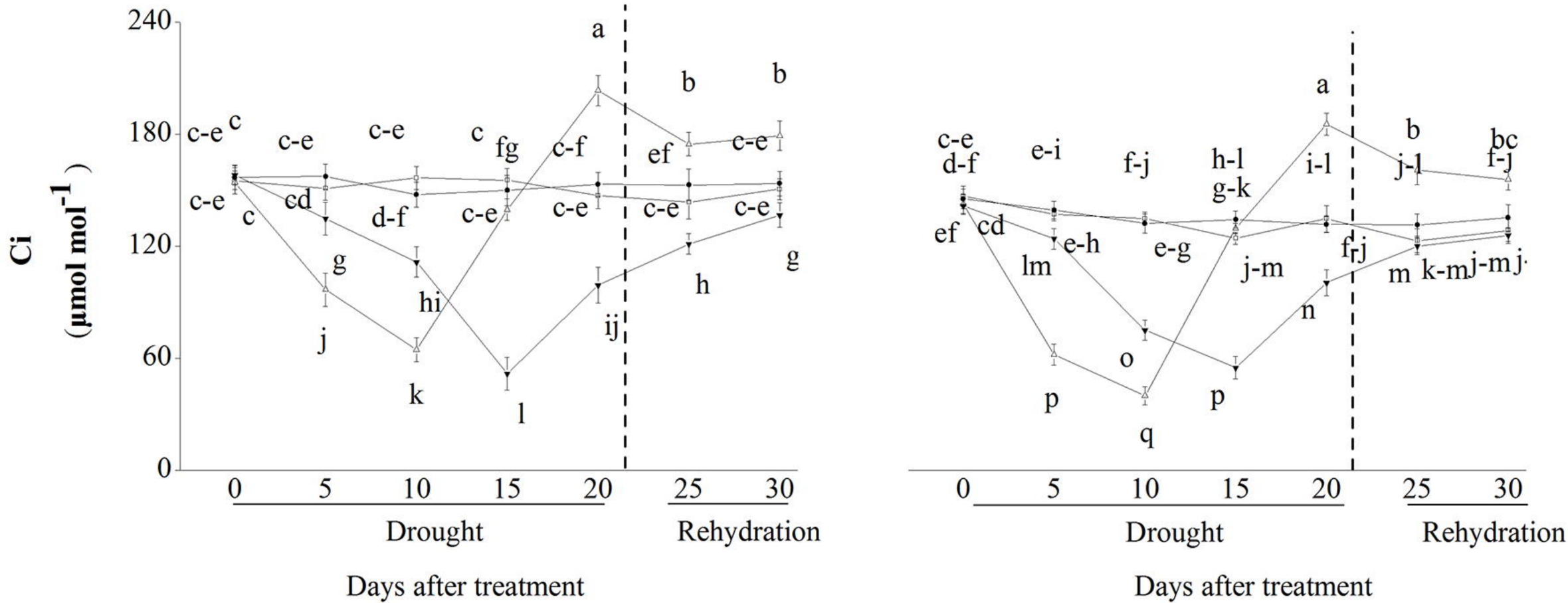


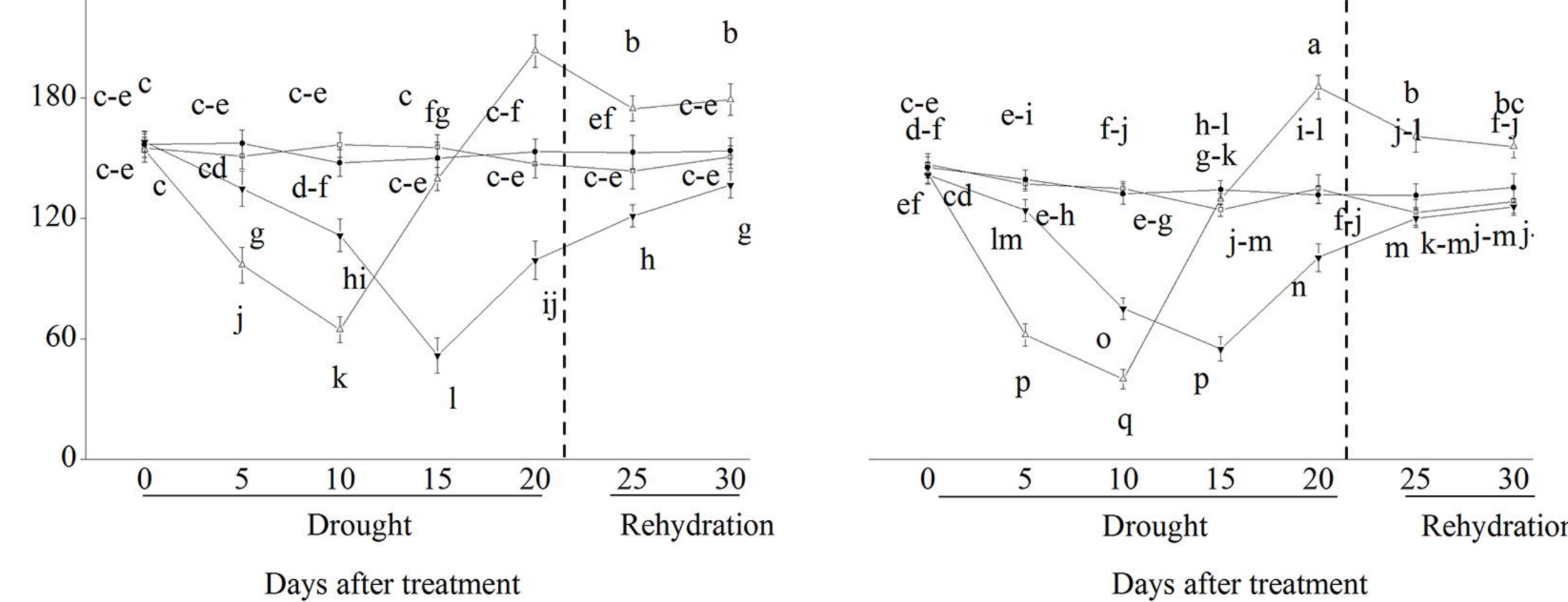




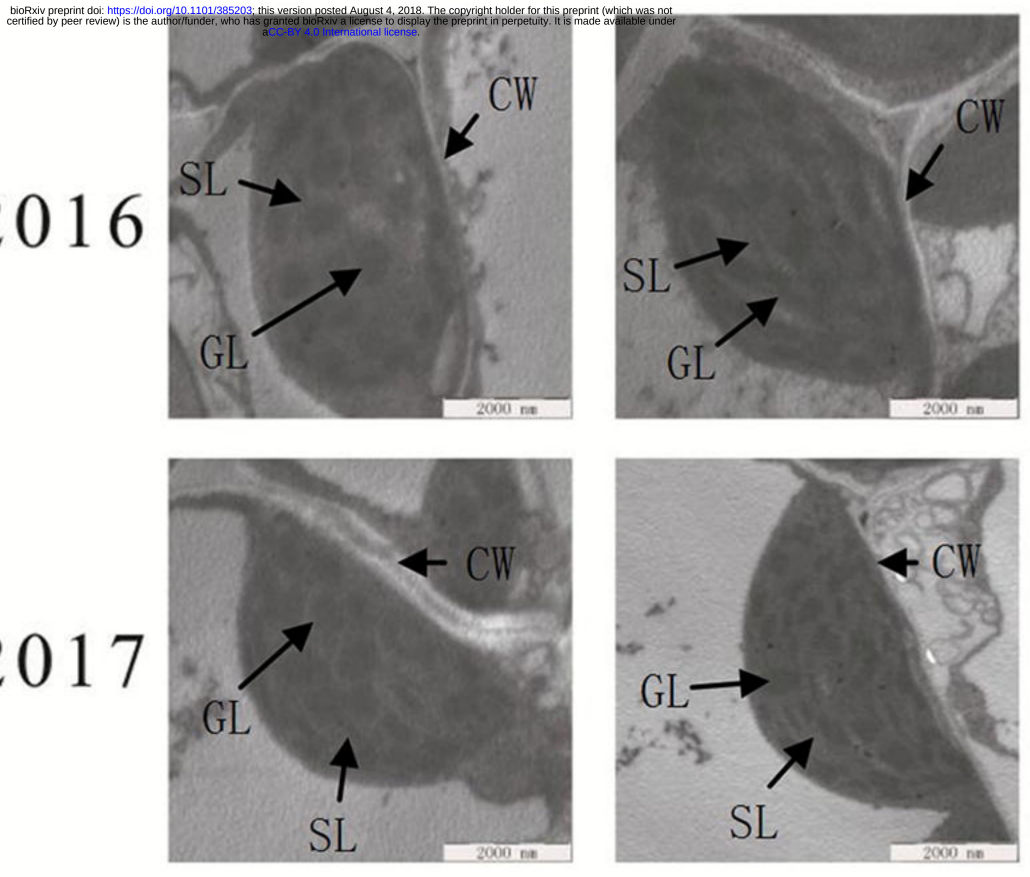


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2017

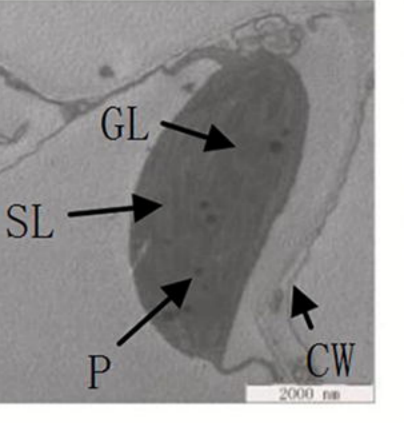


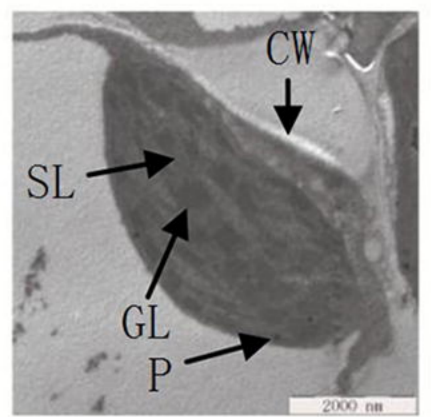
Well-watered

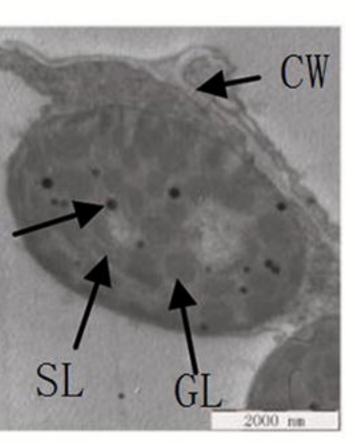
#### Well-watered +DCPTA

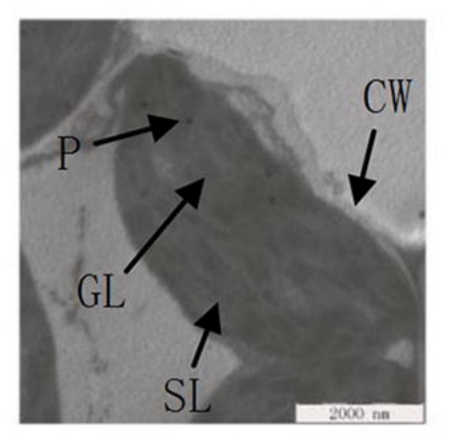
#### Drought +DCPTA

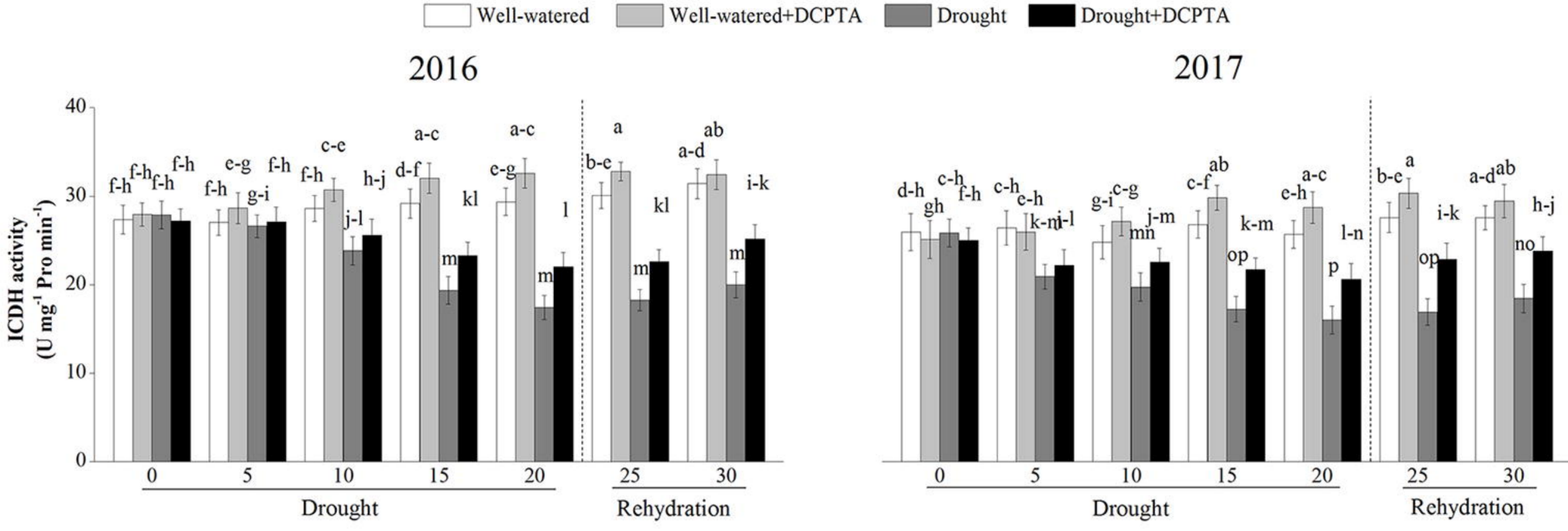
#### Drought







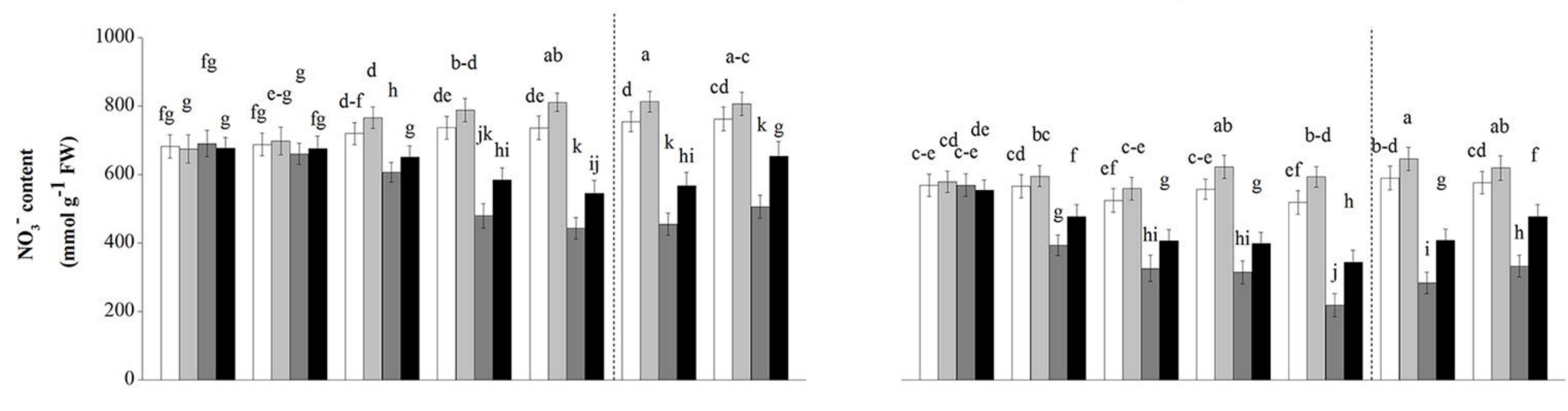


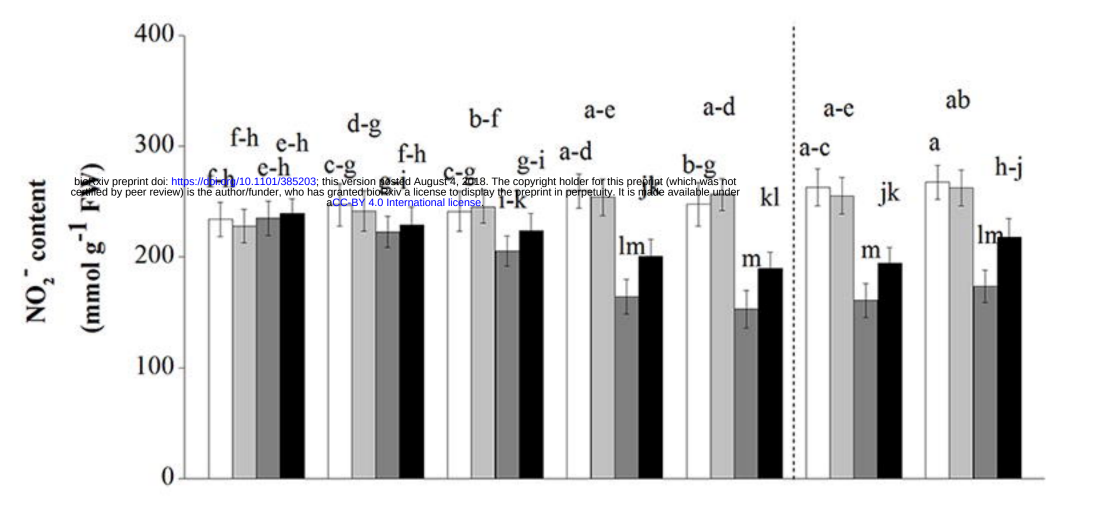


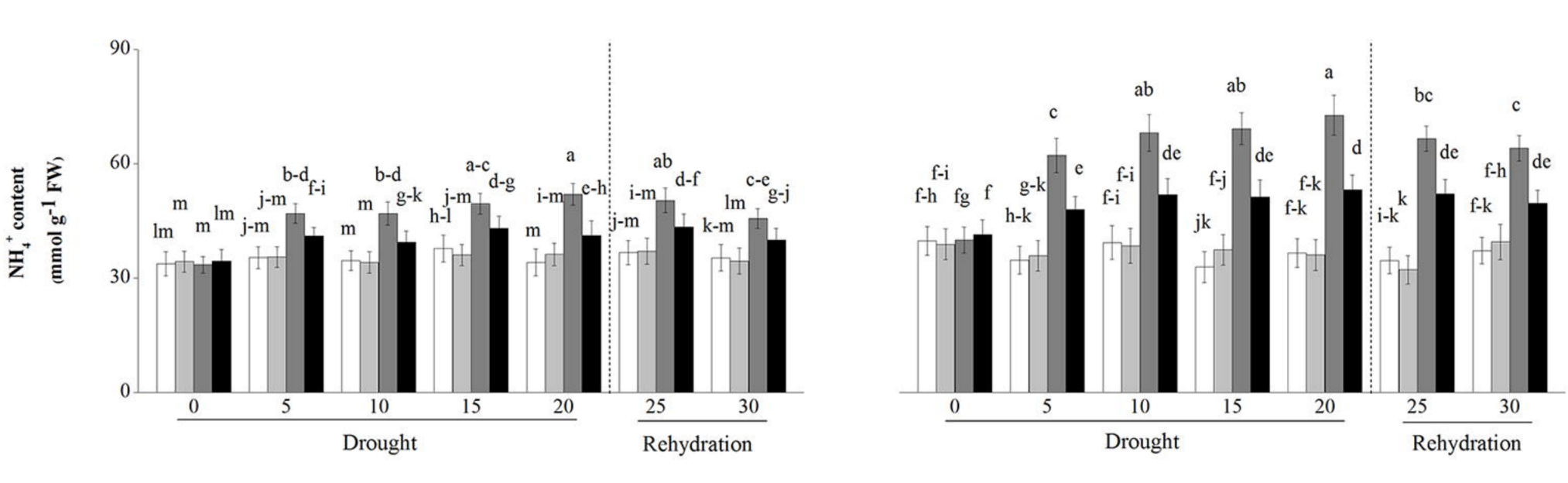
Days after treatment







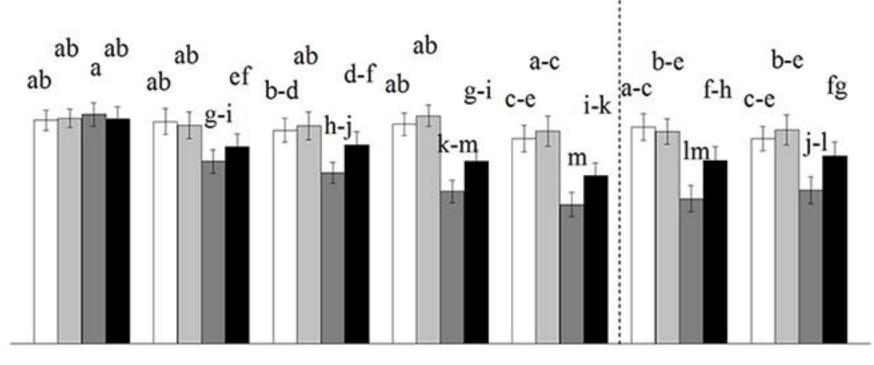




Drought

Drought+DCPTA

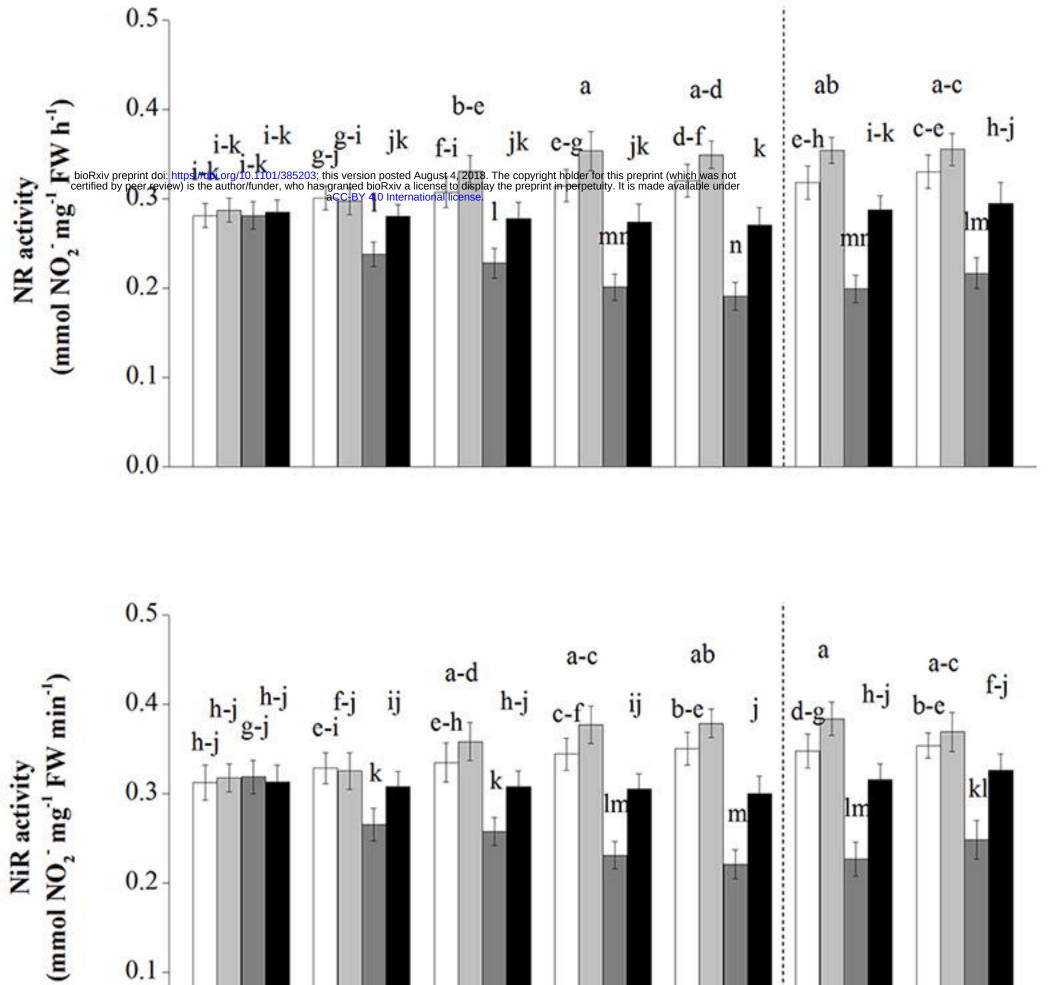
2017

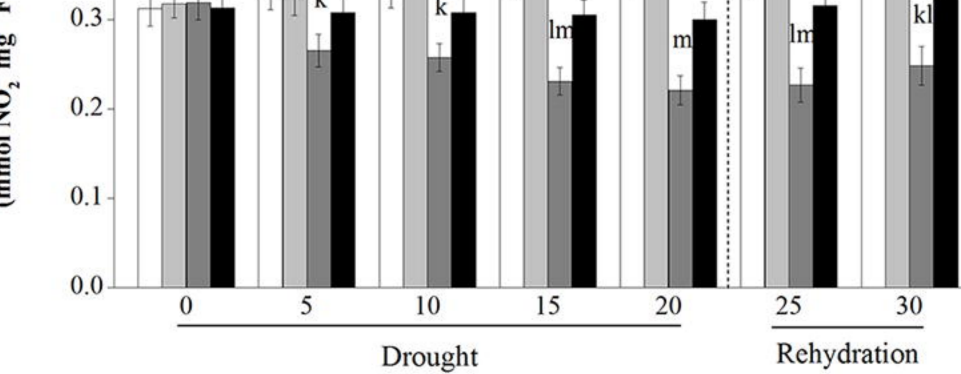


Well-watered

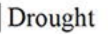
Well-watered+DCPTA





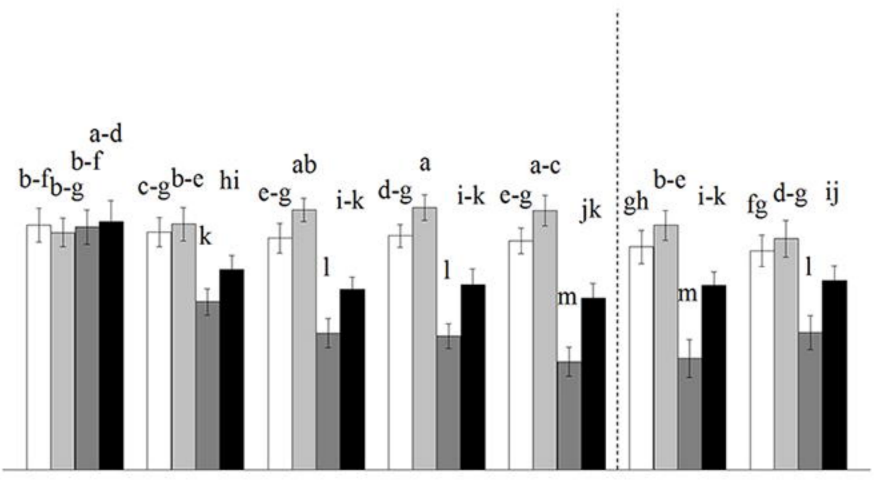


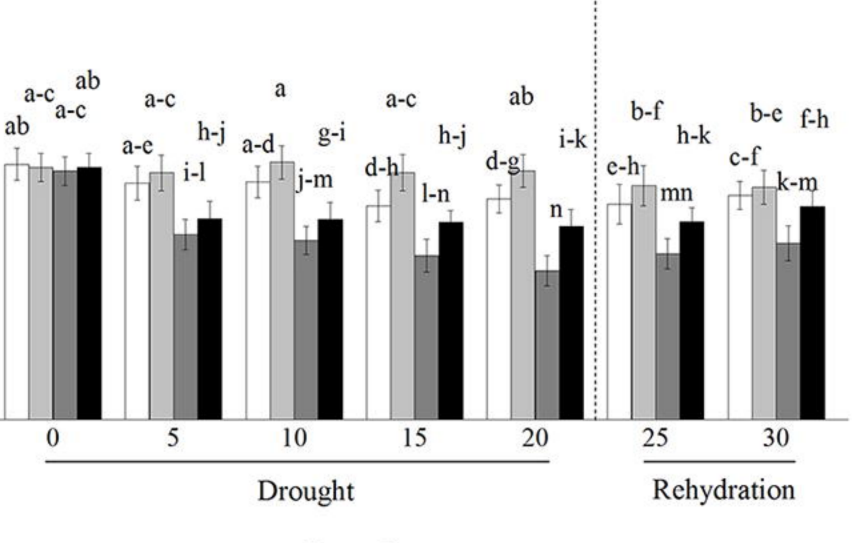
Days after treatment



Drought+DCPTA

#### 2017



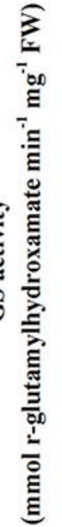




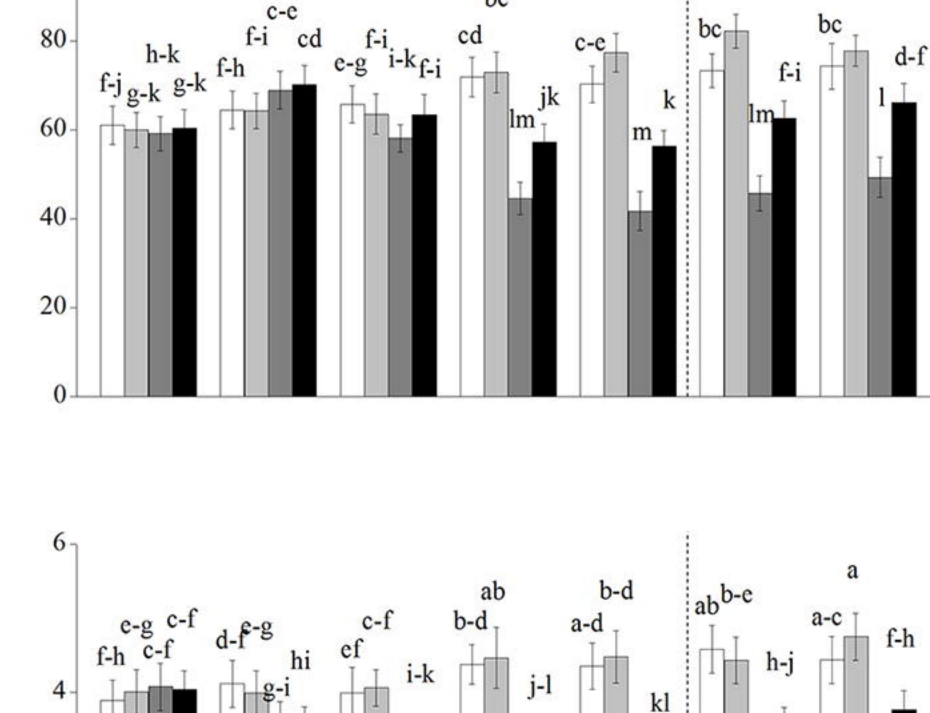
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# GS activity



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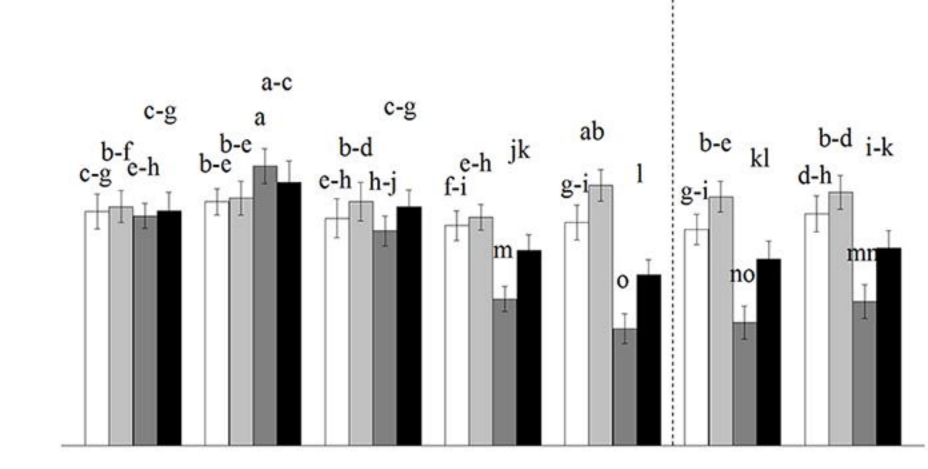
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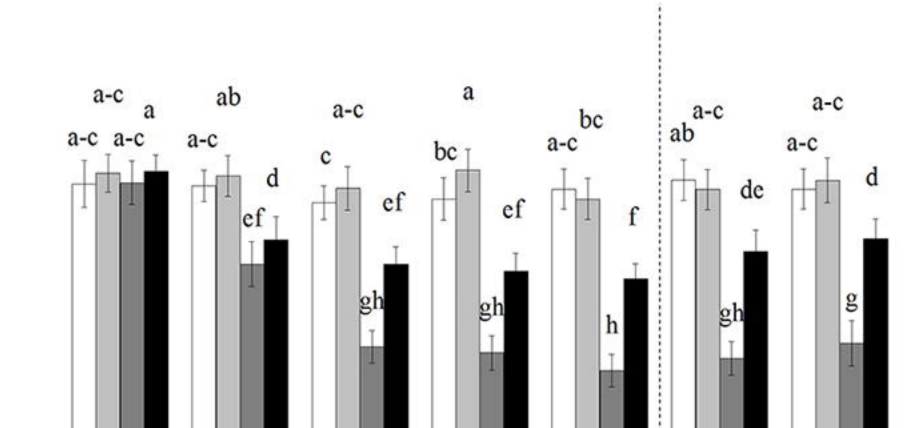
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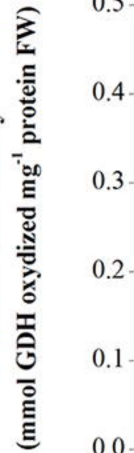


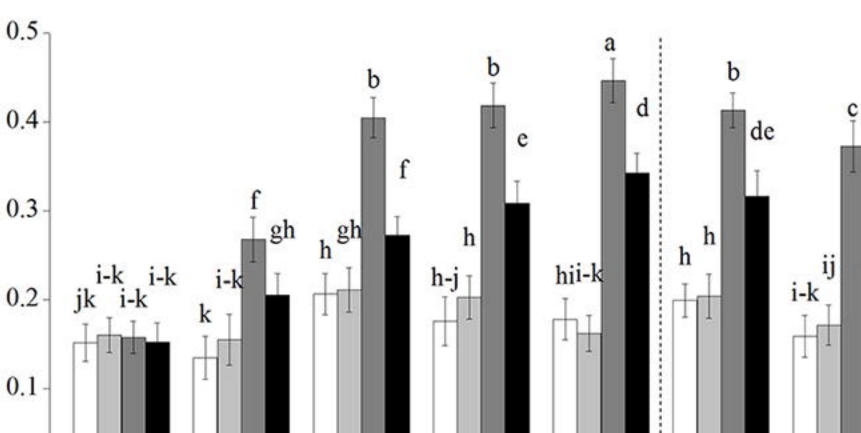


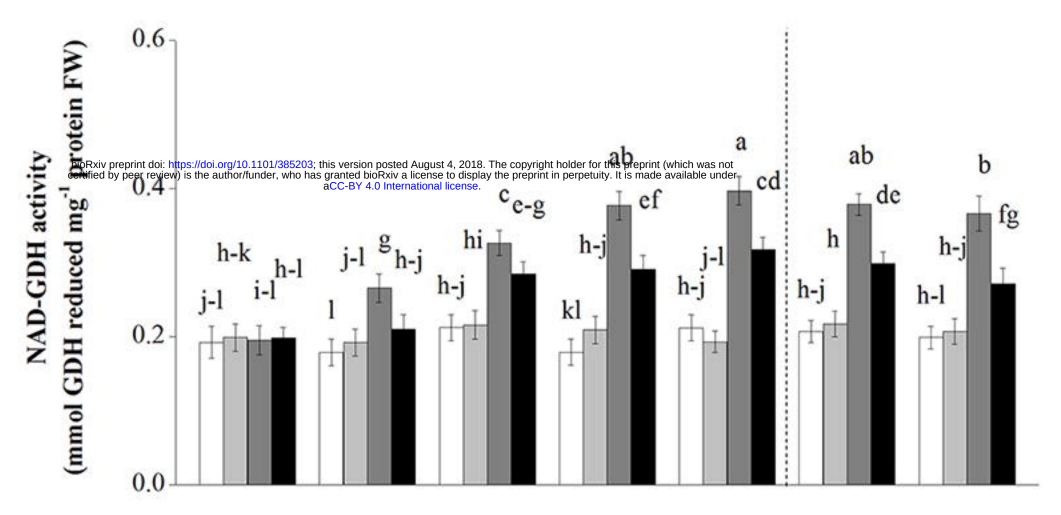


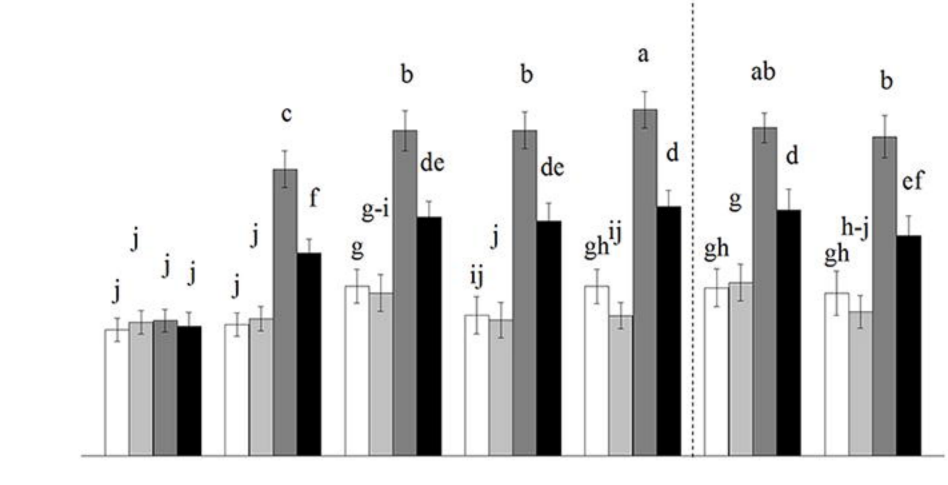


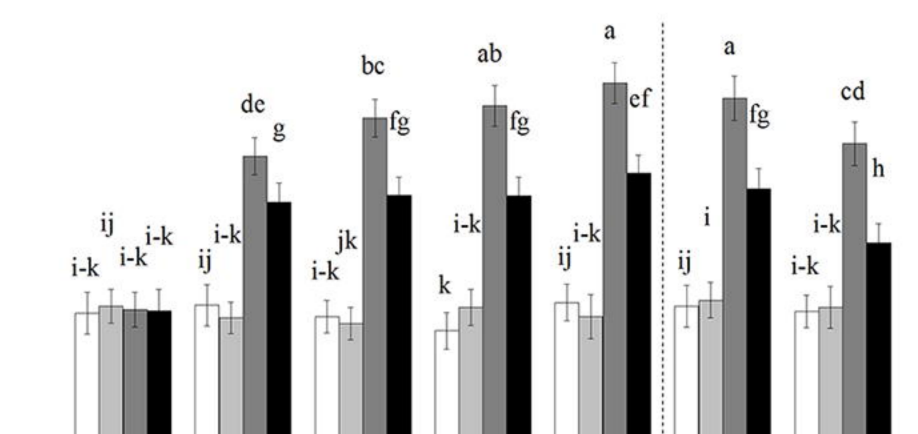
# NADH-GDH activity

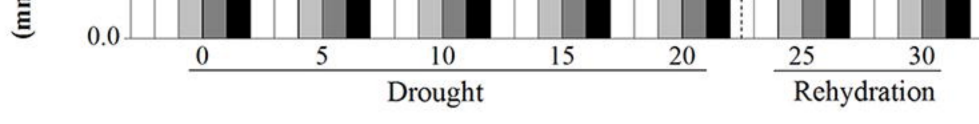


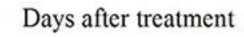


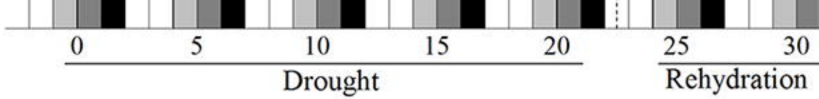


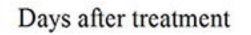


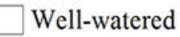






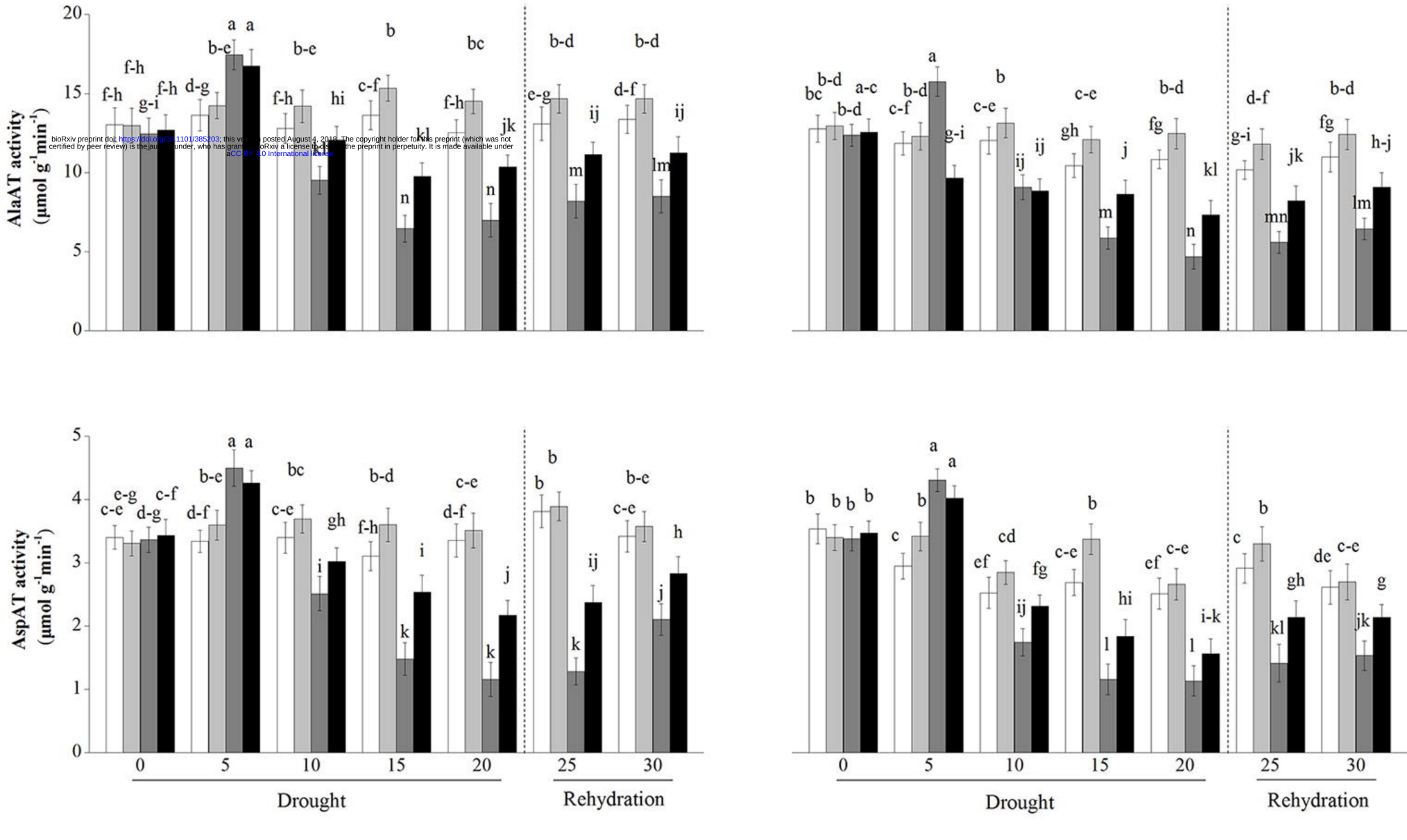




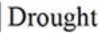


Well-watered+DCPTA

2016

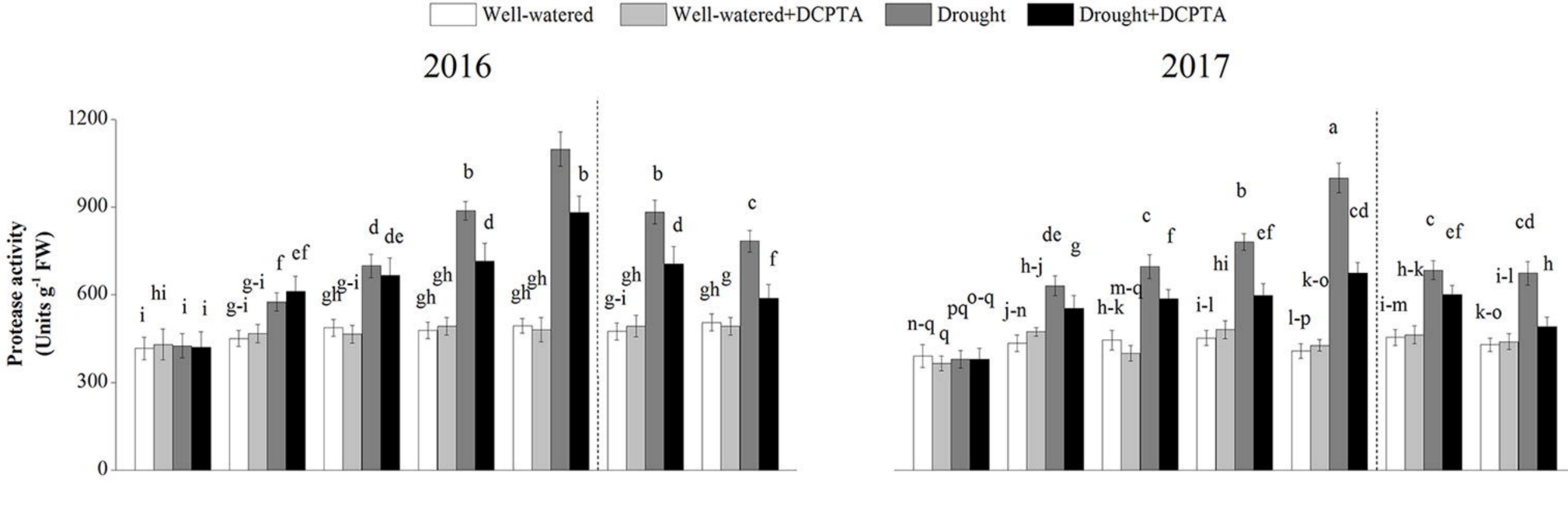


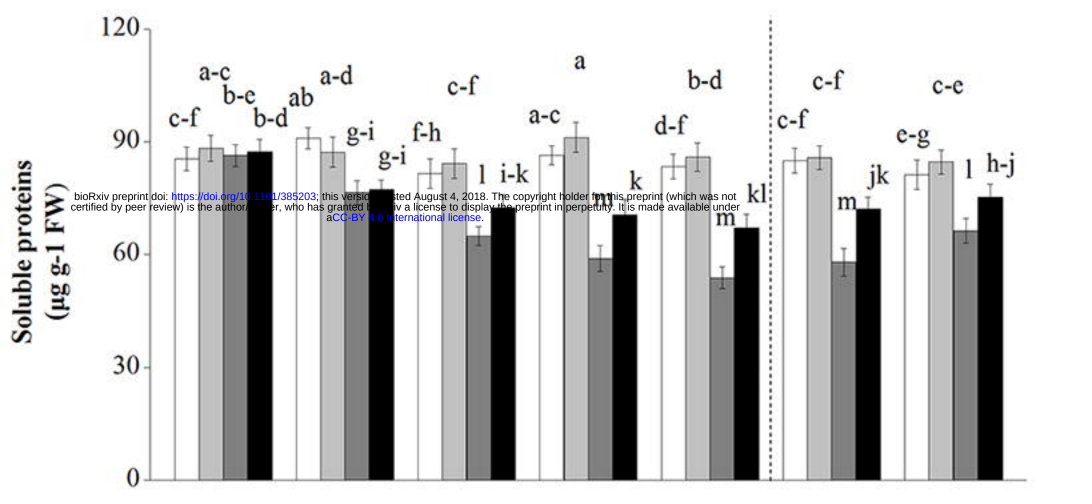
Days after treatment

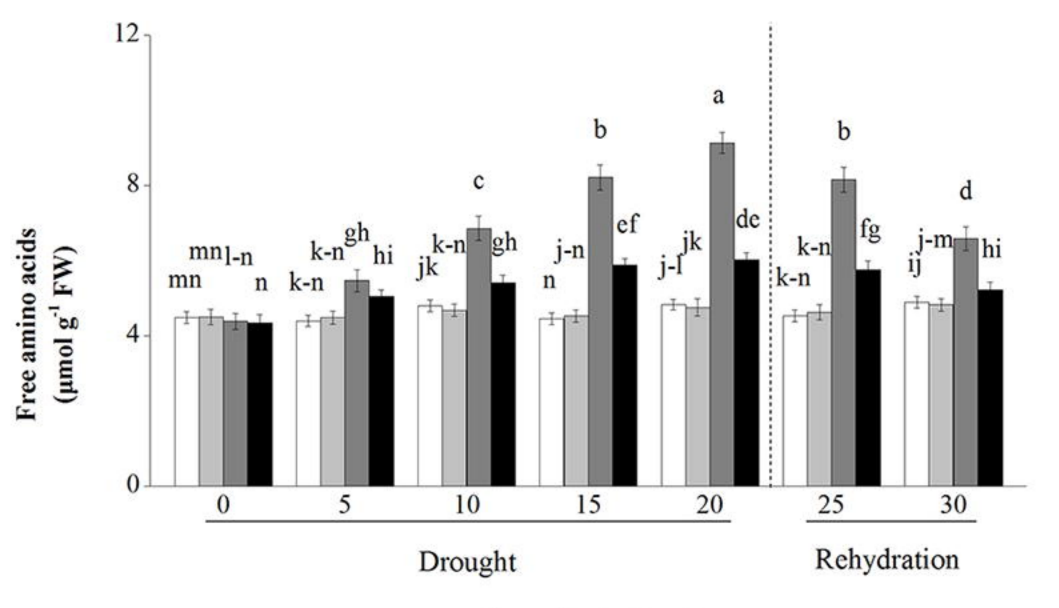


Drought+DCPTA

2017

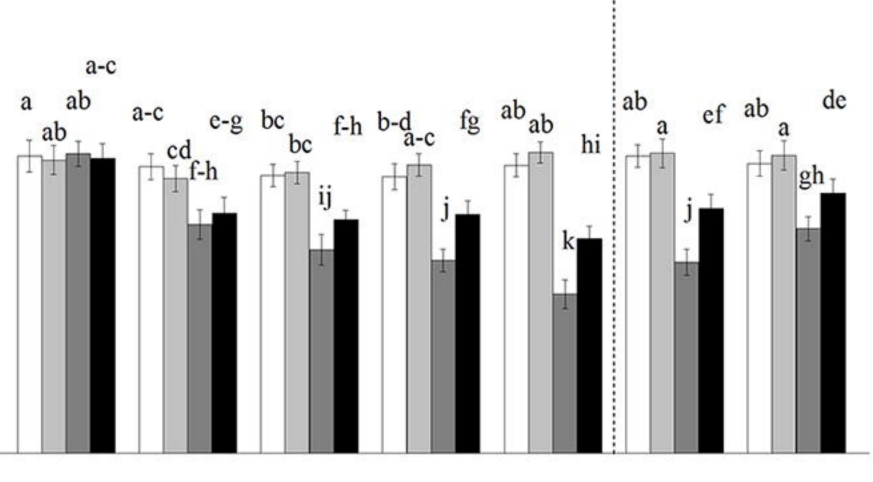


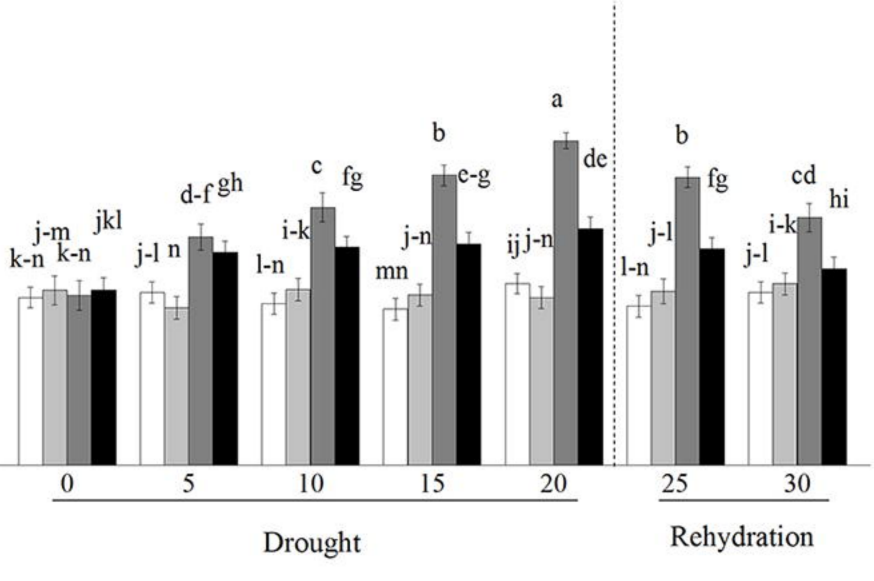




Drought

Drought+DCPTA





Days after treatment