ACCEPTED ARTICLE 1 Alleviation of drought stress in German Chamomile (Matricaria chamomilla L.) in 2 3 response to suppressive oxidative stress and water deficit-induced stomatal closure by 4 exogenous polyamines 5 6 Mohammad Javad Ahmadi-Lahijani^{1*}, Jafar Nabati², Saeed Moori³, Mohammad Kafi¹ 7 8 ¹ Department of Agrotechnology, Ferdowsi University of Mashhad, Iran. 9 ² Research Center for Plant Sciences, Ferdowsi University of Mashhad, Iran.

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Running head: Polyamines alleviates drought stress in German Chamomile ABSTRACT

18 Polyamines (PAs) are signaling molecules that exhibit promising roles in improving stress 19 tolerance in plants. Limited information is available concerning the effects of the exogenous PAs on medicinal plants including chamomile. This experiment was carried out to study the 20 21 effects of foliar application of PAs [putrescine (Put), spermidine (Spd), and spermine (Spm)] 22 on physiological and biochemical processes to understand the possible mechanisms concerning the water deficit stress [soil field capacity (FC) as control, 80% of FC (FC₈₀), and 60% of FC 23 24 (FC₆₀)] alleviation in German Chamomile. We found that PAs partially inhibited water deficitinduced stomatal closure and induced antioxidant enzymes to eliminate the increased H_2O_2 . 25 Spd increased stomatal conductance (g_s) by 66, 65, and 35% at FC, FC₈₀, and FC₆₀, respectively, 26 27 compared with the control. The increased g_s enhanced leaf net photosynthesis (A_N) by 52 and 28 86% at FC_{80} and FC_{60} , respectively, compared with the control. The role of PAs in oxidative damage alleviation was approved by the negative correlation of leaf antioxidant activities and 29 30 malondialdehyde (MDA) and H_2O_2 content. According to the results, PAs function as stress-31 protecting compounds to instigate the antioxidative enzymes to scavenge stress-induced H_2O_2 , 32 improve membrane stability, and enhance water deficit tolerance. Generally, our results suggested that PAs could be potential growth regulators to alleviate mild to severe water deficit 33 34 stress.

Keywords: Chlorophyll fluorescence; enzymatic antioxidant; hydrogen peroxide; gas exchange variables; non-photochemical quenching.

INTRODUCTION

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39 Drought is considered the most crucial worldwide factor in plant production systems (Hafezi 40 Ghehestani *et al.*, 2021). Water deficit stress affects the metabolism and growth of plants, 41 agricultural ecosystems, and human societies (Tezara et al., 1999). Various physiological and 42 metabolic responses such as stomatal closure and decline in photosynthesis and growth rate are 43 induced in plants under water deficiency (Flexas and Medrano, 2002). Plant response to 44 stressful conditions is initiated when the stress is recognized at the cellular level. In both 45 unstressful and stressful environments, plants produce reactive oxygen species (ROS) that react with proteins, lipids, and DNA and impair normal cellular functions (Apel and Hirt, 2004). 46 47 Water deficit stress disturbs the balance between ROS production and scavenging in plants, leading to the accumulation of ROS and accelerating cell membrane damage and lipid 48 49 peroxidation (Farooq et al., 2009).

Polyamines (PAs) are classified as a group of phytohormone-like aliphatic amine natural 50 51 compounds with aliphatic nitrogen structure and are considered secondary messengers in 52 signaling pathways (Liu et al., 2023). Generally, naturally occurring PAs in the higher plants 53 including putrescine (Put), spermidine (Spd), and spermine (Spm) are not only involved in 54 numerous cellular and molecular processes in plants but also have been shown to improve plant 55 tolerance to abiotic stresses (Baghalian et al., 2011). PAs trigger several molecular, 56 biochemical, and physiological responses of plants including increasing membrane stability and 57 osmolyte accumulation, protection of photosynthetic apparatus, activation of antioxidant machinery, regulation of redox homeostasis, upregulation of stress-related genes, and 58 59 promotion of plant stress tolerance (Alcázar et al., 2020).

60 The interaction of PAs with membrane phospholipids induces membrane stability under stressful conditions. PAs play a vital role as signaling molecules that regulate several metabolic 61 62 pathways. The abiotic stress adaptations are enhanced by the PAs' functions as stress signaling 63 molecules (Pál et al., 2015). Exogenously applied PAs increased antioxidant enzyme activities 64 under various stressful conditions; which could reduce cell damage and enhance the stress tolerance of plants (Hassan et al., 2018; Alcázar et al., 2020). The accumulation of PAs under 65 adverse conditions can directly act as an antioxidant in eliminating ROS or may activate the 66 **ROS-scavenging enzyme system (Liu** *et al.***, 2023).** Previous studies showed that exogenous 67 68 PAs significantly increased the activity of antioxidants such as SOD, POD, and CAT and decreased ROS synthesis in Vicia faba, Citrus reticulata, Arabidopsis thaliana, and Rosa 69 70 damascene (Hasan et al., 2021; Liu et al., 2023).

71 German chamomile (Matricaria chamomilla) is one of the most valuable medicinal plants of 72 the Asteraceae (Compositae) family with many applications in the pharmaceutical, nutritional, 73 and cosmetic industries. Chamomile is relatively adaptable to a wide range of climates including arid and semi-arid regions (Das et al., 1998). However, drought negatively affects 74 75 chamomile performance and productivity. Although studies of PAs have been performed on 76 various crops (Alcázar et al., 2006; Farooq et al., 2009; Liu et al., 2023), the available 77 information concerning the effects of PAs on medicinal plants is still limited. The effect of foliar application of polyamines on leaf gas exchanges, chlorophyll fluorescence, and 78 79 physiological and biochemical processes was studied to understand the possible mechanisms 80 concerning water stress alleviation in German chamomile.

81 MATERIALS AND METHODS

82 Experimental site and procedure

83 The experiment was carried out at the Greenhouse of the Research Center for Plant Sciences, 84 Ferdowsi University of Mashhad, in 2021. Chamomile seeds (cv. Presov, obtained from Isfahan 85 Natural Resources Research Center) were surface sterilized with 0.2% sodium hypochlorite 86 solution for 5 min and rinsed three times with tap water. Ten chamomile seeds were sown in 87 each 10 kg pot (20 and 25 cm in diameter and depth, respectively) filled with clay loam soil at 88 a depth of 1 cm and thinned to 5 plants per pot after establishment. Plants were grown under 89 greenhouse conditions with day/night temperatures of 20/15±2 °C, natural light (~800 µmol.m⁻ 1 .s⁻¹ PPFD), photoperiod of 13/11 h day/night, and relative humidity of 50±10%. 90

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Soil preparation and treatments

Irrigation was applied until the full establishment (4-leaf), and then, water deficit was applied 93 94 at three levels of control (FC₁₀₀), moderate stress (80% of FC; FC₈₀), and severe stress (60% of 95 FC; FC_{60}) according to the method of Topp and Davis (1985). Plants were fertilized with the 96 Hoagland nutrient solution once a week along with irrigation water. Polyamines were foliar 97 applied as (a) spermine [Spm], (b) spermidine [Spd], (c) putrescine [Put], and (d) control, at a concentration of 10 µM (Farooq et al., 2009; Ali et al., 2009). 10 mL of the solution was applied 98 99 to each plant using a handheld sprinkler. The first spray was made at the 5-leaf stage and 100 repeated at 15-day intervals until the flowering onset. The control plants were sprayed with 101 distilled water. The experiments were carried out in three replicates. All physiological and 102 biochemical data were taken from fully expanded leaves at the middle of the flowering stage. 103

104 Gas exchange parameters

Net photosynthetic rate (A_N), intercellular CO₂ concentration (C_i), transpiration rate (T_r), and stomatal conductance (g_s) were measured between 9:00–11:00 h using a portable photosynthetic system (ADC Bio Scientific Ltd, UK). Photosynthetically active radiations (PAR), air temperature, relative humidity, and CO₂ concentration inside the sensor head were set at 800 µmol.m⁻².s⁻¹, 25±2 °C, 50±5%, and 400±20 ppm, respectively. Instantaneous (WUE_i) and intrinsic (A_N/g_s) water use efficiency were calculated by dividing A_N by T_r and g_s , respectively. Mesophyll conductance (g_m) was also calculated as A_N/C_i (Fischer *et al.*, 1998).

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116 Chlorophyll fluorescence (Chf)

A portable fluorometer (PAM-2500, Walz, Effeltrich, Germany) was used to measure the dark-117 118 and light-adapted leaf chlorophyll fluorescence between 10:00-12:00 h. After 30 min of dark adaptation, Fv/Fm was calculated as (Fm-Fo)/Fm, where Fm and Fo were the maximum 119 120 fluorescence elicited by a saturating light pulse and steady-state chlorophyll fluorescence, 121 respectively (Genty et al., 1989). The maximum (Fm') and the steady-state (Fs) fluorescence 122 signals were measured from the light-adapted leaves after 4 min of illumination with 123 continuous red wavelength, non-saturating actinic light, and saturating pulses every 25 sec 124 (Murchie and Lawson, 2013). To measure the minimal fluorescence after the PSI excitation 125 (Fo'), the actinic light was then turned off, and far-red pulses were applied. Photochemical 126 quenching (qP) was calculated as $(F_m'-F_s)/(F_m'-F_o')$. Non-photochemical quenching, NPQ, which is a proportion of the rate of the thermal energy dissipation, was estimated as (F_m-127 F_m')/F_m' (Van Kooten and Snel, 1990). 128

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Electrolyte leakage (EL)

Leaf EL was measured to determine leaf membrane damage using an electrical conductivity
(EC) meter (Jenway Model 4510) according to Eq. 1 (Lutts *et al.*, 2016):

$$EL(\%) = \frac{EC1}{EC2} \times 100 \tag{1}$$

Here, EC₁ and EC₂ are the EC of the solution after 24 h and the autoclaved (120 °C for 20 min)
samples, respectively.

137 Relative water content (RWC)

138 Leaf RWC was estimated using Eq. 2 (Smart and Bingham, 1974):

139	$RWC (\%) = \left[\frac{Wf - Wd}{Wt - Wd}\right] \times 100 $ (2)
140	LWt - Wa Here, W _f , W _t , and W _d are fresh weight, turgid weight, and oven-dried weight (at 70 °C until
141	constant mass), respectively.
142	
143	Leaf osmotic potential (<mark>#0</mark>)
144	Leaf $\frac{1}{2}$ was determined according to the freezing point depression method using an osmometer
145	(Wogel, model OM802.D). The leaf osmolytes content was calculated based on the van't Hoff
146	equation, and the leaf water content was measured by Eq. (3):
147	$\frac{mMol}{g} = \left[\left(-\frac{Op}{RT} \right) \times \left(\frac{WC}{1 - WC} \right) \right] $ (3)
148	Where the osmolytes content is based on mM g ⁻¹ dry weight, R is the universal gas constant
149	(0.00831-liter MPa mol ⁻¹ °K ⁻¹), T is the absolute temperature (273 °K), Op is the leaf osmotic
150	potential (MPa), and WC is the leaf water content". The solute potential was determined at
151	room temperature (25 °C).
152 153	Photosynthetic pigments
154	Fresh leaves (100 mg) were homogenized in ethanol 70% and kept at 4 °C for 24 h. Leaf
155	pigments content (Chlorophylls a, b, and carotenoids) were determined spectrophotometrically
156	(U-2000, Hitachi Instruments, Tokyo, Japan) according to Lichthentaler and Wellburn (1983).
157 158	Loof ontiovidant ongrange
158 159	Leaf antioxidant enzymes 100 mg leaf fresh weight was ground in liquid nitrogen, and 1 ml potassium phosphate (0.1 M,
160	pH = 7.8) containing 1 mM EDTA was added. The insoluble solids were removed by
161	centrifuging the mixture in 12,000 g at 4 °C (Sigma, model K18-3). The supernatant was kept
162	at -80° C to assay the enzymatic antioxidant activities (Yamaguchi <i>et al.</i> , 1995). Leaf enzymatic
163	antioxidants activity, including Ascorbate peroxidase (APX, EC 1.11.1.11), Superoxide
164	dismutase (SOD, EC 1.15.1.1), Catalase (CAT, EC 1.11.1.6), and Peroxidase (POD, EC
165	1.11.1.7), were assayed by the methods described by Nakano and Asada (1981), Giannopolitis
166	and Ries (1977), Cakmak and Horst (1991), and Ghanati <i>et al.</i> (2002), respectively.
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168	Malondialdehyde (MDA) and H2O2 content
169	One hundred mg of leaf fresh weight was used to measure leaf MDA by the methods described
170	by Jiang and Zhang (2001). Leaf MDA was measured by homogenizing leaf fresh weight in 5
171	ml of trichloroacetic acid (100 g ⁻¹) containing 250 g l ⁻¹ thiobarbituric acid. The supernatant

- absorbance was read at 532 nm spectrophotometrically (Jenway UV-Visible, Model 6305) and
- 173 was corrected at A600. For H_2O_2 content measurement, leaf tissues (500 mg) were 174 homogenized in an ice bath with 5 ml 0.1% (w:v) TCA. The homogenate was centrifuged at
- 175 12000×g for 15 min and the supernatant absorbance was read at 390 nm. The content of H_2O_2
- 176 was given on a standard curve (Sergiev *et al.*, 1997).
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178 Statistical analysis

The experiment was carried out as a factorial (3 levels of water deficit \times 4 levels of PAs) arrangement in a randomized complete block design with three replications. The experiment was carried out twice and the pooled data were analyzed. Data were subjected to a two-way analysis of variance, and the LSD p \leq 0.05 was the test criterion for assessing differences between the means of the main and/or interaction effects using SAS *v*.9.4 software. Data presented as ±SE.

185

186 **RESULTS**

187 **Photosynthetic parameters**

188 Although the gas exchange parameters were reduced by the water deficit, they were 189 significantly improved by the application of PAs. Water deficit at FC₆₀ diminished the untreated 190 plant A_N by 40% compared with FC₁₀₀ (Fig. 1A). PAs application increased A_N by ~60% 191 compared with the untreated plants under FC_{100} (Fig. 2A). However, under FC_{80} and FC_{60} , Spd 192 showed the greatest improving effect on A_N; Spd application enhanced leaf A_N by 52 and 86% 193 at FC₈₀ and FC₆₀, respectively, compared with the untreated plants (Fig. 1A). Chamomile leaf 194 T_r reduced by 1.1 and 1.6 times, respectively, at FC₈₀ and FC₆₀ compared with FC₁₀₀ (Fig. 1B). 195 The highest leaf T_r was observed in Spd-treated plants at FC₁₀₀; 41, 17, and 21%, respectively, 196 higher than the untreated, Put, and Spm-treated plants (Fig. 1B).

197 Foliar application of PAs reduced the diminishing effects of water deficit on g_s. Spd increased 198 leaf g_s by 66, 65, and 35% at FC₁₀₀, FC₈₀, and FC₆₀, respectively, compared with the untreated 199 plants (Fig. 1C). The lowest leaf gm was observed when Spd and Spm were applied, 200 respectively, at FC₁₀₀ and FC₈₀ (Fig. 1D). Although water deficit decreased C_i and C_i:C_a, PAs 201 significantly increased Ci and Ci:Ca compared with the untreated plants (Fig. 3A and B). At 202 FC_{100} and FC_{80} , C_i was the highest in the Spd-treated plants by 1.4 and 1.1 times higher than 203 the untreated plants, respectively. The highest WUE_i was observed in the untreated plants at 204 FC₈₀ (Fig. 3C). Spm-treated plants showed 43% higher WUE_i at FC₁₀₀ compared with the 205 untreated plants; however, WUE_i reduced when PAs were applied under water deficit 206 conditions. A_N/g_s decreased by reducing the soil moisture. The highest A_N/g_s was observed in 207 Spm and Spd-treated plants at FC₁₀₀ and FC₆₀, respectively; 24 and 38% higher than the 208 untreated plants (Fig. 3D).

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210 Leaf chlorophyll fluorescence (Chf)

211 Water deficit at the level of FC₆₀ decreased F_v/F_m by 22% compared with FC₁₀₀ (Fig. 4A). At 212 FC_{80} , Put-treated plants showed the highest F_v/F_m , which was 13% higher than the untreated 213 plants. Non-photochemical quenching (NPQ) increased by increasing the water deficit intensity 214 (Fig. 4B). Spd-treated plants showed 85 and 65% lower leaf NPQ than the untreated plants, respectively, at FC₈₀ and FC₆₀. Photochemical quenching (qP) reduced by 26 and 33% at FC₈₀ 215 216 and FC₆₀, respectively, compared with FC₁₀₀ (Fig. 4C). The highest qP was observed in the Put-217 treated plants; 36, 69, and 34% higher than the untreated plants, respectively, at FC₁₀₀, FC₈₀, 218 and FC₆₀. The linear electron transport rate, ETR, decreased by 24 and 48%, respectively, at 219 FC₈₀ and FC₆₀ compared with FC₁₀₀ (Fig. 4D). The greatest ETR was recorded in the Spm-220 followed by Put-treated plants at FC₈₀ by on average $\sim 25\%$ higher than the untreated plants 221 (Fig. 4D and 2A). However, at FC₆₀, Spd increased ETR by 39% compared with the untreated 222 plants.

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224 Leaf RWC, *wo*, and EL

225 Water deficit at FC₈₀ and FC₆₀ reduced leaf RWC by 13 and 22%, respectively, compared with 226 FC₁₀₀ (Fig. 5A). Put, Spm, and Spd enhanced leaf RWC by an average of ~25% compared with 227 the untreated plants at FC₁₀₀ (Fig. 5A and 2A). Water deficit at the level of FC₆₀ reduced $\frac{\psi_0}{\psi_0}$ by 228 45% compared with FC₁₀₀ (Fig. 5B). In contrast, leaf ψ_0 was enhanced in the PAs-treated 229 plants. Put application increased leaf ψ_0 35% compared with the untreated plants at FC₆₀ (Fig. 230 5B). Leaf EL was increased by 1.6 and 2.8 times at FC_{80} and FC_{60} , respectively, compared with 231 FC₁₀₀ (Fig. 5C). However, Spm and Spd decreased leaf EL by 17 and 28%, respectively, 232 compared with untreated plants at FC_{60} (Fig. 5C).

234 Leaf photosynthetic pigment

A significant decrease was observed in leaf photosynthetic pigments content exposed to water deficit (Table 1). However, Spd increased leaf Chlt by 51, 60, and 79%, respectively, compared with the untreated plants at FC₁₀₀, FC₈₀, and FC₆₀ (Table 1). The highest Chl a:b was observed in the put-treated plants at FC₈₀; 32, 13, and 18% higher than the untreated, Spm-, and Spdtreated plants, respectively. Water deficit increased leaf carotenoid content on average by ~18% 240 compared with FC_{100} (Fig. 2B). The highest leaf carotenoid content was observed in Spd-treated 241 plants at all water-deficit levels (Table 1).

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243 Enzymatic antioxidant

244 Leaf enzymatic antioxidant activity was significantly influenced by the water deficit, foliar application of PAs, and their interaction (Fig. 6A). Water deficit and PAs increased leaf 245 246 antioxidant activity. Spd-treated plants showed the highest CAT at FC₈₀ and FC₆₀ than the untreated plants. Spd increased CAT by 82 and 100% compared with the untreated plants at 247 248 FC₈₀ and FC₁₀₀, respectively (Fig. 6A). Leaf POD activity showed an increasing trend by 249 increasing the water deficit intensity and PAs application. At FC₈₀ and FC₆₀, Spd- and Spm-250 treated plants showed the greatest POD, respectively, which were nearly double the untreated 251 plants at the respective water deficit level. At FC₈₀, the highest APX activity was recorded in 252 Spm- followed by Spd-treated plants by an average of ~35% over the untreated plants (Fig. 6C 253 and 2B). Leaf SOD activity showed a similar trend as POD. The highest SOD activity was 254 recorded in Spd- and Spm-treated plants at FC₈₀ and FC₆₀ by 42 and 36%, respectively, over 255 the untreated plants at their respective water deficit levels (Fig. 6D).

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257 Leaf MDA and H₂O₂ content

258 Water deficit increased leaf MDA content; water deficit at FC₈₀ and FC₆₀ increased leaf MDA 259 content by 85% and 1.6 times, respectively, compared with FC₁₀₀ (Fig. 6E). Although PAs-260 treated plant MDA also increased at FC₈₀, it remained almost unaltered at FC₆₀ compared with FC₈₀. Leaf MDA content of Put, Spm, and Spd-treated plants were 38, 34, and 31%, 261 respectively, lower than the untreated plants (Fig. 6E). Leaf H_2O_2 content was almost doubled 262 263 at FC₆₀ compared with FC₁₀₀. However, PAs treatments reduced leaf H_2O_2 content compared with the untreated plants (Fig. 6F). Spd application reduced leaf H₂O₂ by 18 and 10% compared 264 265 with the untreated plants at FC_{80} and FC_{60} , respectively.

267 DISCUSSION

Abiotic stresses such as drought, cold, and K deficiency stresses simultaneously stimulate abscisic acid (ABA) and PAs biosynthesis (Li *et al.*, 2021; Réthoré *et al.*, 2021; Zhu *et al.*, 2020). It has been supposed that PAs and ABA either alone or synergically induce stomatal closure to increase plant tolerance during stressful conditions (Gong *et al.*, 2021). However, most recent findings revealed that the ABA-induced stomatal closure is directly inhibited by PAs in *Vicia faba*, and exogenous applications of PAs could reopen stomata even if they were

- 274 partially closed by ABA treatment (Liu et al., 2023). We observed that exogenously spayed
- 275 PAs stimulated leaf g_s of the water deficit-stressed chamomile plants, resulting in the improved
- leaf A_N, T_r, F_{ν}/F_m , ETR, and qP, and decreased NPQ, which may indicate the alleviating role
- 277 of PAs on drought-induced stomatal closure. Spd or Spm increased g_s and photosynthesis of
- 278 Chinese dwarf cherry but did not affect the F_{ν}/F_m under drought stress (Yin *et al.*, 2014). Those
- 279 observations indicate that PAs can enhance photosynthesis by inhibiting stomatal closure
- 280 without affecting the stability of the photosynthetic system.
- 281 PAs are involved in plant protection against different environmental stresses (Baghalian et al., 282 2011; Farooq et al., 2009). PAs with acid-neutralizing, antioxidative, and membrane-stabilizing 283 properties positively influence photosynthetic efficiency under stressful conditions (Mapelli et 284 al., 2008). Exogenously application of Put increased the net photosynthetic rate of basil 285 (*Ocimum basilicum* L.) plants under drought stress, while electrolyte leakage was reduced 286 (Darabi *et al.*, 2020). PAs with high net positive charges can stabilize PSII proteins such as D_1 287 and D_2 and by binding to membrane proteins can stabilize the structure of the proteins during 288 stress (Hamdani et al., 2011).
- 289 The reductions in F_{ν}/F_m and qP were correlated with an increase in NPQ (Fig. 7). A decline in 290 F_{ν}/F_{m} indicates photoinhibition damage resulting from the incident PPFD when plants are 291 exposed to environmental stresses (Wang et al., 2018). We found that water-deficit-induced 292 NPQ was ameliorated by the PAs, meanwhile, leaf qP of PAs-treated plants was improved 293 under water deficit conditions. Sang et al. (2016) found that the NPQ of water-stressed 294 cucumber (*Cucumis sativus*) leaves treated with Spd was enhanced, indicating that Spd can 295 accelerate the dissipation of absorbed light under drought conditions. Water deficit leads to a 296 decrease in the ETR and the generation of excess excitation energy (Tezara et al., 2005). The 297 ETR of PAs-treated plants was higher than the untreated plants under water deficit, indicating 298 that the PSII reaction center of PAs-treated leaves maintained high activity. The negative 299 charges of LHCII can be neutralized with the PA's positive charges, resulting in the LHCII 300 complexes quenching by minimizing the exertion between the complexes (Hamdani et al., 301 2011).
- Exogenously applied PAs significantly increased leaf enzymatic antioxidant activities. Enzymatic antioxidant activity and higher ROS scavenging ability were enhanced by exogenously applied Spd in cucumber roots under stressful conditions (Wu *et al.*, 2018). In a study on tomato plants under heat stress, exogenous application of Spd regulated various signal transduction factors; mainly associated with genes related to stress signaling pathways such as

307 hormonal and sugar metabolisms (Cheng et al., 2012). To decrease the ROS content during 308 stressful conditions, the concentration of several nonenzymatic antioxidants, such as ASA and 309 lycopene, as well as enzymatic antioxidants including SOD, POD, APX, and CAT activity can 310 be enhanced by PAs (Hasan et al., 2021). ABA or Spd treatment alone decreased the activities 311 of both POD and CAT, which might be due to their contribution to the ROS accumulation 312 induced by ABA or Spd. However, Spd could activate the antioxidant enzymes to scavenge 313 H₂O₂ induced by ABA in the presence of ABA (Liu *et al.*, 2023). Our results also showed that 314 the increase in the antioxidant enzymes activities by PAs, when plants were fully irrigated, was 315 not as high as the water-stressed plants, which might be due to their lower ABA content. 316 However, under stressful conditions, higher ABA content probably activates the antioxidant 317 enzymes to scavenge higher H_2O_2 . Alcázar *et al.* (2006) also found that Put accumulation was 318 mainly an ABA-dependent metabolic response during drought.

319 PAs are regulators of redox homeostasis that play a dual role in plant oxidative stress (Saha et 320 al., 2015). Although PAs might be responsible for cellular breakdown due to generating the 321 strong oxidizers H_2O_2 under stressful conditions, H_2O_2 can act as a signaling molecule involved 322 in stress signal transduction (Groppa and Benavides, 2008). The H_2O_2 -mediated signaling 323 pathway, which is involved in salt stress-responsive genes (SIWRKY1, SIHKT1, SIDREB2, and 324 *SIMYB102*), was induced by exogenous Spd in tomato plants and significantly reduced the 325 adverse effects of salt stress (Raziq et al., 2022). Working on Salvia officinalis L. under drought 326 stress revealed that the enzymatic activities of APX and CAT were strongly coordinated 327 (Mohammadi-Cheraghabadi et al., 2021).

328 PAs foliar application alleviated the ROS-induced membrane damage and reduced leaf MDA, 329 EL, and H_2O_2 . MDA and H_2O_2 accumulation can be indicators of cell damage. A lower leaf 330 MDA and H_2O_2 content led to lower membrane damage and leaf EL. PAs can accelerate the 331 antioxidant enzyme activities to protect plants against the oxidative damages and membrane 332 injury or may enhance the biosynthesis of protective substances under stressful conditions. 333 Tomato (Solanum lycopersicum) leaf MDA content and EL were decreased by exogenous Spd 334 under drought stress (Sang *et al.*, 2016). Besides their properties as free radical scavengers, PAs 335 also stabilize biological membranes by binding to membrane phospholipids under stressful 336 conditions (Pál et al., 2015). Moradi Peynevandi et al. (2018) also observed that exogenously 337 applied PAs significantly decreased leaf H_2O_2 and MDA contents and enhanced the membrane 338 stability of cold-stressed stevia (*Stevia rebaudiana* Bertoni) plants.

339	Leaf Chla was decreased greater than Chlb content under water deficit conditions. This might
340	be due to a greater susceptibility of Chla than Chlb to stress or the generation of Chlb through
341	the degradation of Chla products (Sen et al., 2014; Shahba et al., 2010). This eventually led to
342	a decrease in Chla:b. It has been reported that PAs could protect the functional and structural
343	integrity of chloroplasts and slow down the rate of photosynthetic pigment degradation (Li et
344	al., 2014; Nahar et al., 2015). In our experiment, Spd foliar application increased the carotenoid
345	content of leaves. Carotenoids are also among the essential compounds playing a role in
346	protecting photosynthesis and stress-signaling pathways. Leaf carotenoid content was
347	positively correlated with leaf A_N , chlorophyll fluorescence, and antioxidative enzyme
348	activities (Fig. 7). Due to an increase in leaf carotenoid content under PAs application, it seems
349	that carotenoids are likely to prevent chlorophyll degradation due to their protective role (Dhar
350	<i>et al.</i> , 2020).

352 CONCLUSIONS

353 Our results revealed that exogenously applied PAs improved the drought tolerance of 354 chamomile plants in multiple ways. PAs improved leaf water status and alleviated oxidative 355 damage on the biological membranes by instigating leaf antioxidant content and reducing 356 membrane damage. The results here indicated that exogenously applied PAs act as regulators 357 to prevent chlorophyll degradation, protect the photosynthetic antenna and PSII structure, and 358 improve the photosynthetic efficiency of chamomile plants. Leaf antioxidant activities were 359 found to be determining and effective mechanisms to alleviate water deficit effects on 360 chamomile plants, induced by PAs. PAs inhibit water deficit-induced stomatal closure through 361 antioxidant enzyme-dependent H_2O_2 elimination. PAs can potentially improve water deficit 362 tolerance in chamomile. Furthermore, Spd showed the most effective impact in alleviating the 363 water deficit effects.

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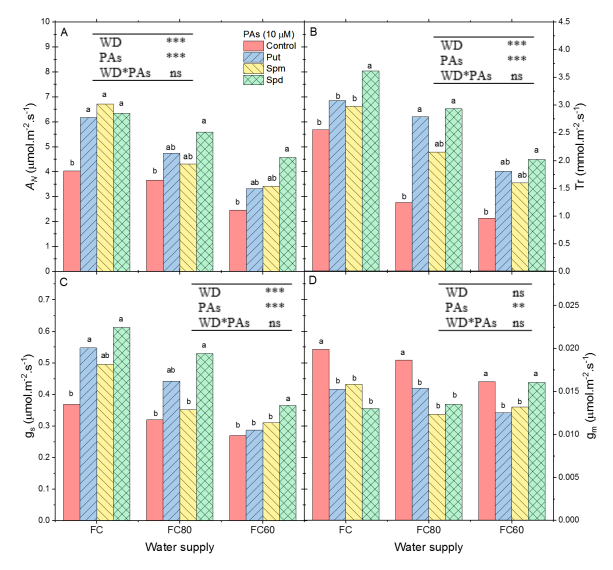
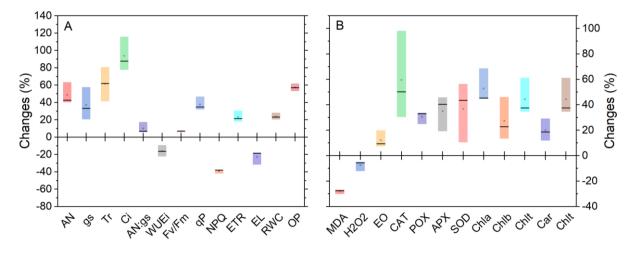
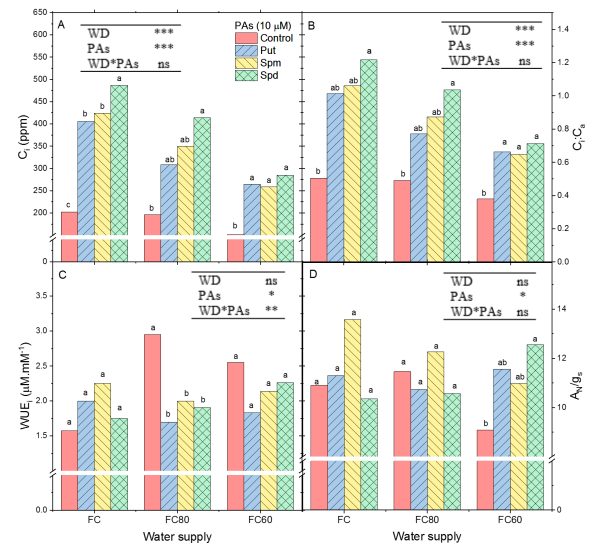


Fig. 1. Changes in (A) Net photosynthetic rate, (B) Transpiration rate, (C) Stomatal conductance, and (D) Mesophyll conductance of German chamomile leaves exposed to water deficit and foliar application of polyamines. WD: water deficit, PAs: polyamines. FC: field capacity, FC₈₀: 80% of FC, FC₆₀: 60% of FC. *, **, ***, and ns: significant at $p \le 0.05$, $p \le$ 0.01, $p \le 0.001$, and non-significant. Means with the same letters are not significantly different. LSD $p \le 0.05$. n=9.

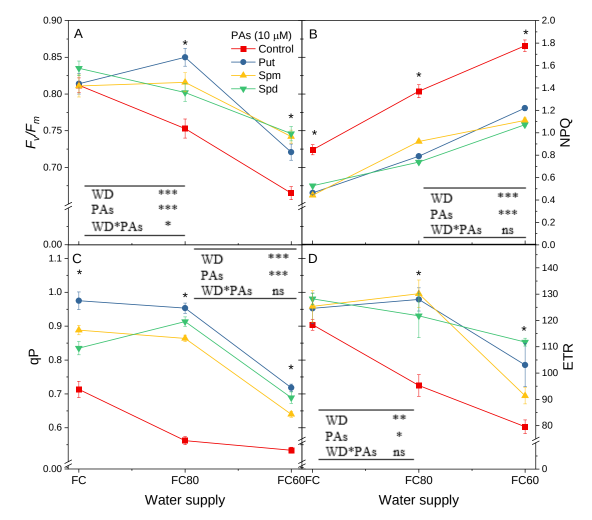


532 Fig. 2. Percent changes of (A) physiological and growth parameters, and (B) biochemical traits of German chamomile leaves affected by polyamines foliar application under water deficit 533 levels relative to the untreated plants. A_{N} : Net photosynthetic rate, g_{s} : Stomatal conductance, 534 535 T_i : Transpiration rate, C_i : Intercellular CO₂ concentration, A_N/g_s : Intrinsic water use efficiency, 536 WUE: instantaneous water use efficiency, F_v/F_m : Maximum photochemical quantum yield of 537 PSII, qP: Photochemical quenching, NPQ: Nonphotochemical quenching, ETR: Electron transport rate, EL: electrolyte leakage, RWC: Relative water contents, OP: Leaf osmotic 538 potential, MDA: Malondialdehyde, H₂O₂: Hydrogen peroxide, EO: Essential oil, CAT: 539 Catalase, POX: Peroxidase, APX: Ascorbate peroxidase, SOD: Superoxide dismutase, Chla: 540 541 Chlorophyll a, Chlb: Chlorophyll b, Chlt: Total chlorophyll, Car: Carotenoids. 542



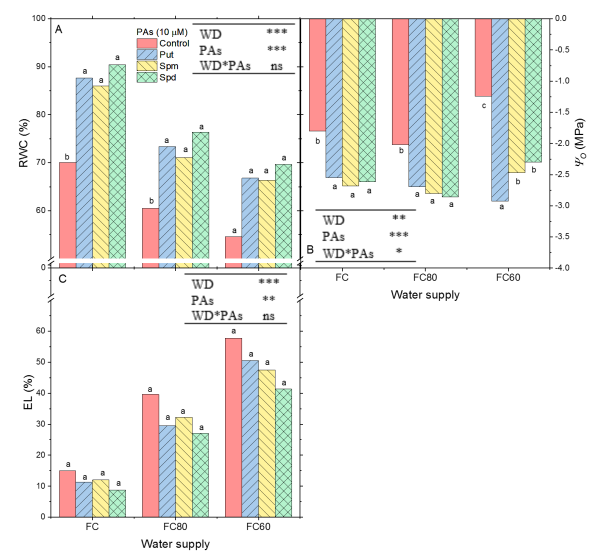
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Fig. 3. Changes in (A) Intercellular CO₂ concentration, (B) Intercellular to ambient CO₂ concentration (C) Instantaneous water use efficiency, and (D) Intrinsic water use efficiency of German chamomile leaves exposed to water deficit and foliar application of polyamines. WD: water deficit, PAs: polyamines. FC: field capacity, FC₈₀: 80% of FC, FC₆₀: 60% of FC. *, **, ***, and ns: significant at $p \le 0.05$, $p \le 0.01$, $p \le 0.001$, and non-significant. Means with the same letters are not significantly different. LSD $p \le 0.05$. n=9.



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Fig. 4. Changes in (A) Maximum photochemical quantum yield of PSII, (B) Nonphotochemical quenching, (C) Photochemical quenching, and (D) Linear electron transport rate of German chamomile leaves exposed to water deficit and foliar application of polyamines. FC: field capacity, FC₈₀: 80% of FC, FC₆₀: 60% of FC. WD: water deficit, PAs: polyamines. *, **, ***, and ns: significant at $p \le 0.05$, $p \le 0.01$, $p \le 0.001$, and non-significant. Error bars indicate the differences between the water deficit levels, ±SE. n=9.



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Fig. 5. Changes in (A) Relative water content, (B) Osmotic potential, and (C) Electrolyte leakage of German chamomile leaves exposed to water deficit and foliar application of polyamines. WD: water deficit, PAs: polyamines. FC: field capacity, FC₈₀: 80% of FC, FC₆₀: 60% of FC. *, **, ***, and ns: significant at $p \le 0.05$, $p \le 0.01$, $p \le 0.001$, and non-significant. Means with the same letters are not significantly different. LSD $p \le 0.05$. n=9.

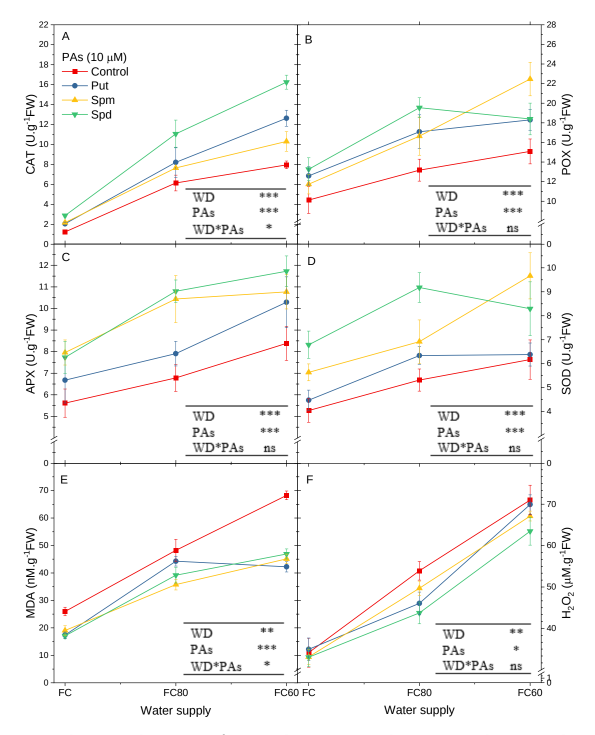


Fig. 6. Changes in the activity of (A) Catalase, (B) Peroxidase, (C) Ascorbate peroxidase, (D) Superoxide dismutase, the content of (E) Malondialdehyde, and (F) Hydrogen peroxide German chamomile leaves exposed to water deficit and foliar application of polyamines. FC: Field capacity, FC₈₀: 80% of FC, FC₆₀: 60% of FC. WD: water deficit, PAs: polyamines. *, **, ***, and ns: significant at p ≤ 0.05 , p ≤ 0.01 , p ≤ 0.001 , and non-significant. Error bars indicate the differences between the water deficit levels, \pm SE. n=9.

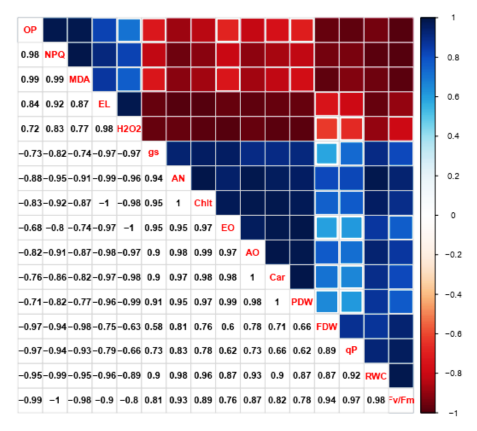


Fig. 7. Corplot analysis of OP: Leaf osmotic potential, NPQ: Nonphotochemical quenching, MDA: Malondialdehyde, EL: electrolyte leakage, H2O2: Hydrogen peroxide, gs: Stomatal conductance, A_N : Net photosynthetic rate, Chlt: Total chlorophyll, EO: Essential oil, AO: Antioxidant enzymes, Car: Carotenoids, qP: Photochemical quenching, RWC: Relative water contents, F_v/F_m : Photochemical quantum yield of PSII german chamomile plants under water deficit and polyamines foliar application.

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		mg.gFW ⁻¹				
Water deficit	PAs [‡] (10 μM)	Chlorophyll a	Chlorophyll b	Total chlorophyll	Carotenoids	Chlorophyll a:b
	Control	<mark>0.71</mark>	<mark>0.31</mark>	<mark>1.03</mark>	<mark>0.64</mark>	<mark>2.41</mark>
FC^{\dagger}	Put	<mark>0.93</mark>	<mark>0.36</mark>	<mark>1.29</mark>	<mark>0.75</mark>	<mark>2.58</mark>
FC'	Spm	<mark>0.98</mark>	<mark>0.39</mark>	<mark>1.38</mark>	<mark>0.78</mark>	<mark>2.50</mark>
	Spd	<mark>1.09</mark>	<mark>0.46</mark>	<mark>1.55</mark>	<mark>0.82</mark>	<mark>2.39</mark>
LSD		<mark>0.46</mark>	<mark>0.23</mark>	<mark>0.46</mark>	<mark>0.23</mark>	<mark>1.32</mark>
	Control	<mark>0.53</mark>	<mark>0.26</mark>	<mark>0.79</mark>	<mark>0.75</mark>	<mark>2.02</mark>
FC ₈₀	Put	<mark>0.83</mark>	<mark>0.27</mark>	<mark>1.11</mark>	<mark>0.86</mark>	<mark>3.38</mark>
ΓC_{80}	Spm	<mark>0.72</mark>	<mark>0.29</mark>	<mark>1.01</mark>	<mark>0.89</mark>	<mark>2.58</mark>
	Spd	<mark>0.90</mark>	<mark>0.37</mark>	<mark>1.27</mark>	<mark>0.96</mark>	<mark>2.42</mark>
LSD		<mark>0.41</mark>	<mark>0.21</mark>	<mark>0.54</mark>	<mark>0.46</mark>	<mark>2.21</mark>
	Control	<mark>0.33</mark>	<mark>0.20</mark>	<mark>0.54</mark>	<mark>0.76</mark>	<mark>1.64</mark>
FC_{60}	Put	<mark>0.54</mark>	<mark>0.25</mark>	<mark>0.78</mark>	<mark>0.80</mark>	<mark>2.07</mark>
1°C60	Spm	<mark>0.58</mark>	<mark>0.27</mark>	<mark>0.84</mark>	<mark>0.88</mark>	<mark>2.14</mark>
	Spd	<mark>0.66</mark>	<mark>0.30</mark>	<mark>0.96</mark>	<mark>0.99</mark>	<mark>2.25</mark>
LSD		<mark>0.35</mark>	<mark>0.15</mark>	<mark>0.43</mark>	<mark>0.51</mark>	<mark>1.29</mark>
Water deficit		***	***	<mark>***</mark>	ns	ns
Polyamines		***	**	<mark>***</mark>	*	<mark>ns</mark>
WD*PAs		<mark>ns</mark>	<mark>ns</mark>	<mark>ns</mark>	ns	ns

Table 1. Effect of water deficit and foliar application of polyamines on photosynthetic pigments of German chamomile leaves.

[†] FC: field capacity, FC₈₀: 80% of FC, FC₆₀: 60% of FC, ‡ PAs: polyamines. \Box Put: putrescine, Spm: spermine, Spd: spermidine. WD: water deficit, PAs: polyamines. *, **, ***, and ns: significant at p ≤0.05, p ≤ 0.01, p ≤ 0.001, and non-significant. Means with the same letters are not significantly different. LSD p≤0.05.

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کاهش تنش خشکی در بابونه آلمانی (.Matricaria chamomilla L) در پاسخ به تنش اکسیداتیو و بسته شدن روزنه 586 ناشی از کمبود آب توسط کاربرد خارجی پلی آمین ها

$$588$$
 محمد جواد احمدی لاهیجانی 1^* ، جعفر نباتی 1 ، سعید موری 2 ، محمد کافی 1

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چکیدہ

591 یلی آمین ها مولکول های پیامرسانی هستند که نقش های امیدوار کنندهای در بهبود تحمل به تنش در گیاهان نشان دادهاند. اطلاعات 592 محدودي در مورد اثرات كاربرد خارجي يلي آمينها بر روي گياهان دارويي از جمله بابونه در دسترس است. اين آزمايش بهمنظور 593 بررسی اثرات محلولپاشی پلیآمینها [پوترسین (Put)، اسپرمیدین (Spd) و اسپرمین (Spm)] بر فرآیندهای فیزیولوژیکی و 594 بیوشیمیایی برای در ک مکانیسمهای احتمالی مربوط به کاهش اثرات تنش کمبود آب [ظرفیت مزرعه (FC) به عنوان شاهد، 80 درصد ظرفیت مزرعه (FC80) و 60 درصد ظرفیت مزرعه (FC60)] در بابونه آلمانی انجام شد. نتایج نشان داد که پلی آمین ها تا حدی 595 596 بستهشدن روزنه ناشی از کمبود آب را مهار می کند و آنزیمهای آنتی اکسیدانی را برای از بین بردن افزایش پر اکسیدهیدروژن القا می کند. اسپرمیدین هدایت روزنهای را در FC80 ،FC و FC60 به ترتیب 66، 65 و 35 درصد در مقایسه با شاهد افزایش داد. افزایش 597 هدايت روزنهاي فتوسنتز خالص بر گ را در FC80 و FC60 به ترتيب 52 و 86 درصد در مقايسه با شاهد افزايش داد. نقش يلي آمين ها 598

599	در کاهش تنش اکسیداتیو با همبستگی منفی فعالیتهای آنتیاکسیدانی برگ و محتوای مالوندیآلدئید و پراکسیدهیدروژن تایید
600	شد. با توجه به نتایج، پلیآمینها به عنوان ترکیبات محافظ تنش عمل میکنند تا آنزیمهای آنتیاکسیدانی را برای حذف
601	پراکسیدهیدروژن ناشی از تنش، بهبود پایداری غشاء و افزایش تحمل کمبود آب تحریک کنند. بهطورکلی، نتایج نشان داد که
602	پلی آمین ها می توانند تنظیم کننده های بالقوه رشد برای کاهش تنش کم آبی خفیف تا شدید باشند.
603	کلیدواژ گان: فلورسانس کلروفیل؛ آنتی اکسیدان آنزیمی؛ پراکسیدهیدروژن؛ متغیرهای تبادل گاز؛ خاموشی غیر فوتوشیمیایی

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