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Distinctive phytohormonal and metabolic profiles of *Arabidopsis thaliana* and *Eutrema salsugineum* under similar soil drying

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1 **Distinctive phytohormonal and metabolic profiles of *Arabidopsis thaliana* and *Eutrema***
2 ***salsugineum* under similar soil drying**

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42 25 Running title: **Arabidopsis and Eutrema responses to soil drying**

43 26

27 **Abstract**

28

29 Although plants perceive and respond to soil drying via a series of concurrent physiological
30 and molecular events, drought tolerance differs greatly within the plant kingdom. While
31 *Eutrema salsugineum* (formerly *Thellungiella salsuginea*) is regarded as more stress tolerant
32 than its close relative *Arabidopsis thaliana*, their responses to soil water deficit have not been
33 compared. To ensure a similar rate of soil drying for the two species, daily soil water
34 depletion was controlled to 5-10 % of the soil water content. While partial stomatal closure
35 occurred earlier in *Arabidopsis* (Day 4) than *Eutrema* (from Day 6 onwards), thereafter both
36 species showed similar stomatal sensitivity to drying soil. Nonetheless, both targeted and
37 untargeted metabolite analysis showed larger responses in *Arabidopsis* in both early/mild
38 drought (Days 1, 3, 5 – no significant change in leaf relative water content, RWC) and
39 late/severe drought (Day 12 – circa 80 % decrease in leaf RWC). *Arabidopsis* (but not
40 *Eutrema*) showed early peaks in foliar abscisic acid (ABA), jasmonic acid (JA) and salicylic
41 acid (SA) contents. *Arabidopsis* showed greater metabolic adjustment than *Eutrema* during
42 both early drought stress (428 versus 35 variables significantly changing) and severe drought
43 (607 versus 171 variables significantly changing). Different sugar profiles between species
44 were accompanied by opposing patterns in the bioactive cytokinin profile responses. The
45 distinctive metabolic responses of each species during early drought, which occurred prior to
46 leaf water status declining, were apparently independent of later stomatal closure in response
47 to drought. These biochemical differences can have implications in regulating transpiration,
48 since *Eutrema* reduced whole plant water use very early (Day 3) while this occurred later
49 (Day 6 onwards) in *Arabidopsis*. *Arabidopsis* provides a promising model to evaluate the
50 mechanisms responsible for stress-induced growth inhibition under the mild/moderate soil
51 drying that crop plants are typically exposed to.

52

53 Keywords: bioactive cytokinins; rewatering; redox state; stomatal conductance; unsupervised
54 multivariate analysis

55 Introduction

56 In view of climate change, a major goal for the plant biology community is to understand the
57 mechanisms that allow some plants to withstand drought or hot weather. Knowledge of how
58 plants survive and reproduce in challenging environmental conditions can allow novel targets
59 to be tested in crop-breeding programs. The well-known model species *Arabidopsis thaliana*
60 provides information that can be applied to crop systems (Piquerez *et al.*, 2014; Gilliam *et*
61 *al.*, 2017). Using the Columbia accession (Col-0) and its mutants has allowed many stress
62 regulatory and responsive pathways to be deciphered (Koornneef and Meinke, 2010; Osakabe
63 *et al.*, 2014), although its stress resilience has not been fully established. Despite wide
64 ecotypic variation (Montesinos-Navarro *et al.*, 2011; Clauw *et al.*, 2016), *Arabidopsis* is not
65 expected to cope well in extreme environments (Zhu *et al.*, 2015). Instead, *Arabidopsis*
66 relatives such as *Eutrema salsugineum* have been proposed as stress-tolerant models (Orsini
67 *et al.*, 2010; Zhu *et al.*, 2015). *Eutrema* seems prepared for stress, as its stress-related genes
68 are upregulated in comparison to *Arabidopsis* even when grown under optimal conditions
69 (Taji *et al.*, 2004; Gong *et al.*, 2005). As in *Arabidopsis*, *Eutrema salsugineum* ecotypes from
70 different geographical regions show significant genetic variation (Lee *et al.*, 2016). However,
71 physiological and metabolic responses of *Arabidopsis* and its stress tolerant relatives to soil
72 water deficit have not been directly compared.

73 Physiological responses to water deficit are modulated by the intensity, duration, and rate of
74 progression of imposed drought (Pinheiro and Chaves, 2011). Extensive research on the
75 stomatal regulation of water loss demonstrates a trade-off between carbon assimilation,
76 efficient water use and leaf cooling capacity (Chaves *et al.*, 2016). Plants can be grouped
77 according to whether they avoid heat (keeping their stomata open for longer) or use water
78 efficiently (closing their stomata sooner, a typical drought-avoidance strategy). However, if
79 plants can avoid the deleterious effects of heat by keeping their stomata open for longer,
80 while maintaining a favourable water status by extracting more water (e.g. by having deep
81 roots), this strategy benefits carbon uptake in addition to the cooling effect. Under drought,
82 *Arabidopsis* Col-0 closes its stomata at higher soil moisture levels than other *Arabidopsis*
83 genotypes (Meyre *et al.*, 2001). The two well-studied ecotypes of *Eutrema*, Shandong and
84 Yukon, can grow under limited soil water availability (Xu *et al.*, 2014; Macleod *et al.*, 2015),
85 but their drought performance, relative to *Arabidopsis*, is unknown.

86 The two plant species seemingly have distinct water consumption strategies, although it may

87 be difficult to separate species *versus* accession variation. Arabidopsis (Col-0) had relatively
88 higher total transpiration than Eutrema (Shandong) under non-challenging conditions, which
89 was related to its higher relative growth rate (Orsini *et al.*, 2010). Salinity decreased
90 transpiration to a larger extent in Arabidopsis than Eutrema. [In addition to these different](#)
91 [water consumption strategies](#), Eutrema and Arabidopsis also had different biochemical
92 composition under non-challenging growth conditions, with foliar sucrose and glucose
93 content higher in Eutrema, while the hormones salicylic acid (SA) and jasmonic acid (JA)
94 were higher in Arabidopsis (Arbona *et al.*, 2010; Pilarska *et al.*, 2016). [Furthermore, Eutrema](#)
95 [expressed more stress and defence genes than Arabidopsis under non-challenging conditions,](#)
96 [which is described as stress priming \(e.g. Zhu *et al.*, 2015; Lee *et al.*, 2016\).](#) It is uncertain
97 [whether these biochemical differences regulate differences in transpiration, and consequently](#)
98 [different rates of soil water depletion.](#)

99 However, the metabolic features associated with the initial stages of soil drying are not clear.
100 In Arabidopsis, soil drying partially closes the stomata well before any decrease in carbon
101 assimilation rate (Hummel *et al.*, 2010; Bechtold *et al.*, 2016) or any significant increase in
102 foliar abscisic acid (ABA) content (Bechtold *et al.*, 2016). ABA is described as the main
103 driver controlling plant performance under limited water availability since it induces stomatal
104 closure, but more comprehensive recent studies demonstrated that most of the plant hormones
105 are involved in stress signalling (Müller and Munné-Bosch, 2015). In addition, during the
106 very early stages of water limitation, effects on carbon metabolism (CO₂ assimilation, and
107 sucrose and starch formation and allocation) may be decoupled from stomatal closure
108 (Pinheiro *et al.*, 2011; Bechtold *et al.*, 2016). Many players are involved in stress perception
109 and signal transduction leading to large alterations in carbon metabolism, and in the
110 transcription program (Golldack *et al.*, 2014; Urano *et al.*, 2017). The metabolic balance
111 between several molecules triggers adjustment mechanisms, and when several thresholds are
112 achieved, physiological responses to drought occur (Pinheiro *et al.*, 2011). The integration of
113 multiple environmental signals by sugars, hormones, and reactive oxygen species (ROS)
114 adjusts plant growth and determines whether plants survive or perish under given
115 environmental conditions (Pinheiro and Chaves, 2011; Osakabe *et al.*, 2014). The precise
116 chain of events is not yet defined, and although some pathways and interactions are
117 understood, others are more elusive (Rivas-San Vicente and Plasencia, 2011; Munné-Bosch
118 and Müller, 2013; Ruan, 2014; Considine and Foyer, 2014; de Ollas and Dodd, 2016).
119 Although recent reports highlight that stomatal closure is one of the initial events in response

120 to soil drying, many other metabolic adjustments also take place.

121 This work aimed to explore the metabolic adjustments prior to significant stomatal closure in
122 *Arabidopsis* (Col-0) and *Eutrema* (Shandong) responding to slowly-imposed soil water
123 deficit. Our working hypothesis is that *Arabidopsis* and *Eutrema* show distinctive responses
124 to progressive and slow soil drying. In addition, untargeted analysis of LC-MS metabolite
125 data was used to detect similarities and differences between these two species.

127 **Materials and Methods**

128 *Arabidopsis thaliana* (Col-0) and *Eutrema salsugineum* (Shandong) seeds were soaked and
129 stratified at 4 °C for 4 or 14 days, respectively. *Eutrema salsugineum* is the current
130 designation of *Thellungiella salsuginea* (Integrated Taxonomic Information System on-line
131 database, www.itis.gov; The International Plant Names Index, www.ipni.org). Seeds were
132 then transferred to pots (300 mL) containing a 1:1 mixture of coarse sand and peat
133 (Shamrock). Plants were grown under controlled conditions, under a 12 h photoperiod,
134 temperatures ranging from 20 to 24 °C, with a 60-70 % relative humidity and
135 photosynthetically active radiation (PAR) of 250-300 $\mu\text{mol m}^{-2} \text{s}^{-1}$ (SON-T Agro 400w,
136 Phillips). Plants were watered every day with demineralized water to 85 % of soil water
137 content (SWC). SWC was monitored daily and is defined as: $\text{SWC} = [(\text{pot weight} - \text{pot}$
138 $\text{weight with totally dried substrate})] / [(\text{pot weight at drained capacity} - \text{pot weight with}$
139 $\text{totally dried substrate})] \times 100$. Drought stress treatments were imposed when plants had 8 to
140 10 fully expanded leaves (40 days for *Eutrema* and 36 days for *Arabidopsis*) and had covered
141 the surface of the pots (thereby minimising evaporation from the soil). During the experiment
142 plant growth increased, on average by 2.9 g fresh weight (FW) for *Arabidopsis* and 2.5 g FW
143 for *Eutrema*, corresponding to less than 0.8 % error in estimating SWC (Fig. 1).

144 Preliminary drought experiments, in which water was withheld, showed faster soil water
145 depletion and more rapid stomatal closure in *Arabidopsis* (supplementary Fig. S1). Similarly,
146 higher transpiration rates of *Arabidopsis* were previously reported (Orsini *et al.* 2010).
147 Stomatal conductance of the two species was differentially sensitive to soil drying
148 (supplementary Fig. S1B and C). Within the 45-55 % SWC range, *Arabidopsis* showed
149 greater stomatal closure than *Eutrema*, but below 40 % SWC both species showed similar
150 stomatal sensitivity to soil water deficit and were severely affected by drought. However,

151 analysis of covariance demonstrating no significant species x SWC interaction, both species
152 showed a similar relationship between % gs vs soil water content, with (supplementary Fig
153 S1C).

154 To compare stress duration and intensity effects on plant responses, the rate of soil water
155 depletion was controlled to 5-10 % of the SWC per day by pre-dawn irrigation (Fig. 1). Even
156 when controlling the SWC, Arabidopsis consumed more water than Eutrema, as indicated by
157 the greater divergence between SWC measured at maximum soil water deficit (symbols) and
158 the SWC to which the pot was re-turned to pre-dawn (“stress” line in Fig 1). This greater
159 water use of Arabidopsis was most prominent between Days 3 and 7.

160 Plants were harvested 0 (last day of watering), 1, 3, 5 and 12 days after the beginning of the
161 assay, corresponding to 75 %, 66 %, 45 %, and 12 % SWC, respectively. Samples were also
162 taken the day after re-watering (1 d). Six biological replicates were obtained at each time-
163 point, except for Day 1 controls for which there were only five biological replicates,
164 providing 65 samples of each plant species. At the beginning of the assay, the most recently
165 expanded two-three leaflets were identified and used for physiological and water status
166 measurements. For the biochemical analysis, and when analysing severe drought and early
167 rewatering, only non-senescent leaflets were used, i.e. the younger leaflets. Samples for
168 biochemical (hormone, carbohydrate, pigment and oxidative status) analysis were
169 immediately frozen in liquid nitrogen and kept at -80 °C until further extraction and analysis.
170 Samples for osmotic potential and for RWC were then collected.

171 *Leaf conductance, water status and osmotic adjustment*

172 Stomatal conductance was measured 2-3 h after the beginning of the photoperiod in five
173 plants per treatment using a portable gas exchange photosynthesis system coupled to a 6400-
174 15 chamber (1 cm² diameter cuvette, Li-6400, Li-Cor, Lincoln, Nebraska, USA). Three to
175 five measurements were made per plant on the most recently expanded leaf.

176 Leaf and root samples were taken 4 h after the beginning of the photoperiod. Leaf discs (3
177 mm diameter) and total roots were weighed to obtain fresh weight (FW), placed in darkened
178 petri dishes containing distilled water for 2 h to fully hydrate, then re-weighed to obtain
179 turgid weight (TW), and then dried at 80 °C for 48 h to obtain dry weight (DW). Leaf
180 (LRWC) and root (RRWC) relative water content were calculated as: $RWC = [(FW - DW) \times$
181 $100 / (TW - DW)]$.

182 Leaf osmotic potential (ψ_s) was evaluated from leaf disks (8 mm, n = 5–6), frozen and stored
183 at -80 °C. The leaf osmotic potential was measured with an HR-33T dew point
184 microvoltmeter and C-52 sample chambers (Wescor, Inc., Logan, UT, USA). The osmotic
185 potential was adjusted to the LRWC to calculate the osmotic potential at full turgor (OP100),
186 and the osmotic adjustment was calculated as previously described (Turner *et al.*, 2007).

187 *Phytohormone quantification via LC-MS targeted analysis*

188 Freeze-dried shoots (50 mg) were used to extract and quantify the following hormones
189 (Müller and Munné-Bosch, 2011): auxin (indole-3-acetic acid: IAA), gibberellins (GA1, 4, 9,
190 19, 20, 24), cytokinin (CK) compounds (trans-zeatin: Z; trans-zeatin riboside: ZR; 2-
191 isopentenyl adenine: 2iP; isopentenyl adenosine: IPA; dihydrozeatin: DHZ; dihydrozeatin
192 riboside: DHZR), and stress-related phytohormones (ABA; JA; SA; and ethylene precursor
193 1-amino-cyclopropane-1-carboxylic acid: ACC).

194 Extraction was performed in methanol solutions containing 1 % glacial acetic acid, using the
195 following standards: d₅-IAA, d₆-2-isopentenyl adenine (d₆-2iP), d₆-IPA, d₆-ABA, d₅-JA, d₄-
196 SA, d₄-ACC, d₂-GA₁, d₂-GA₄, d₂-GA₉, d₂-GA₁₉, d₂-GA₂₀ and d₂-GA₂₄; d₅-Z and d₅-ZR were
197 used as standards for Z, DHZ, ZR, and DHZR. After adding 170 μ L of the extraction solution
198 and 30 μ L of a solution containing 100 ppm of the standards in the same solvent, the
199 materials were mixed in a vortex mixer for 5 s and exposed to ultrasound for 30 min,
200 followed by centrifugation at 9,500 g for 10 min. The supernatant was removed and the
201 residue was washed twice with 100 μ L of the solvent solution. The supernatant and washes
202 were combined and filtered through PTFE 0.22 μ m filter paper (Waters, Milford, MA) and 5
203 μ L aliquots were analysed using a UPLC-ESI-MS/MS (Acquity UPLC System from Waters,
204 Milford, MA) and tandem MS/MS experiments were performed on an API 3000 triple
205 quadrupole mass spectrometer (PE Sciex, Concord, Ont., Canada) using a HALO™ C18
206 column (2.1 \times 75 mm, 2.7 μ m) (Advanced Materials Technology, Inc. Wilmington, DE) and
207 a binary mobile phase system composed of (A) water modified with 0.05 % glacial acetic
208 acid and (B) acetonitrile modified with 0.05 % glacial acetic acid. Quantification was
209 performed by preparing a calibration curve including each of the analysed compounds and
210 calculating the compound/standard ratio using Analyst™ software (Applied Biosystems, Inc.,
211 Foster City, CA). The results were expressed on a dry weight (DW) basis.

212 *Ascorbate oxidative status*

213 Ascorbate reduced and oxidized forms were determined by a plate-reader method (Queval
214 and Noctor, 2007) with slight modifications. Briefly, lyophilised leaves (20 mg DW) were
215 placed in a microcentrifuge tube with two tungsten balls and ground under liquid nitrogen in
216 a Retsch MM300 Bead Mill Cell Disrupter (Retsch GmbH & Co Haan Germany).
217 Subsequently, 1 mL of extraction buffer (6 % meta-phosphoric acid) was added, vortexed for
218 1 min and clarified by centrifugation at 10,000 g (10 min, 4°C). Finally, extracts were
219 neutralized and adequately diluted before spectrophotometric readings on a 96 well quartz
220 microplate (Hellma Hispania SL, Badalona, Spain). The levels of ascorbate (AscA) (reduced)
221 and dehydroascorbate (DHA) (oxidized) were determined using ascorbate oxidase (AO) and
222 dithiothreitol (DTT), respectively (Foyer *et al.*, 1983). AO specifically oxidizes all AscA in
223 the sample. Therefore, the decrease in O.D. at 265 nm is related to AscA content.
224 Alternatively, when the samples are incubated with DTT, DHA is reduced to AscA and the
225 increase in O.D. is proportional to the initial DHA content. The ascorbate oxidative status
226 was estimated as $DHA/(DHA + AscA)$.

227 ***Photosynthetic pigments quantification***

228 For pigment extraction, lyophilised leaf samples (15 mg DW) were placed in a
229 microcentrifuge tube with two tungsten balls, ground under liquid nitrogen in a Retsch
230 MM300 Bead Mill Cell Disrupter (Retsch GmbH & Co Haan Germany), and extracted with
231 ice-cold 80 % acetone (v/v). After centrifuging at 6,500 g for 10 min at 4 °C, the supernatant
232 was collected and the pellet was re-extracted with the same solvent until it was colourless.
233 Then, supernatants were pooled and analysed spectrophotometrically. Specific absorption
234 coefficients in 80 % acetone previously reported were used to quantify chlorophyll a,
235 chlorophyll b and carotenoids (Lichtenthaler and Buschmann, 2001).

236 ***Extraction of water soluble carbohydrates and starch***

237 Water-soluble carbohydrates were extracted from freeze-dried leaf material following a
238 chloroform:methanol method previously described (Antonio *et al.*, 2008). Briefly, 50 mg DW
239 of leaf material was ground in liquid nitrogen and extracted with 250 µL ice-cold chloroform:
240 methanol (3:7, v/v), vortex-mixed and incubated at -20 °C for 2 h. After incubation, samples
241 were extracted twice with ice-cold water, and after centrifugation at 17,900 g at 4 °C for 10
242 min, the upper phases were collected and pooled. The combined supernatants containing the
243 water-soluble carbohydrates were evaporated to dryness using a centrifugal concentrator
244 (Savant SpeedVac Plus SC110A, Thermo Electron Corporation, Runcorn, UK). Samples

245 were reconstituted in 100 μ L water and centrifuged at 6,800 g at 20 $^{\circ}$ C for 30 min, followed
246 by LC-MS analysis.

247 For starch analysis, the pellet resulting from the chloroform:methanol extraction was washed
248 twice with water. Ten volumes of water were added to the pellet, boiled for 3 min, and
249 autoclaved at 130 $^{\circ}$ C for 1 h. After cooling, samples were incubated with 6 U
250 amyloglucosidase (Roche Applied Science, Amadora, Portugal) for 2 h at pH 4.8 and 60 $^{\circ}$ C.
251 Starch was quantified in the supernatant using a starch enzymatic quantification kit (n $^{\circ}$
252 10207748035, R-Biopharm Aktiengesellschaft, Darmstadt, Germany) and by making use of
253 the Hatterscheid and Willenbrink modification as previously described (Pinheiro *et al.*,
254 2001).

255 *Untargeted LC-MS analysis of the water-soluble carbohydrate fraction*

256 Arabidopsis and Eutrema samples were analysed as separate cohorts. In each case, samples
257 were randomized and run in batches of eight or nine with the injection of a pooled sample
258 between batches for quality control (QC). LC-MS analyses were performed on a Dionex
259 U3000 2D HPLC system coupled to a Bruker maXis UHR-Q-TOF MS with an ESI interface.
260 Analytes were detected in the negative ion mode using the following MS parameters:
261 capillary voltage, 4500 V; nebulizer gas, 2 Bar; drying gas, 8.0 L/min; drying temperature,
262 200 $^{\circ}$ C, and collision energy, -10.0 eV. Mass spectra were acquired over the scan range m/z
263 50-1000. Chromatographic separation was carried out using a porous graphitic carbon (PGC)
264 HypercarbTM column (5 μ m, 100 mm \times 4.6 mm; Thermo Electron, Runcorn, Cheshire, UK)
265 at a flow rate of 600 μ L min⁻¹. All samples were reconstituted with 500 μ L deionised water
266 with a further 50-fold dilution in deionised water to prevent signal saturation and to minimise
267 matrix effects. The sample injection volume was 20 μ L and the PGC column was used at
268 ambient temperature (25 $^{\circ}$ C). The binary mobile phase was composed of (A) water modified
269 with 0.1 % (v/v) formic acid (FA) and (B) acetonitrile modified with 0.1 % FA. The gradient
270 elution was as follows: 0-4 min maintained at 2 % B; 4-7 min, 2 to 8 % B; 7-10 min 8-25 %
271 B and maintained for 3 min, followed by column regeneration and re-equilibration: 13-19
272 min, 25 to 40 % B; 19-19.5 min, 40 to 50 % B held for 1 min; 20.5-21 min 50 to 99 % B held
273 for 2 min; 23-25 min 99 to 2 % B and maintained for 10 min. All solvents were purchased
274 from Fisher Scientific except FA, which was purchased from Sigma Aldrich.

275 *Statistical analysis*

276 Raw LC-MS data were pre-processed using Progenesis QI (Nonlinear Dynamics, Newcastle
277 Upon Tyne, UK). Mass spectra were aligned by retention time and normalized to the same
278 total ion count before peak picking was performed to provide a matrix of potential
279 metabolites for each observation, annotated by the accurate mass (m/z between 50 and 1000)
280 and retention time (between 1 and 30 min) of the corresponding peak. In total, 53208 and
281 33032 peaks were recorded for Arabidopsis and Eutrema, respectively, and were used as
282 variables in multivariate and univariate analyses.

283 When the Eutrema data were scaled to unit variance to allow smaller variables to contribute
284 to the analysis, differences between batches became apparent, with the last two batches
285 differing substantially from the rest (supplementary Fig. S2A). Liquid chromatography-mass
286 spectra are often acquired batch-wise to allow necessary calibrations and cleaning of the
287 instrument. However, this may introduce further sources of variation, such as differences in
288 the conditions under which data for individual batches is acquired. Quality control (QC)
289 samples are frequently employed to both judge and correct for this variation.

290 However, batch correction using the QC observations increased inter-batch variation as the
291 change in observations between batches was often not well-represented by the change in
292 corresponding QCs. Therefore, background correction for each variable was performed
293 (Rusilowicz *et al.*, 2016; Wehrens *et al.*, 2016). This method identifies a background trend,
294 using experimental observations as well as the QCs, with which to adjust the intensities. The
295 run order for data collection was randomized, but by chance a disproportionate number of
296 early-stress observations occurred in batch 3 and several late-stress observations in batch 4.
297 With the exception of these two batches, which were combined, we used a separate trend for
298 each batch, obtained as a moving median with a window width of 5 observations. The
299 effectiveness of batch correction was assessed using the Bhattacharyya distance (Wehrens *et*
300 *al.*, 2016). In addition, an outlier that dominated the variance after scaling was removed
301 before calculating the trend. Control correction was also performed on each variable to
302 remove differences due to growth. For each day of harvest, this was achieved by subtracting
303 the median over the six control replicates from the corresponding variable in the water-
304 stressed observations for that day. The Arabidopsis data showed no obvious differences
305 between batches (supplementary Fig. S3), and therefore, batch correction was deemed
306 unnecessary but control correction was performed to prevent differences due to growth from
307 masking early-stress characteristics.

308 Principal components analysis (PCA) was used for unsupervised multivariate analysis with
309 both unscaled data and after scaling to unit variance to prevent high content metabolites
310 dominating the analysis. To identify patterns in metabolites over time, k-means cluster
311 analysis was performed with the control-corrected time-series for both datasets. The obtained
312 initial clusters were filtered using the sum of squared values to remove the time-series for
313 metabolites that did not differ appreciably between drought and control observations, i.e.
314 where all values in the control-corrected time-series were close to zero. Cluster analyses of
315 the remaining time-series (with various values of k) showed the largest cluster to consist of
316 less interesting time-series and an iterative filtering process was used to reduce the number of
317 time-series, leaving only those with the most interesting patterns. In each iteration, k-means
318 clustering with $k = 15$ was performed and the largest cluster removed before the next
319 analysis. After four iterations, 46 time-series remained and were clustered using k-means
320 with $k = 9$.

321 Univariate analyses were performed using the non-parametric Mann-Whitney U-test with
322 Benjamini-Hochberg correction for multiple comparisons (Benjamini and Hochberg, 1995).
323 Three-way group comparisons were carried out (early stress/late stress/rewatered and Days 1,
324 3 and 5 for each species) with one-way ANOVA and Tukey's honest significant difference
325 (HSD) correction for multiple pairwise testing. Data correction methods were implemented
326 using C code written in-house and statistical analyses were performed in the R platform,
327 version 2.13.1 (R Core Team, 2016) or in Matlab (The MathWorks Inc., Natick, MA, USA).

328 Analysis of covariance (ANCOVA) discriminated possible species difference in stomatal
329 sensitivity to drying soil.

331 **Results and discussion**

332 *Stomatal sensitivity to drying soil and plant water status*

333 Under well-watered conditions, stomatal conductance (g_s) of both species exceeded 0.11 mol
334 $\text{m}^{-2} \text{ s}^{-1}$ (supplementary Fig S4A). Since g_s of well-watered plants varied from day to day, g_s
335 of plants in drying soil was normalised according to the average well-watered values of each
336 species. As the soil dried (Fig. 2), partial stomatal closure of Arabidopsis and Eutrema was
337 detected on Days 4 and 6, respectively (Fig. 2). Within the 45-55 % SWC range, Arabidopsis
338 showed greater stomatal closure than Eutrema, but below 40 % SWC both species showed

339 similar stomatal sensitivity to soil water deficit and were severely affected by drought.
340 Stomatal conductance responded sluggishly to re-watering, with limited recovery
341 (supplementary Fig S4A). Across the entire experiment, both species showed a similar
342 relationship between % gs vs soil water content, with analysis of covariance demonstrating
343 no significant species x SWC interaction (supplementary Fig S4B). Thus, both species
344 showed similar stomatal sensitivity to drying soil.

345 Initial stomatal closure was not associated with decreased leaf water status, i.e. lower cell
346 volume did not trigger early stomatal closure (Sack *et al.*, 2018). On imposing soil water
347 deficit, no significant differences in *Eutrema* leaf (and root) RWC were detected until Day 5
348 (supplementary table S2). In *Arabidopsis*, leaf RWC transiently decreased on Day 3
349 (supplementary table S1). Although statistically significant (at the 95% confidence level), its
350 small magnitude (~4%) could be within the method error or due to daily fluctuations.

351 In contrast to plant water status, the water consumption patterns changed very early on, but
352 were not temporally correlated with stomatal closure. Compared with its well-watered
353 control, *Eutrema* started to lose less water from Day 3 onwards (3 days before any significant
354 stomatal closure), as indicated by the slope of the soil RWC% line for plants in drying soil
355 (Fig. 1). In contrast, *Arabidopsis* started to use less water on Day 6 onwards (two days after
356 partial stomatal closure occurred). This suggests that earlier growth inhibition of *Eutrema*,
357 decreased whole plant water loss independent of changes in plant water status.

358 By Day 12, leaf RWC of both species had declined to very low values (< 20 %) and leaflets
359 selected for water status measurements (those most recently expanded at the onset of the
360 assay) were severely wilted and exhibited senescence symptoms. Lower leaf chlorophyll
361 fluorescence (Fv/Fm) and lower chlorophyll a content indicated photo-inhibition and/or leaf
362 senescence (Kalaji *et al.*, 2016).

363 Despite the severity of the stress imposed, root water status of both species recovered within
364 24 h of re-watering. Root RWC of *Eutrema* was similar to those of the well-watered controls,
365 while the root RWC of *Arabidopsis* was ~90 % of that of the controls. However, leaf RWC
366 remained low, only ~50 % and ~40 % of the well-watered control values in *Eutrema* and
367 *Arabidopsis* respectively (supplementary tables S1 and S2). In addition, Fv/Fm tended to
368 increase in *Eutrema*, but values were unaffected in *Arabidopsis* (supplementary tables S1 and
369 S2).

370 Traditionally, it has been argued that only resurrection plants can survive such severe
371 drought, i.e. recover from leaf RWC values below 20% (Dinakar and Bartels, 2013). Since
372 leaf RWC was determined in the most recently expanded leaves at the beginning of the assay
373 (see Material and Methods), these older leaves were severely wilted and senescent after 12
374 days, while younger leaves visually maintained turgor. Several reports indicate that
375 Arabidopsis Col-0 plants are able to recover from severe drought, with 30% of Col-0 plants
376 surviving exposure to 15% SWC and severe wilting (Sun et al. 2013) while 20% of severely
377 wilted Col-0 plants survived SWCs < 20% (Zhao *et al.* 2016). Moreover, Col-0 plants with
378 40-50% leaf RWC recovered from drought (Meyre *et al.* 2001; Tran *et al.* 2007; Kosma *et al.*
379 2009; Koffler *et al.* 2014) while some plants recovered from 20% leaf RWC although the
380 survival percentage was very low (Lü *et al.* 2012; Nguyen *et al.* 2016). In contrast to
381 Arabidopsis, Shandong was able to recover from drought if the leaf RWC declined to 50%,
382 but not 30% (Dedrick 2007). Since our measurements were made only 1 Day after rewatering
383 and no plants were available to evaluate long-term recovery, irreversible damage cannot be
384 ruled out.

385 As small changes in soil water content (10-15 %) affect not only leaf conductance but also
386 plant metabolism (Davies *et al.*, 1990; Pinheiro *et al.*, 2011), both untargeted metabolite
387 analysis and targeted metabolite/biochemical analysis aimed to determine the impact of
388 gradually declining soil water availability on leaf metabolism. This approach aimed to detect
389 species differences in metabolism when plant water status was not affected (up to Day 5).

390

391 *Untargeted metabolite analysis*

392 The responses to soil water depletion in Arabidopsis and Eutrema were analysed via
393 untargeted LC-MS, making use of the water-soluble fraction. After batch correction of the
394 Eutrema data, PCA of the control corrected and scaled data grouped according to drought-
395 stress duration for both species (Fig. 3). Moreover, PCA of unscaled data showed that most
396 of the variance is due to large differences between early-stress (Days 1, 3 and 5) and late-
397 stress (Days 12 and 13) observations. Statistical separation of late-stress effects was not
398 related to differing sample water content, since comparable dry weights were used and the
399 resulting data normalised before statistical analysis.

400 When considering only the early-stress observations, the PCA scores plot shows clear

401 grouping of the observations by stress duration for both plant species (Fig. 4). Distinctive
402 metabolic signatures were obtained even for early days with limited soil drying (< 20 %
403 change in SWC at Day 3). The PCA loadings identified variables that most influenced the
404 separation between Days 1, 3 and 5, and early- and late-stress observations. In addition, an
405 iterative k-means algorithm filtered out the largest clusters to leave those comprising more
406 unusual, and potentially more informative, patterns (supplementary Fig. S5). Hierarchical
407 clustering with the 46 time-series selected by the k-means analysis (Fig. 5; supplementary
408 Fig. S6) allowed the similarities (or differences) between the associated metabolites to be
409 visualised.

411 *Severe drought causes larger metabolic alterations in Arabidopsis than in Eutrema*

412 Late-stress markers for both Arabidopsis and Eutrema included peaks that were identified as
413 the carbohydrates sucrose and raffinose, by comparison with authentic standards of these
414 molecules. Sucrose significantly increased and raffinose significantly decreased ($p < 0.001$)
415 in late stress (Day 12) and on re-watering (Day 13). A significant decrease was found for
416 features with m/z values of 341 and 387, most probably a hexose disaccharide. A feature with
417 m/z 711, also decreasing significantly, is tentatively assigned to stachyose, known to co-elute
418 with raffinose (Antonio *et al.*, 2008). Sucrose (detected in a range of ionic forms) observed at
419 high levels in the leaf blade tissue (Table 1) was not unexpected. Drought stress upregulated
420 sucrose synthesis in different resurrection plants (Peters *et al.*, 2007; Whittaker *et al.*, 2007;
421 Gechev *et al.*, 2013) as well as in herbaceous and woody plants (e.g Antonio *et al.*, 2008;
422 Pinheiro and Chaves, 2011; Granda and Camarero, 2017). Although stress downregulated
423 raffinose synthesis in several resurrection plants (Muller *et al.*, 1997; Moyankova *et al.*,
424 2014), raffinose is typically described as being upregulated under stress (Elsayed *et al.*,
425 2014), including in Arabidopsis (Taji *et al.*, 2002). Soil water deficit significantly ($p <$
426 0.00001) decreased two co-eluting features (with m/z 191 and m/z 405) in both plant species
427 The feature with m/z 191 was assigned to citric acid, following tandem mass spectrometry
428 (MS^2) analysis and comparison of the fragmentation pattern in both METLIN
429 (www.metlin.scripps.edu) and PRIME (www.prime.psc.riken.jp) metabolomics databases.
430 The co-eluting feature at m/z 405.0019 on MS^2 produced a single fragment at m/z 191.0185,
431 that was tentatively assigned as the $[2M-2H+Na]^-$ charge-sharing dimer of citric acid
432 (accurate mass 405.0287). Univariate analyses (after multiple test correction) indicated that

433 607 variables significantly ($p < 0.0001$) differed between late-stress observations and controls
434 in Arabidopsis, in comparison to just 171 in Eutrema, suggesting that Arabidopsis more
435 extensively adjusts its metabolism. An alternative view is that larger changes in Arabidopsis
436 indicate less active metabolism, since metabolites accumulate because the plant has no
437 capacity to use them.

438 In the cluster analysis, three clusters tend to decrease over time, including the response of
439 raffinose (supplementary Fig. S5C, S5D, S5G), which was more extreme in Arabidopsis than
440 Eutrema, therefore occurring in a different cluster. Severe drought stress decreased the levels
441 of raffinose in lupins, although this was preceded by a transient increase during early stress
442 (Antonio *et al.*, 2008; Pinheiro *et al.*, 2011). Although the different ionic forms of citric acid
443 from both Arabidopsis and Eutrema group together in cluster D, a difference in the trend
444 between the two different plant species can be seen, with Eutrema showing an early increase
445 before the overall decrease. Citric acid decreased in response to late and severe drought, as
446 previously observed in lupin and Eutrema (Pinheiro *et al.*, 2004; MacLeod *et al.*, 2015). The
447 final two clusters (Fig. S5H and S5I) show the response profiles of (unknown) compounds
448 that are significantly greater than or lower than the controls throughout the time-series,
449 notably all from Arabidopsis, and are good candidates for further studies.

450 In contrast, four clusters tended to increase rapidly in late drought; the scale of the response
451 accounts for the difference between these four clusters. They mostly comprise the differing
452 ionic forms of sucrose. In Arabidopsis, unknown compounds with m/z 133 and m/z 288
453 exhibited a very similar pattern to sucrose (supplementary Fig. S5A, S5B, S5E and S5F). The
454 most extreme responses result in separate clusters consisting of just one or two observations
455 (Fig. S5E-F). For each sucrose ionic species, the response for Days 12 and 13 is more
456 extreme for Arabidopsis than for Eutrema. In both plant species, a further unknown with m/z
457 195 also clusters with sucrose, and re-watering causes a greater response than during late
458 stress.

459

460 ***Severe drought causes larger biomass reduction in Eutrema than in Arabidopsis***

461 Although severe drought decreased the biomass of both species, Arabidopsis (22% decrease)
462 was less sensitive than Eutrema (38% decrease) (supplementary tables S1 and S2). The
463 mechanisms that limit biomass accumulation are poorly understood (Pinheiro and Chaves,

464 2011; Skirycz *et al.*, 2011). Under drought, the higher availability of sugars indicates that
1
2 465 CO₂ assimilation is not limited as much as growth, with carbon being available but plants
3
4 466 unable to use it, termed “sink limitation” (Wiley and Helliker 2012; Granda and Camarero,
5
6 467 2017). This also supports the concept of passive accumulation. On the other hand, higher
7
8 468 sugar content may reflect their use in osmoregulation, maintaining cell integrity and
9
10 469 providing readily available carbon to resume growth (active reserve storage concept; Granda
11
12 470 and Camarero, 2017). Although osmotic adjustment was detected under severe drought and
13
14 471 rewatering in *Eutrema* (supplementary table S2), it was only detected in *Arabidopsis* on
15
16 472 rewatering (supplementary table S1). The growth reduction was accompanied by starch
17
18 473 remobilization, supporting the hypothesis of carbon reserve reallocation. A regulatory
19
20 474 mechanism that integrates carbon availability and its use within the plant (Smith and Stitt,
21
22 475 2007; Pinheiro and Chaves, 2011) partitions photoassimilates to other biochemical pathways
23
24 476 (than growth) to withstand severe drought and/or resume growth whenever possible. To
25
26 477 determine whether sucrose accumulation represents an active or passive process, it would be
27
28 478 necessary to determine if and how carbon limits growth (Wiley and Helliker 2012). A better
29
30 479 understanding of these mechanisms is crucial to select genotypes with more stable growth
31
32 480 under stress. In that sense, a favourable ideotype will depend on where the plant is to be
33
34 481 grown. Higher survival, largely due to conservative water consumption can be the most
35
36 482 relevant selection criterion in arid or semi-arid regions. On the other hand, in moderate
37
38 483 climates with milder droughts, plant production can be boosted if stress has little impact on
39
40 484 growth (Skirycz *et al.*, 2011; Tardieu, 2012). Higher stomatal conductance is a positive trait
41
42 485 when plants are under moderate/mild drought since it favors growth maintenance (Tardieu,
43
44 486 2012). In that scenario, *Arabidopsis* can provide a suitable model for biomass production
45
46 487 under moderate drought.

488

489 ***Severe drought decreased ascorbate content of both species***

490 Under severe stress, some parameters, including ascorbic acid (AscA), leaf chlorophyll
491
492 fluorescence (Fv/Fm) and chlorophyll a (Chla), have similar patterns in the two species.
493
494 Lower Fv/Fm indicates a lower photochemical harvest, with a larger proportion of photons
495
496 being diverted to non-photochemical quenching (heat and ROS production). Although two
497
498 hormones (IAA and SA) are proposed to up-regulate ROS production (Rivas-San Vicente
499
500 and Plasencia, 2011; Considine and Foyer, 2014; Khokon *et al.*, 2017), severe drought

496 significantly decreased SA content but did not alter IAA content. Thus, these hormones are
497 unlikely to promote ROS accumulation. Water deficit has variable effects on carotenoid
498 content (Koffler *et al.*, 2014; Uarrota *et al.*, 2018), but we did not detect quantitative
499 alterations. However, decreased chlorophyll a content indicates that chlorophyll degrades
500 faster than carotenoids (Lichtenthaler and Buschmann, 2001). A higher carotenoid to
501 chlorophyll content can favour photo-protection per amount of light received. On the other
502 hand, non-photochemical quenching via carotenoids requires ascorbate for correct function
503 (Koffler *et al.* 2014). Severe drought significantly decreased ascorbate content (43% in
504 *Eutrema*; 24% in *Arabidopsis*), which could have compromised such quenching. Noctor *et al.*
505 (2014) state that ascorbate content is only significantly decreased by drought when stress-
506 induced senescence processes are activated. Although the sampled leaves did not show
507 visible symptoms of senescence, their ascorbate levels suggest senescence programs were
508 already activated. Under drought, both *Arabidopsis Col-0* and *Col-0* mutants with lower
509 ascorbate content show senescence symptoms (Koffler *et al.*, 2014), with some mutant leaves
510 showing necrosis. Rewatering further decreased the ascorbate content (55% in *Eutrema*; 52%
511 in *Arabidopsis*), indicating the senescence program is still active. Changes in ascorbate
512 reduction status can also modulate signalling pathways, affecting stress-response mechanisms
513 (Considine and Foyer, 2014).

514 *Arabidopsis and Eutrema are metabolically distinct*

515 To characterize in more detail the responses to soil water depletion in *Arabidopsis* and
516 *Eutrema*, various biochemical parameters (Table 2) were measured during early drought.
517 Some metabolites showed minimal (< two-fold) differences between species, while starch, JA
518 and ZR were more abundant in *Arabidopsis*, and IAA and DHA were more abundant in
519 *Eutrema* (supplementary tables S1 and S2). Distinctive phytohormone profiles n under non-
520 challenging conditions were previously reported (Arbona *et al.*, 2010; Pilarska *et al.*, 2016).

521 To remove the effects of differential growth between species, data were control-corrected by
522 subtracting the median control value from the measurements for each day. PCA analysis with
523 all biochemical parameters for both species (supplementary Fig. S7) showed the greatest
524 source of variance to be the separation of late/severe drought and re-watered (RW)
525 observations, as in the untargeted analyses. Without variable scaling, loadings plots showed a
526 large influence of the variables with the greatest mean values (leaf RWC, osmotic potential

528 (OP) and starch) in the total variance. After scaling to unit-variance, the separation of late
529 stress/re-watered observations is still seen along the first principal component, although
530 accounting for far less of the total variance. In Arabidopsis, variables from re-watered
531 samples were closer to those from early-day observations. In Eutrema, the difference between
532 late stress and re-watering is only apparent along the second component, which represents
533 less variance and more similar metabolic status. These findings suggest: 1) Arabidopsis
534 responds faster to soil water availability; and/or 2) Eutrema requires prolonged stimulus to
535 reprogram its metabolism. However, whole plant water use of Eutrema is more conservative
536 than Arabidopsis (with soil drying decreasing water use after 3 and 6 days respectively – Fig
537 1) in spite of an opposite stomatal response (with soil drying decreasing g_s after 4 and 6 days
538 in Arabidopsis and Eutrema respectively – Fig. 2). Thus vegetative growth and stomatal
539 closure seem independently regulated, with Eutrema reacting faster to soil water availability
540 and Arabidopsis requiring prolonged stimulus. The “optimistic” strategy of Arabidopsis Col-
541 0 maintains biomass production under mild stress and/or under deficit irrigation (Skirycz *et*
542 *al.*, 2011). An alternative hypothesis could be that Eutrema slows its metabolism much earlier
543 while Arabidopsis reprograms metabolism in a different way. It will be important to
544 determine whether growth is maintained, both above and below ground, and if reserves are
545 reallocated.

546 547 *Eutrema responds with small changes to early drought*

548 In both Arabidopsis and Eutrema, inspection of the PCA loadings showed that many
549 variables contribute to the separation of each of the early stress days (Fig. 4). Thus metabolic
550 separation between sampling dates is due to the cumulative changes arising from small
551 contributions of many metabolites. However, the two species react differently to similar
552 decrease in the soil water availability. More metabolites responded to early drought stress in
553 Arabidopsis, with 428 variables showing statistically significant differences between Days 1,
554 3 and 5 ($p < 0.01$; 36 with $p < 0.001$) in comparison to 35 in Eutrema ($p < 0.01$; 4 with $p <$
555 0.001). However, none of the variables that consistently differed between the early days
556 corresponded to those identified as late-stress markers (such as sucrose), showing different
557 metabolism during early and late drought.

558 The larger changes in Arabidopsis suggest different metabolic strategies to deal with the
559 progressive decline in soil water availability. This is consistent with greater stomatal closure

560 of *Arabidopsis* than *Eutrema* on Day 5 (Fig. 2), supporting a drought avoidance strategy for
561 *Arabidopsis*. Partial stomatal closure can limit carbon assimilation and carbon availability for
562 respiration and growth (Pinheiro and Chaves, 2011). Acclimation of *Arabidopsis* to optimise
563 carbon use (Smith and Stitt, 2007) would increase carbon storage (e.g. via changes in starch
564 synthesis and turnover) and adjust the growth rate. It could be that the changing conditions
565 did not achieve the threshold required to induce the acclimatory response as, although
566 stomata were starting to close, plant water status was not yet affected. A similar rationale can
567 be used for *Eutrema* although its perception as a drought avoider (MacLeod *et al.*, 2015),
568 suggests a different soil water availability threshold for acclimation. Transpiration data
569 indicate more conservative water use in *Eutrema* than *Arabidopsis* (Fig. 1) although
570 *Arabidopsis* had greater stomatal sensitivity to drying soil (Fig. 2). Together, these data
571 suggest species differences in regulating water consumption, implying distinct integration of
572 environmental signals to limit vegetative growth in *Eutrema* and induce stomatal closure in
573 *Arabidopsis*

574

575 *Species-dependent hormonal responses during early stress*

576 ABA, JA, SA and GA profiles clearly discriminated the plant species between Days 1 and 5
577 (Table 3; Fig. 5 and supplementary Fig. S8). Despite daily irrigation to ensure a similar rate
578 of soil drying in the two species, soil water deficit increased foliar ABA content of
579 *Arabidopsis*, but not *Eutrema*, on Day 5 (Fig. 6). Although daily irrigation increases leaf
580 water status and decreases ABA accumulation of plants exposed to reduced soil water
581 availability (Puértolas *et al.*, 2017), significantly higher water consumption of *Arabidopsis*
582 between Days 3 and 7 (Fig. 1) enhanced foliar ABA accumulation, potentially mediating
583 stomatal closure. Nevertheless, temporal decoupling of foliar ABA accumulation from
584 stomatal closure (Pinheiro *et al.*, 2011; Bechtold *et al.*, 2016) suggests that ABA
585 quantification at the guard cell level (Harris and Outlaw, 1991) is needed to better understand
586 the regulation of stomatal conductance. Alternatively, direct hydraulic regulation of stomatal
587 conductance (that was not detected in RWC measurements), or water-deficit stimulation of
588 localised foliar ABA accumulation (that was not detected in bulk leaf ABA measurements)
589 provide alternative hypotheses for stomatal closure.

590 Also, increased JA content may stimulate stomatal closure (Daszkowska-Golec and Szarejko,
591 2013), either independently or in association with ABA accumulation (de Ollas and Dodd,

592 2016). Foliar JA content transiently increased on Day 3 only in Arabidopsis, preceding
1 593 increased ABA accumulation on Day 5 (Fig. 6, supplementary Fig. S8). Transient increases
2
3 594 in foliar JA accumulation preceded ABA accumulation in *Solanum lycopersicum* leaves
4
5 595 (Muñoz-Espinoza *et al.*, 2015) and Arabidopsis (Arbona *et al.*, 2010). Thus, transient JA
6
7 596 accumulation may be necessary to induce ABA accumulation thereby contributing to
8
9 597 stomatal closure.

10
11 598 Similarly to JA, SA content transiently increased on Day 3 in Arabidopsis (Fig. 6). In
12
13 599 Arabidopsis, SA promotes ROS production and accumulation, and activates S-type anion
14
15 600 channels in guard cells involved in stomatal closure (Khokon *et al.*, 2017). SA-mediated
16
17 601 stomatal closure is considered to be ABA-independent, but a positive cross-talk is also
18
19 602 accepted (Rivas-San Vicente and Plasencia, 2011; Miura and Tada, 2014). On the other hand,
20
21 603 SA has also been proposed to modulate redox homeostasis, by regulating antioxidant
22
23 604 enzymes (Rivas-San Vicente and Plasencia, 2011).

24
25 605 In Eutrema, changes in leaf RWC and ABA occurred after Day 5, with foliar ABA
26
27 606 accumulation in Eutrema occurring below 45 % SWC. Species differences could be
28
29 607 associated with the osmotic potential (OP) and the redox state regulation, as significant
30
31 608 changes were observed in Eutrema, but not in Arabidopsis (Fig. 7 and supplementary Fig.
32
33 609 S8). Decreased OP in Eutrema at Day 3 may maintain turgor, thereby removing the stimulus
34
35 610 for ABA synthesis (Sack *et al.*, 2018). The opposing trends seen in AscA and DHA for Days
36
37 611 3 and 5 in Eutrema may induce signalling patterns that prevented ABA accumulation. In
38
39 612 Arabidopsis, ABA increased at Day 5, but there were no significant changes in AscA or DHA
40
41 613 until Day 5. Diminished stomatal closure of Eutrema may be related to its limited metabolic
42
43 614 response to soil drying (Fig. 2), since hormonal activation of stomatal closure may require
44
45 615 specific metabolic thresholds to be achieved. [On the other hand, since Eutrema limits whole](#)
46
47 616 [plant water consumption earlier than Arabidopsis, its limited metabolic response can be](#)
48
49 617 [related to slower metabolism, reflecting stress avoidance \(Tardieu, 2012\).](#) Altered GA
50
51 618 metabolism also supports the hypothesis that Arabidopsis responds differently than Eutrema
52
53 619 to soil water availability. Two precursors of the bioactive GA4 (GA24, GA9; Fig. 6&7, Table
54
55 620 3) showed altered profiles in Arabidopsis but not in Eutrema; with increased GA24 and GA9
56
57 621 contents at Day 5 indicating GA4 deactivation, a growth inhibitory signal.

58 622 Despite similar stomatal response to drying soil, phytohormonal responses substantially
59
60 623 differed between the two species. The multiplicity of possible stomatal effectors suggests that

624 the most parsimonious stomatal regulation is via leaf hydraulics (Tardieu, 2016), yet until
625 Day 5 there was scant evidence that soil drying altered leaf water status, at least in *Eutrema*.
626 Redundancy in stomatal regulation between closely-related species may confer an
627 evolutionary advantage under specific environmental conditions.

628

629 *Severe drought and rewatering induce opposite ACC and CK profiles in the two species*

630 With the exception of SA, hormone responses to soil drying are quite distinct in the two
631 species (Table 2). Severe drought increased content of the ethylene precursor ACC by 70 %
632 in *Eutrema*, but had no effect in *Arabidopsis*, suggesting ethylene-independent stomatal
633 closure as both species showed similar stomatal sensitivity to drying soil. In contrast, re-
634 watering *Eutrema* returned ACC levels to well-watered values, while profoundly increasing
635 ACC content in *Arabidopsis*. Enhanced foliar ethylene emission following re-watering has
636 been attributed to additional root ACC export (Gomez-Cadenas *et al.*, 1996) and may be
637 involved in citrus leaf abscission. Moreover, assuming these increases in foliar ACC content
638 are coincident with enhanced ethylene emission, they may also be involved in growth
639 regulation.

640 Several CK species including ZR and 2-iP, long distance translocation forms of CKs (Kieber
641 and Schaller, 2014), as well as the 2-iP precursor IPA accumulated in *Arabidopsis* but not in
642 *Eutrema* during late stress (Fig. 8; supplementary Fig. S9). In contrast, re-watering returned
643 content of these CKs to well-watered values in *Arabidopsis*, while stimulating their
644 accumulation in *Eutrema*. IPA and 2-iP are precursors of Z, one of the most active CK forms
645 (Jameson, 2017). However, the mobilization (metabolism and/or transport) of these CKs in
646 *Arabidopsis* was not reflected in higher Z levels. Decreased levels of bioactive CKs due to
647 severe and prolonged drought stress have been associated with better performance under
648 drought, in mutants with decreased levels of bioactive CKs achieved via overexpression of
649 *CKX* genes or by inactivating *IPT* genes (Ha *et al.*, 2012). Since these mutant lines show
650 reduced growth under optimal conditions, it can be argued that their water requirements are
651 lower than those of the WT. However, while lower evapotranspiration is described for some
652 *CKX* mutants (Farber *et al.*, 2016), *ipt* mutants show similar water consumption (Nishiyama
653 *et al.*, 2011). On the other hand, senescence-induced *IPT* overexpression, maintained
654 bioactive CK content as the soil dries (Rivero *et al.*, 2007; Xu *et al.*, 2017), without reducing
655 growth (Rivero *et al.*, 2007). Nevertheless, re-watering increased bioactive CKs in these

1 656 drought-tolerant transgenics (Rivero *et al.*, 2007). Although differential CK profiles may
2 657 mediate plant drought stress responses, the two species showed similar stomatal sensitivity to
3
4 658 soil drying and re-watering, suggesting CK-independent stomatal regulation.
5

6 659

7 8 660 **Conclusions**

9
10 661 Arabidopsis and Eutrema responded physiologically (stomatal closure) and metabolically
11 662 with temporal differences to slowly imposed drought. Following soil drying (Days 1 to 5),
12
13 663 Arabidopsis rapidly increased JA and SA content (Day 3) and later increased ABA levels
14
15 664 (Day 5), whereas in Eutrema, ABA levels did not change until Day 5. Untargeted metabolite
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17 665 analysis demonstrated larger responses in Arabidopsis than in Eutrema. In contrast, greater
18
19 666 soil drying was needed to initiate partial stomatal closure in Eutrema (Day 6), although
20
21 667 decreased water use (when compared to controls) was observed earlier than in Arabidopsis.
22
23 668 We hypothesise that growth inhibition is a first response to water deficit in Eutrema. With
24
25 669 severe and prolonged drought, conserved metabolic responses (increased sucrose and
26
27 670 decreased raffinose and citric acid in both species) co-occurred with near-complete stomatal
28
29 671 closure in both species. The different physiological and metabolic responses and their timing
30
31 672 allow these species to utilise alternative pathways to physiologically adjust to soil drying,
32
33 673 likely reflecting adaptations to their respective niches.
34

35 674

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Figure Legends

Figure 1. Soil water content (SWC, %) after imposing water deficit and on re-watering (shaded area). SWC is shown before (symbols) and after (dotted lines) partial water replacement (to regulate the rate of soil water depletion). Data show the means \pm standard error of 6 pots (except Day 1 with 5 pots). For pre-irrigation SWC, significance levels were calculated using the Mann–Whitney U test. Data differing significantly from Day 0 are denoted by asterisks (* $p < 0.05$, ** $p < 0.01$, *** $p < 0.001$).

Figure 2. Leaf stomatal conductance (as a % of the control plants) plotted against SWC. Mean values (of 3 to 5 biological replicates) are shown with only positive standard errors for clarity. Significant results, as determined by Mann-Whitney U test, are denoted by asterisks (* $p < 0.05$, ** $p < 0.01$, *** $p < 0.001$). ANCOVA for each main effect (treatment and species) and their interaction is presented in supplementary Fig S4B.

Figure 3. PCA plots showing the scores for the first two principal components obtained for the untargeted metabolomic analysis coloured by experimental group and the day of harvest, for (A) Arabidopsis and (B) Eutrema. For both plant species, the data have been scaled to unit variance and control corrected. In the case of Eutrema only, batch correction has also been performed.

Figure 4. PCA scores plots for the first two principal components obtained from scaled early-stress observations (Days 1, 3 and 5) in the untargeted analysis after control correction for (A) Arabidopsis and (B) Eutrema. The observations are coloured according to the day of harvest, showing that the clustering of observations is related to drought duration.

Figure 5. Dendrogram obtained from hierarchical clustering of the 46 time-series selected by the iterative k-means analysis of the metabolite data. The clusters are coloured and annotated A-I according to the clusters identified in the k-means analysis (supplementary Fig S5). Metabolites within clusters are labelled as follows: S= sucrose; R = raffinose; St = stachyose; CA = citric acid; U = unassigned hexose disaccharide.

945 Figure 6. Biochemical parameters with a statistically significant change in early drought
1 946 stress in Arabidopsis but not in Eutrema. The mean difference from well-watered plants for
2 947 leaf RWC and the hormones ABA, JA, SA and GA24 are shown with error bars representing
3 948 the standard error. The mean values after control correction (i.e. the mean value for the
4 949 controls has been subtracted) are represented. Arabidopsis: dark grey; Eutrema: light grey.
5 950 **Significant results** are shown in Table 3.

951 Figure 7. Biochemical parameters with a significant change in early stress in Eutrema but not
12 952 in Arabidopsis. The mean measurement for osmotic potential, DHA, AscA, 2iP and GA9 are
13 953 shown with error bars representing the standard error of the observations. The mean values
14 954 after control correction (i.e. the mean value for the controls has been subtracted) are
15 955 represented. Arabidopsis: dark grey; Eutrema: light grey. **Significant results** are shown in
16 956 Table 3.

957 Figure 8. Cytokinins during early (Days 1, 3, 5) and late (Day 12) stress and on re-watering
24 958 (Day 13). Mean values and \pm standard error of 6 biological replicates (except for Day 1
25 959 where $n=5$). The mean values after control correction (i.e. the mean value for the controls has
26 960 been subtracted) are represented. In Arabidopsis, ZR, 2iP and IPA peak at late stress and
27 961 decrease on re-watering. However, in Eutrema, these hormones show a slight decrease in late
28 962 stress and increase dramatically on re-watering. Arabidopsis: dark grey; Eutrema: light grey.
29 963 **Significant results** are shown in Table 2.

37 964

965 **Supplementary table S1.** Biochemical parameters for both control (WW) and stressed (WD)
1 966 observations in Arabidopsis. Data are the means \pm standard error of 6 biological replicates,
2 967 except for day 1 (n = 5). Asterisks in the third row show parameters with a significant
3 968 difference between WW and WD for a particular day (obtained using Mann-Whitney tests).
4 969 Asterisks in the final column for each day show days that are significantly different from
5 970 earlier days (using Tukey's HSD test) with the specific days given in parentheses. Here,
6 971 asterisks denote *** $p < 0.001$, ** $p < 0.01$ and * $p < 0.05$.

972 **Supplementary table S2.** Biochemical parameters for both control (WW) and stressed (WD)
13 973 observations in Eutrema. Data are the means \pm standard error of 6 biological replicates,
14 974 except for day 1 (n = 5). Asterisks in the third row show parameters with a significant
15 975 difference between WW and WD for a particular day (obtained using Mann-Whitney tests).
16 976 Asterisks in the final column for each day show days that are significantly different from
17 977 earlier days (using Tukey's HSD test) with the specific days given in parentheses. Here,
18 978 asterisks denote *** $p < 0.001$, ** $p < 0.01$ and * $p < 0.05$.

979 **Supplementary table S3.** Biochemical parameters for both control (WW) and stressed (WD)
27 980 observations in Arabidopsis. Samples were control-corrected (see Methods section). Data
28 981 shown are the means \pm standard error of 6 biological replicates, except for day 1 (n = 5).

982 **Supplementary table S4.** Biochemical parameters for both control (WW) and stressed (WD)
34 983 observations in Eutrema. Samples were control corrected (see). Data shown are the means \pm
35 984 standard error of 6 biological replicates, except for day 1 (n = 5).

985 **Supplementary Figure S1.** Preliminary drought assay. **A** - Soil water content (SWC, %)
41 986 progression during the assay for Eutrema and Arabidopsis; **B** - Leaf stomatal conductance (%
42 987 of the control gs) as a function of the SWC. For controls, percentage gs was calculated
43 988 relative to day 0; for treatments, percentage gs was calculated relative to the control for the
44 989 same day. The 80 % gs level was achieved on different days: by day 4 in Arabidopsis and by
45 990 day 6 in Eutrema; **C** - Regression line fit % gs vs soil water content. Each point represents a
46 991 single measurement and p-values were determined by ANCOVA for each main effect
47 992 (treatment and species) and their interaction (ns: not significant; #: $p < 0.1$; ***: $p < 0.001$).

993

994 **Supplementary Figure S2.** PCA plots showing the scores for the first two principal
1
2 995 components obtained for the Eutrema data after scaling to unit variance with the observations
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4 996 coloured by batch. **A-** Before batch correction, clustering within batches can be seen and, in
5
6 997 particular, batches 7 and 8 cluster separately. **B-** After batch correction, differences between
7 998 batches are no longer apparent.

9
10 999 **Supplementary Figure S3.** PCA scores for the first two principal components obtained for
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12 1000 the Arabidopsis data after scaling to unit variance. The observations are coloured by data
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14 1001 collection batch and no obvious differences between batches can be seen, so that batch
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16 1002 correction is not necessary.

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18 1003 **Supplementary Figure S4. A -** Leaf stomatal conductance of Arabidopsis and Eutrema after
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20 1004 imposing water deficit and on re-watering (shaded area); **B -** Regression line fitting % gs vs
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22 1005 soil water content. Each point represents a single measurement and p-values were determined
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24 1006 by ANCOVA for each main effect (treatment and species) and their interaction (ns: not
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26 1007 significant; ***: $p < 0.001$).

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29 1008 **Supplementary Figure S5.** The nine clusters obtained with k-means analysis of the 46 time-
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31 1009 series remaining after iterative filtering of the metabolite data. Cluster A and cluster B
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33 1010 include several sucrose species. Cluster C includes raffinose and cluster D includes citric
34 1011 acid.

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37 1012 **Supplementary Figure S6.** Heatmap showing the similarity of the 46 time-series selected by
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39 1013 iterative k-means analysis of the metabolite data. Metabolites are labelled as follows: S=
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41 1014 sucrose; R = raffinose; St = stachyose; CA = citric acid; U = unassigned hexose disaccharide.

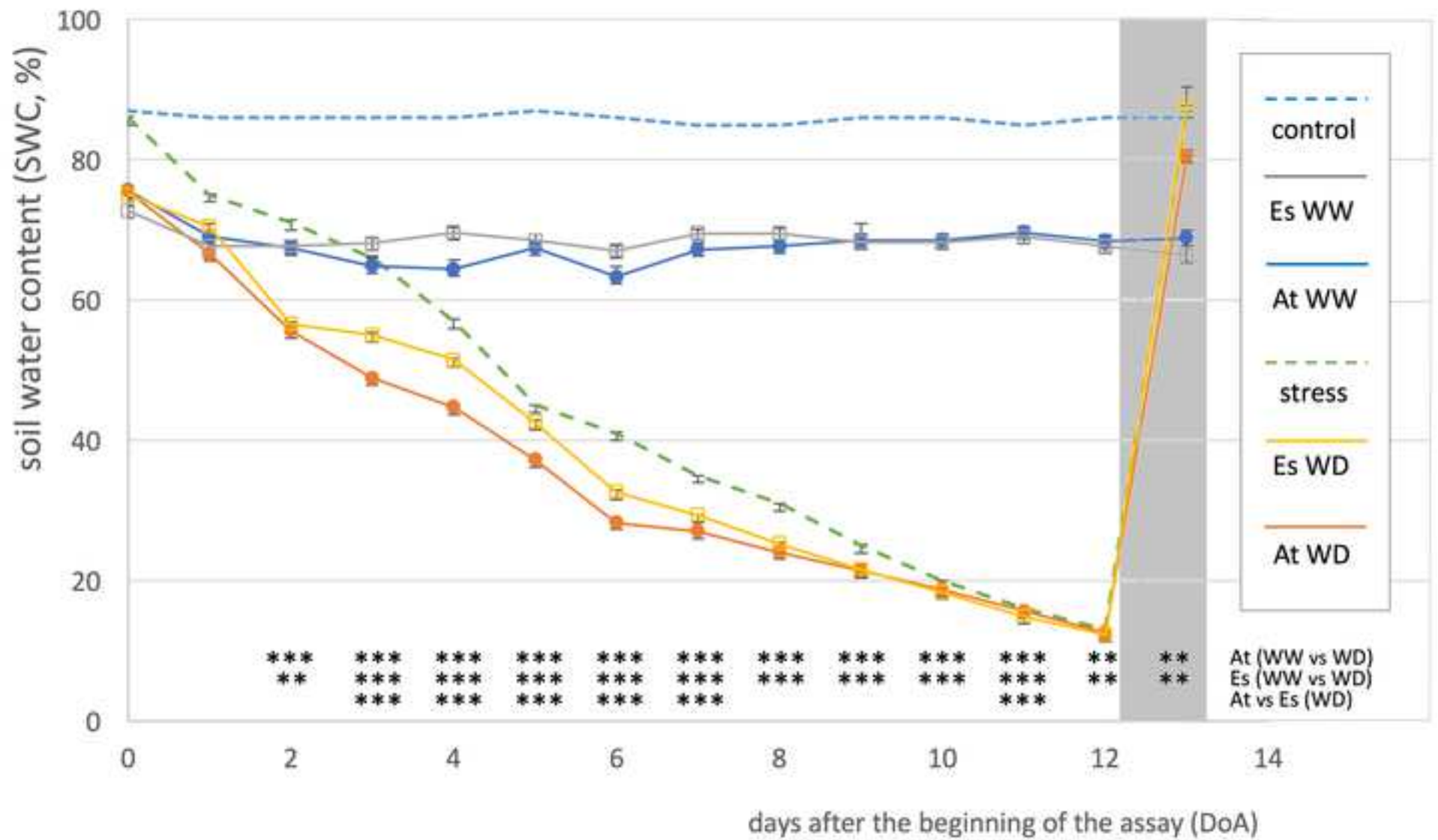
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44 1015 **Supplementary Figure S7.** PCA plots of the biochemical parameters for both control (WW)
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46 1016 and treatment (WD) observations in Arabidopsis and Eutrema after control correction. (A)
47 1017 unscaled variables; (B) scaled variables.

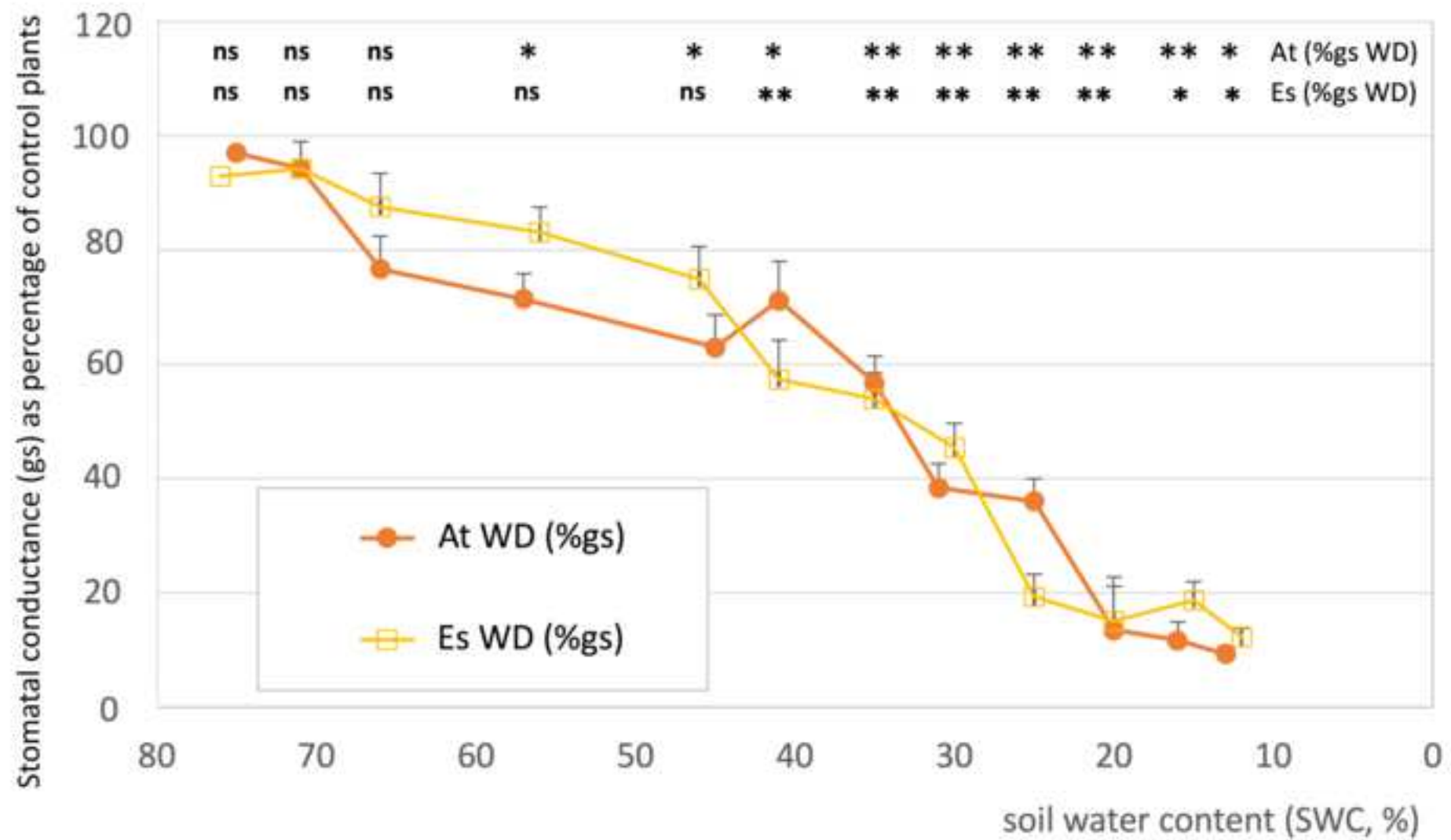
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50 1018 **Supplementary Figure S8.** Line plots showing physiological and biochemical parameters in
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52 1019 early-drought stress (days 1, 3 and 5) after control correction. Error bars show the standard
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54 1020 error between observations ($n = 6$ biological replicates, except for day 1, $n = 5$). Dark grey:
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56 1021 Arabidopsis; light grey: Eutrema. ANOVA results are presented in Table 3.

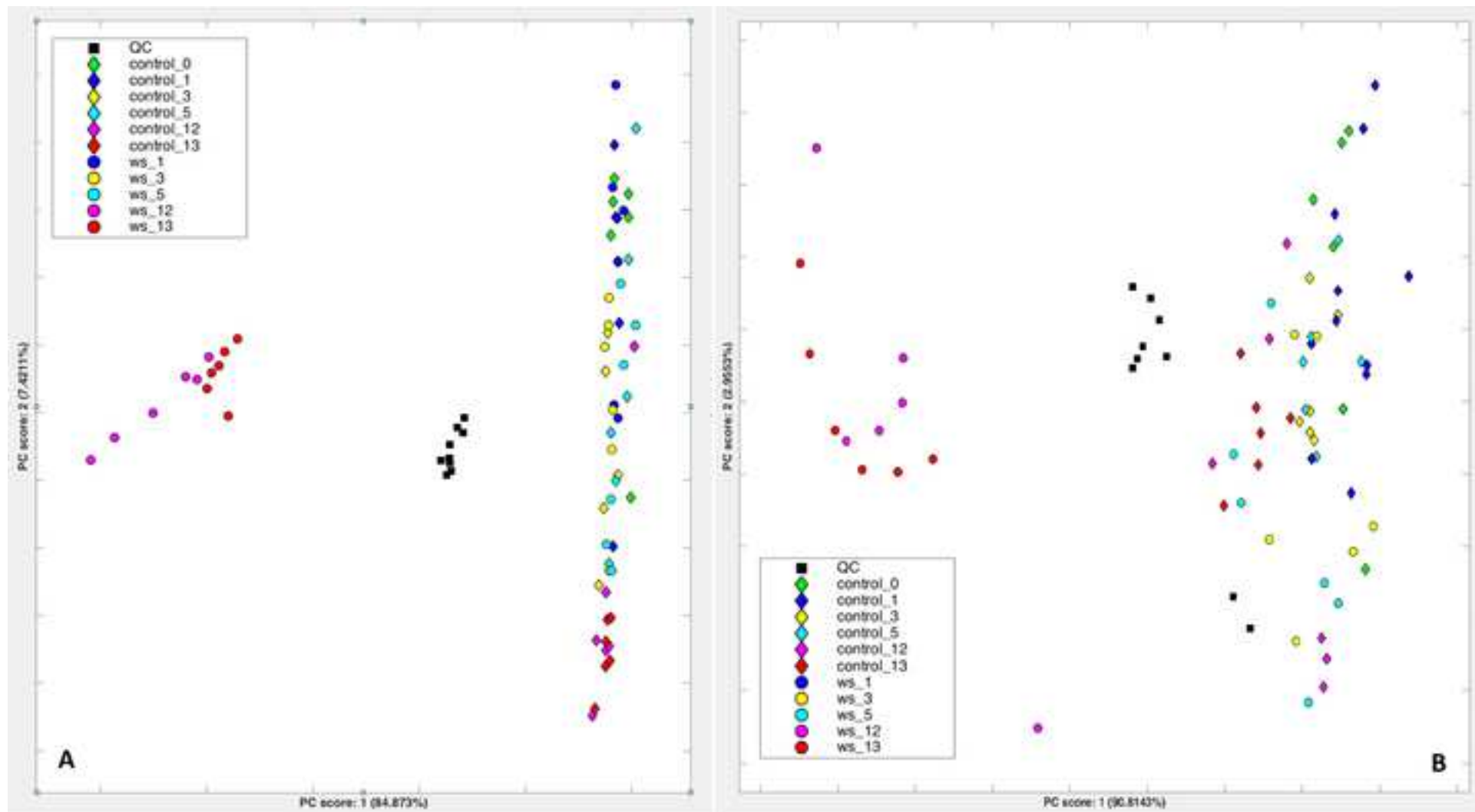
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58 1022 **Supplementary Figure S9.** Bar charts showing physiological and biochemical parameters in
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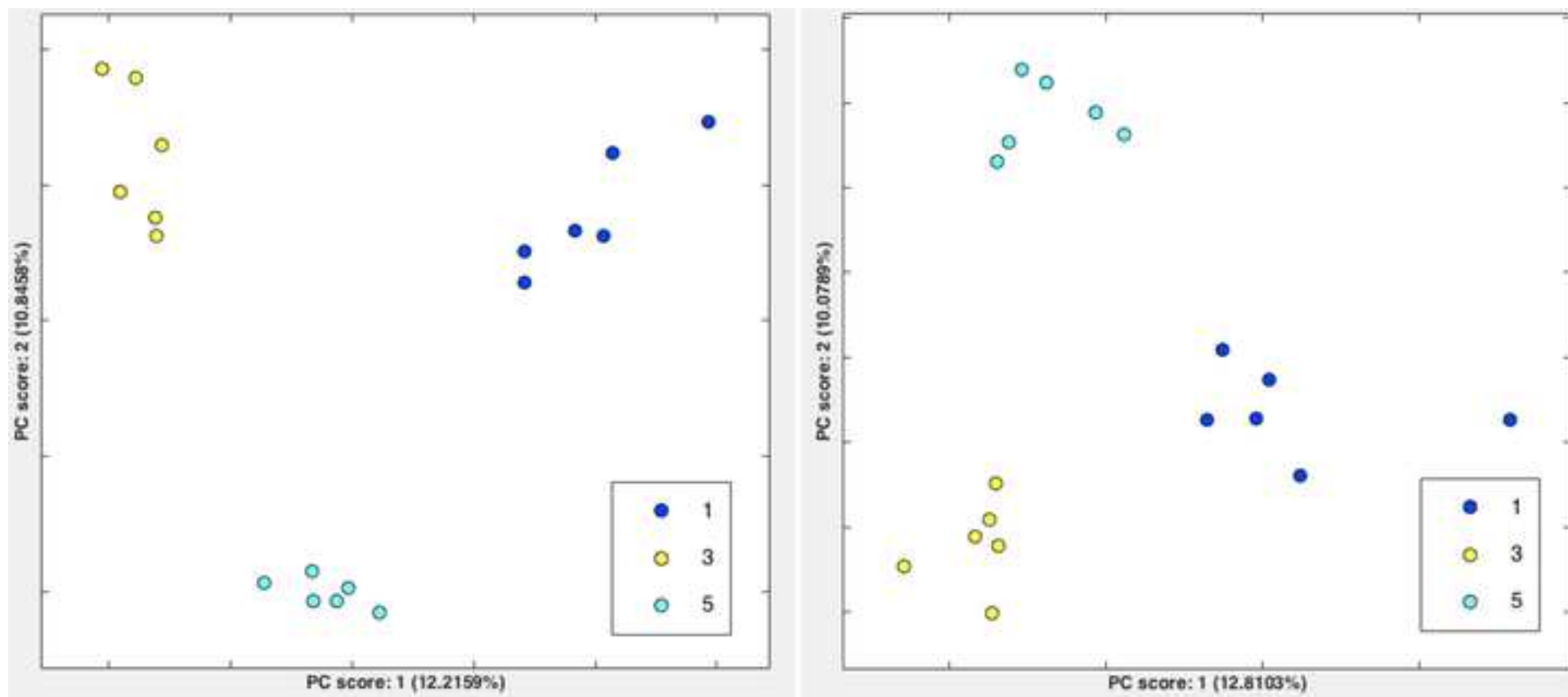
1023 early- (days 1, 3 and 5) and late-drought stress and on re-watering (day 13) after control
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2 1024 correction. Error bars show the standard error between observations (n = 6 biological
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4 1025 replicates, except for day 1, n = 5). Dark grey: Arabidopsis; light grey: Eutrema. ANOVA
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6 1026 results are presented in Table 2.

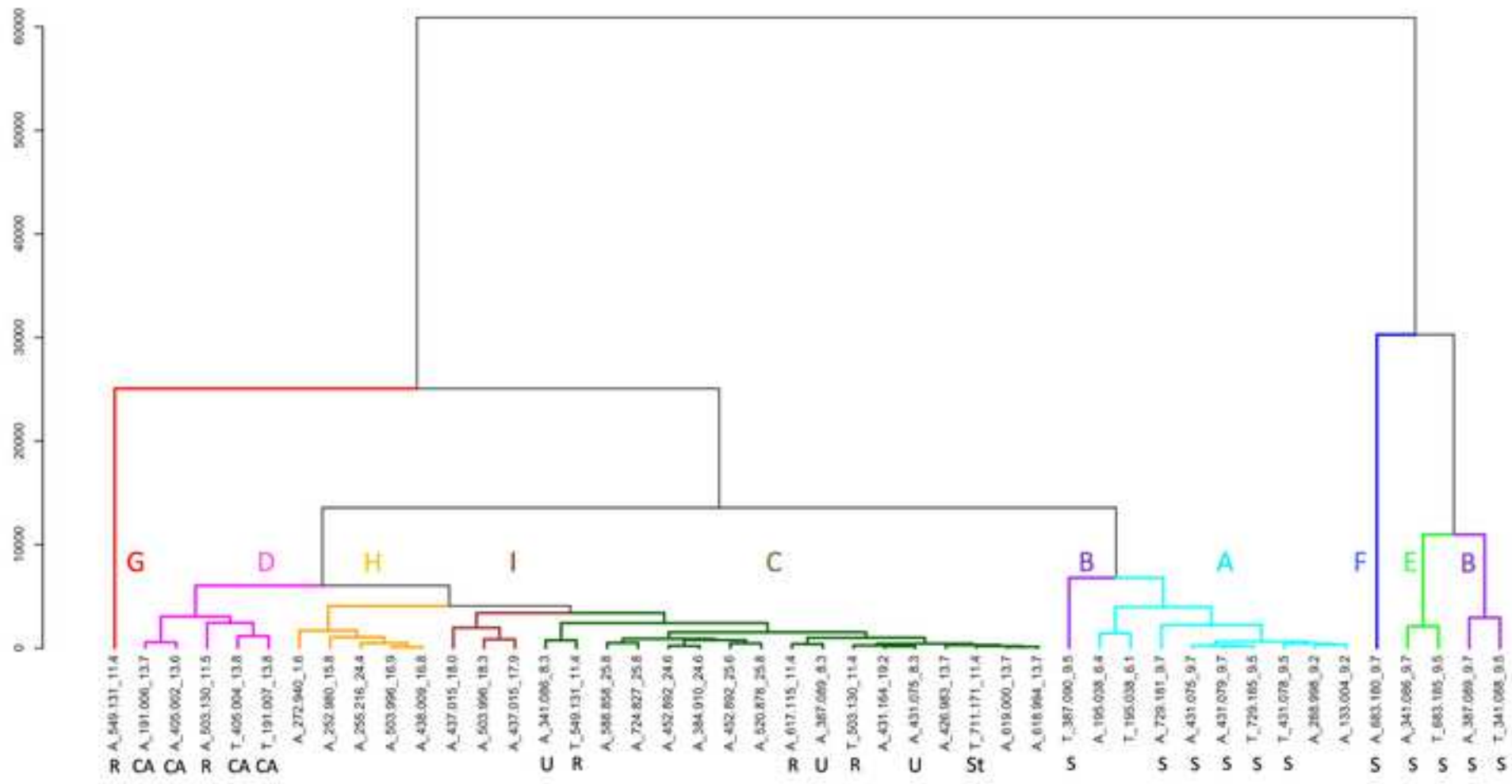
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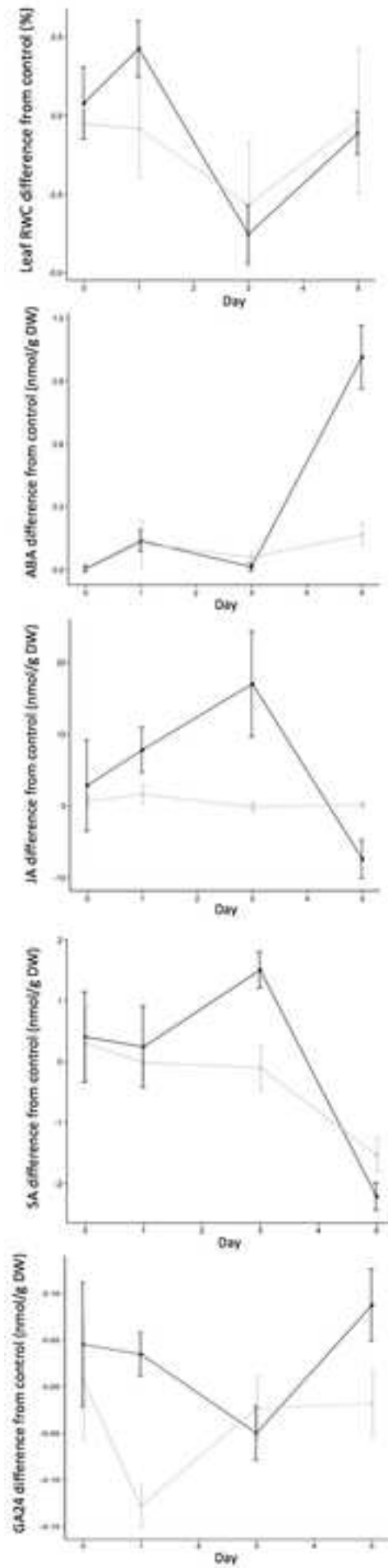


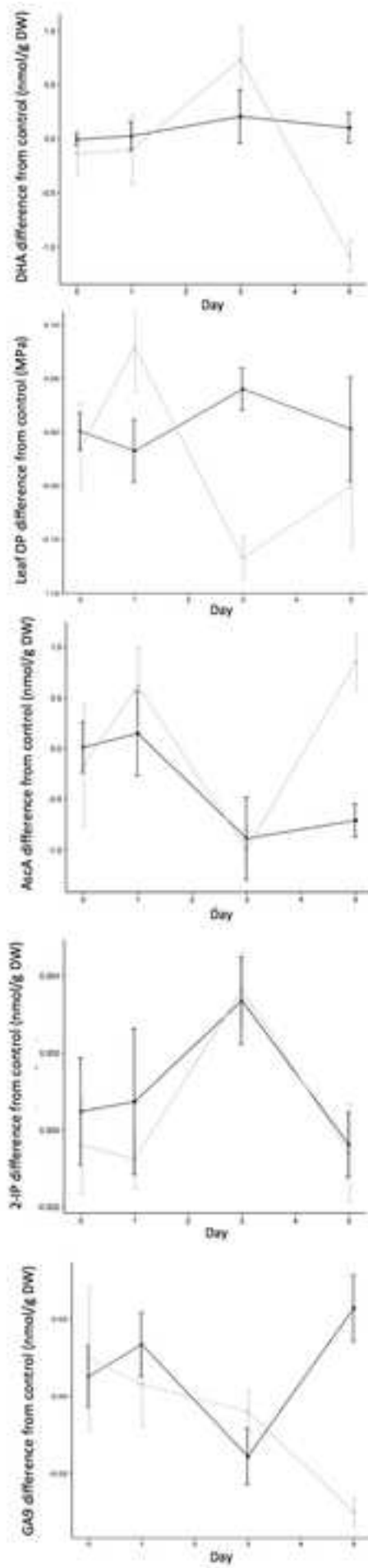












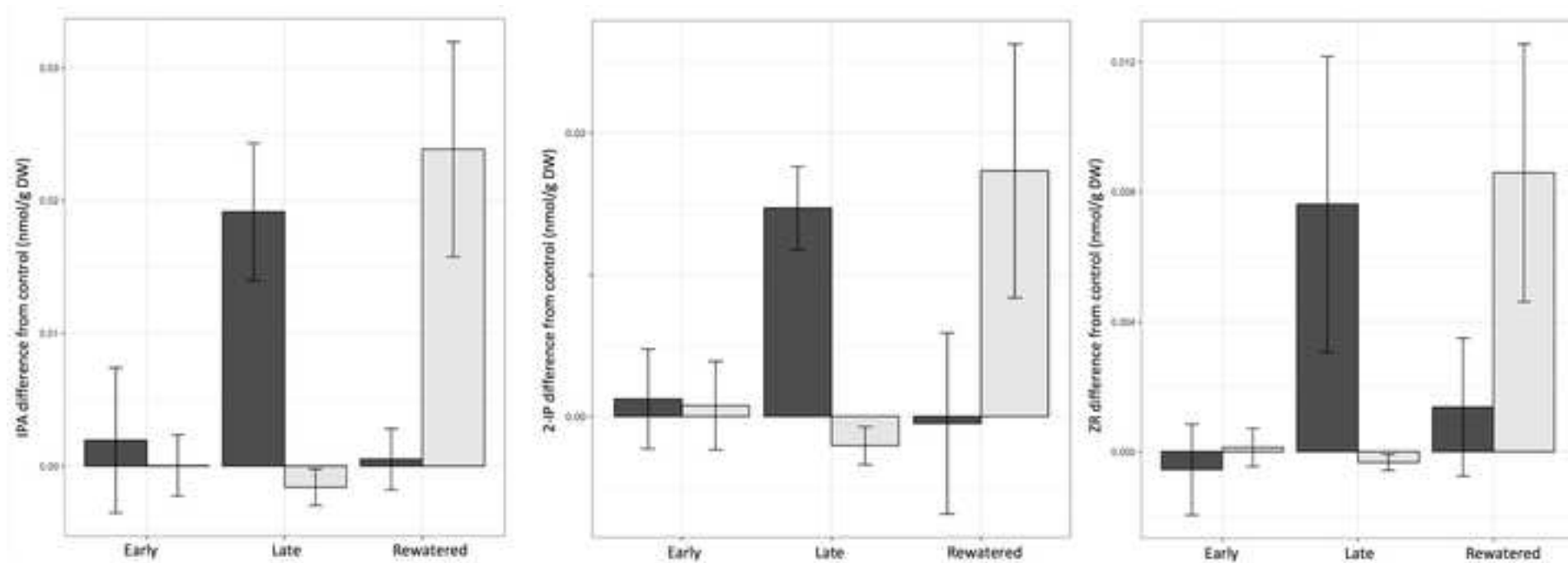


Table 1: Molecular forms assigned to sucrose, raffinose and citric acid that were significantly different ($p < 0.001$) between late drought-stressed and rewatering observations (days 12 or 13) and the corresponding controls.

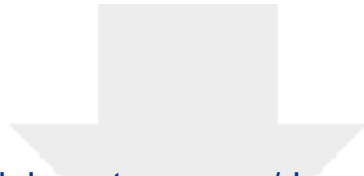
Compound	Molecular form	<i>m/z</i>
sucrose	$[M-H]^-$	341.087
sucrose	$[M+HCOO]^-$	387.092
sucrose	$[M-2H+2Na+HCOO]^-$	431.092
sucrose	$[2M-H]^-$	683.181
sucrose	$[2M-HCOO]^-$	729.186
raffinose	$[M-H]^-$	503.126
raffinose	$[M-HCOO]^-$	549.131
raffinose	$[M+HCOONa+HCOO]^-$	617.119
citric acid	$[M-H]^-$	191.006
citric acid	$[2M-2H+Na]^-$	405.002

Table 2. Representation of ANOVA results after Tukey's HSD correction for pairwise testing between early stress (combined days 1, 3 and 5), late stress (day 12) and re-watered (day 13) observations. Separate ANOVA models were obtained for each species. Light green represents $p < 0.05$, mid-green $p < 0.01$ and dark green $p < 0.001$. White cells indicate no significant difference at the 95 % confidence level. Carotenoids, DHA, IAA, DHZ DHZR, GA19, GA20 and GA24 were measured, but omitted from the table as no significant difference between groups was found for either plant species. Arrows denote an increase (↑) or decrease (↓) in measurement.

Variable	Species	Early - late	Late - rewatered
Biomass	At	↓	
	Th	↓	
Root RWC	At	↓	↑
	Th	↓	↑
Leaf RWC	At	↓	↑
	Th	↓	↑
Starch	At	↓	
	Th	↓	↑
OP	At	↓	
	Th	↓	
OA	At		↓
	Th	↑	↓
OP100	At		↓
	Th	↑	↓
PSII	At	↓	
	Th	↓	↑
Chla	At	↓	↑
	Th	↓	
Chlb	At		
	Th		↓
AscA	At	↓	↓
	Th	↓	
ABA	At		
	Th	↑	
IAA	At		
	Th		
IPA	At	↑	↓
	Th		↑
2-IP	At	↑	↓
	Th		↑
ZR	At	↑	↓
	Th		↑
Z	At		
	Th		↑
DHZ	At		
	Th		
JA	At	↓	
	Th	↓	
SA	At	↓	↓
	Th	↓	↓
ACC	At		
	Th	↑	↓

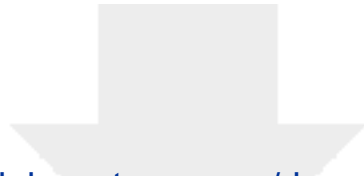
Table 3. Representation of ANOVA results after Tukey's HSD correction for pairwise testing between early stress observations. Light green represents $p < 0.05$, mid-green $p < 0.01$ and dark green $p < 0.001$. White cells indicate no significant difference at the 95% confidence level. Variables that were measured, but for which no significant difference between groups was found for either plant species were omitted from the table. Arrows denote an increase (↑) or decrease (↓) in measurement. Plots are shown in Fig. 7 and 8 and supplementary Fig. S7.

Variable	Species	Day 1 - day 3	Day 3 - day 5
Leaf RWC	At	↓	↑
	Th		
ABA	At		↑
	Th		
JA	At		↓
	Th		
SA	At		↓
	Th		↓
OP100	At	↑	
	Th		
OP	At		
	Th	↓	
DHA	At		
	Th		↓
AscA	At		
	Th	↓	↑
2-IP	At		
	Th	↑	↓
GA24	At		↑
	Th		
GA9	At	↓	↑
	Th		

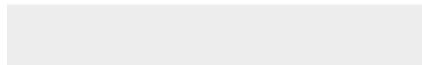


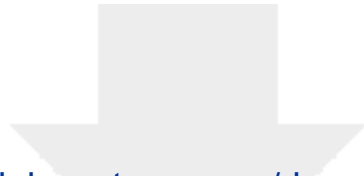
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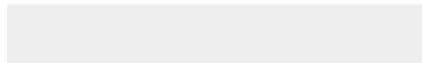


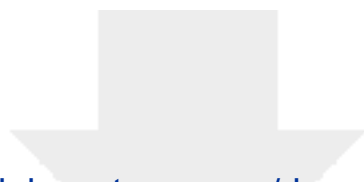
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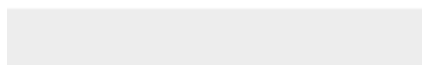
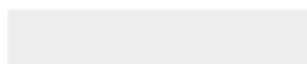


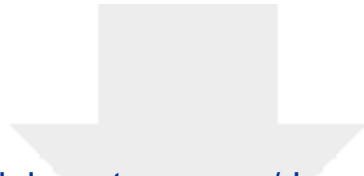
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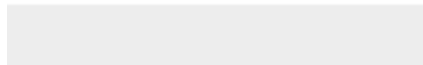


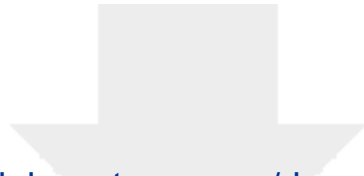
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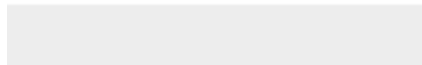
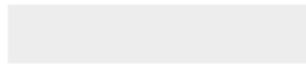


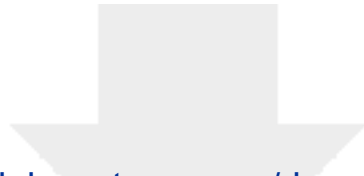
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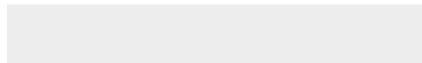
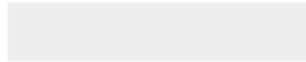


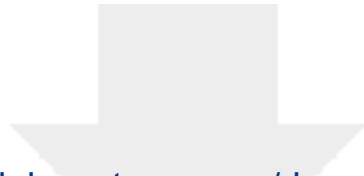
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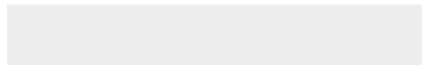


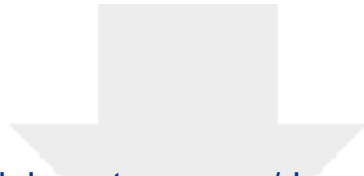
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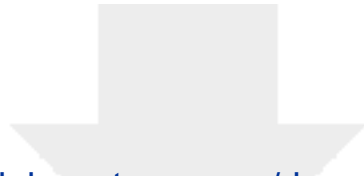
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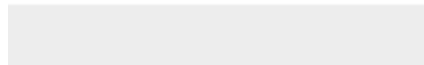
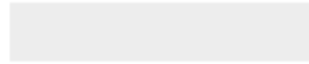


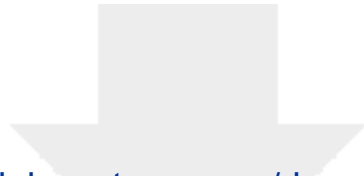
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