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Metabolic plasticity in the hygrophyte *Moringa oleifera* exposed to water stress

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1 **Metabolic plasticity in the hygrophyte *Moringa oleifera* exposed to water stress**

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Abstract

Over the past decades, introduction of many fast-growing hygrophilic, and economically valuable plants into xeric environments has occurred. However, production and even survival of these species may be threatened by harsh climatic conditions unless an effective physiological and metabolic plasticity is available. *Moringa oleifera* Lam., a multi-purpose tree originating from humid sub-tropical regions of India, is widely cultivated in many arid countries because of its multiple uses. We tested whether *M. oleifera* can adjust primary and secondary metabolism to efficiently cope with increasing water stress. It is shown that *M. oleifera* possesses an effective isohydric behavior. Water stress induced a quick and strong stomatal closure, driven by abscisic acid (ABA) accumulation, and leading to photosynthesis inhibition with consequent negative effects on biomass production. However, photochemistry was not impaired and maximal fluorescence and saturating photosynthesis remained unaffected in stressed leaves. We report for the first time that *M. oleifera* produces isoprene, and show that isoprene emission increased three-fold during stress progression. It is proposed that higher isoprene biosynthesis helps leaves cope with water stress through its antioxidant or membrane stabilizing action, and also indicates a general MEP (methylerythritol 4-phosphate) pathway activation that further helps protect photosynthesis under water stress. Increased concentrations of antioxidant flavonoids were also observed in water stressed leaves, and probably cooperate in limiting irreversible effects of the stress in *M. oleifera* leaves. The observed metabolic and phenotypic plasticity may facilitate the establishment of *M. oleifera* in xeric environments, sustaining the economic and environmental value of this plant.

Key words:

Abscisic acid, flavonoids, isoprene, isohydry, MEP (methylerythritol 4-phosphate) pathway, violaxanthin-cycle pigments, water stress.

1. Introduction

There is increasing interest in understanding how plants cope with the severe challenges imposed by climate change. Recurrent droughts and heat waves will likely be amplified in the near future, particularly in mid-latitude and subtropical dry regions (Dai 2013). 'Drought tolerant' plants that are adapted to arid environments (Kozlowsky and Pallardy 2002) invest a large portion of assimilated carbon to increase leaf density and thickness and on the biosynthesis of carbon-based secondary compounds, rather than promoting new growth (Niinemets 2001; Rivas-Ubach et al. 2012). Adverse climate conditions may threaten the survival of fast-growing hygrophilic species which are largely cultivated in xeric environments for ecological restoration and profitable biomass production. This is the case of *Moringa oleifera* Lam., a fast-growing tree native to sub-Himalayan northwest India (Pandey et al. 2011), where mean annual precipitations exceed 1,100 mm (Singh and Mal 2014), mainly concentrated during the monsoon season. *Moringa oleifera* is a multipurpose tree crop utilized for human food and livestock forage because of its high vitamin content (Anwar et al. 2007; Verma et al. 2009; Fuglie 2011; Nouman et al. 2014). This species is also used for many medicinal purposes and is considered a life-saving resource (Fahey 2005; Kasolo et al. 2010; Mbikay 2012; El Sohaimy et al. 2015), while its oleic acid-rich seeds can be used to produce biodiesel (Rashid et al. 2008; Da Silva et al. 2010). Because of these multiple applications, *M. oleifera* has been called a "miracle tree" and its cultivation range has rapidly expanded into sub-tropical dry regions across Africa, South America and Asia, characterized by recurrent droughts combined with both high air temperatures and solar irradiance (Leone et al. 2015). However, if climatic constraints become harsher and more frequent under the influence of climate change, they may threaten the survival and profitable production of *M. oleifera*, which apparently does not possess any conservative functional trait of adaptation to drought (Valladares et al. 2007).

Other adaptive traits related to secondary metabolism could play a determinant role in the process of plant acclimation to harsh environments, which are not explored in *M. oleifera*. There is overwhelming evidence that secondary metabolites derived from both the methylerythritol 4-

1
2 81 phosphate (MEP) and the phenylpropanoid pathway play a key role in the acclimation of 'mesic'
3
4 82 species to low water availability (Tattini et al. 2015; Velikova et al. 2016; Zalindas et al. 2017). For
5
6 83 instance, the emission of isoprene is more widespread in hygrophytes than in xerophytes (Loreto et al.
7
8 84 2014), and isoprene is believed to ameliorate the response of fast-growing species to drought stress
9
10 85 episodes (Loreto and Fineschi 2015; for reviews, see also: Sharkey et al. 2007; Fini et al. 2017).
11
12 86 Isoprene preserves the integrity of thylakoid membranes (Velikova et al. 2011) and scavenges reactive
13
14 87 oxygen species (ROS), particularly singlet oxygen (1O_2) (Velikova et al. 2004; Vickers et al. 2009a;
15
16 88 Zeinali et al. 2016), which are generated at considerable rates in drought-stressed leaves. The benefits
17
18 89 of isoprene biosynthesis on chloroplast membrane-associated processes may improve the use of
19
20 90 radiant energy for carbon fixation under stressful conditions (Pollastri et al. 2014; Vanzo et al. 2015),
21
22 91 thus reducing the risk of photo-oxidative damage (Vickers et al. 2009b).

23
24 92 In drought-stressed leaves, the enhancement of carbon flow through the MEP pathway also
25
26 93 promotes the synthesis of isoprene and non-volatile isoprenoids such as carotenoids and abscisic acid
27
28 94 (ABA) (Tattini et al. 2014; Marino et al. 2017). Carotenoids are known to protect photosynthesis under
29
30 95 drought stress (Beckett et al. 2012; Tattini et al. 2015). The photoprotective functions of carotenoids
31
32 96 include: quenching of triple state chlorophyll ($^3Chl^*$); thermal dissipation of excess energy through de-
33
34 97 epoxidation of xanthophylls (nonphotochemical quenching, NPQ) (Brunetti et al. 2015); and an
35
36 98 antioxidant function of zeaxanthin (Zea) in the chloroplasts by strengthening thylakoid membranes
37
38 99 under heat stress (Havaux et al. 2007; Dall'Osto et al. 2010; Esteban et al. 2015). Notably, biosynthesis
39
40 100 of Zea throughout β -hydroxylation of β -carotene (β -car) may also enhance drought resistance
41
42 101 (Davison et al. 2002; Du et al. 2010), possibly because Zea interacts with light harvesting complex b
43
44 102 (LHCb), thus reducing the production of 1O_2 and sustaining NPQ in high light conditions (Johnson et al.
45
46 103 2007; Dall'Osto et al. 2010). In turn, β -car (like isoprene, see Velikova et al. 2004) is an effective
47
48 104 chemical quencher of 1O_2 (Ramel et al. 2012).

49
50 105 A relationship between isoprene and foliar ABA has been repeatedly observed (Barta and
51
52 106 Loreto 2006; Tattini et al. 2014; Marino et al. 2017). ABA plays a major role in the regulation of
53
54 107 stomatal movements in plants capable of maintaining leaf water potential and relative water content
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1
2 108 unchanged under drought stress conditions (isohydric behavior) (Brodribb and McAdam 2013;
3
4 109 McAdam and Brodiribb, 2014; Coupel-Ledru et al. 2017).

5
6 110 Phenylpropanoid metabolism is another complex “metabolic grid” highly modulated by
7
8 111 environmental constraints (Laurson et al. 2015). Strong evidence has been provided that enhanced
9
10 112 biosynthesis of dihydroxy B-ring-substituted flavonoids is induced under drought, when the use of
11
12 113 light for photosynthesis is reduced (Tattini et al. 2004; Treutter 2006; Agati et al. 2012; Ma et al.
13
14 114 2014). Alterations in ROS homeostasis and/or in the electron transport chain are main drivers for
15
16 115 flavonoid biosynthesis (Taylor and Grotewold 2005; Fini et al. 2012; Fini et al. 2014). Flavonoids
17
18 116 accumulate to a large extent in the mesophyll cells of sun-exposed leaves and may complement the
19
20 117 functions of primary antioxidants in plants, both acting at different places and at different times
21
22 118 (Brunetti et al. 2015; Tattini et al. 2015). In fact, flavonoids are found in sub-cellular compartments,
23
24 119 such as the nucleus, vacuole and outer chloroplast membrane, where other antioxidants do not
25
26 120 effectively operate (Agati et al. 2007; Ferreres et al. 2011).

27
28 121 In plants exposed to drought, the modulation of secondary metabolism may be mostly
29
30 122 intended to reduce excess of ROS by increasing the production of metabolites with antioxidant
31
32 123 properties, including isoprenoids and phenylpropanoids (Nakabayashi et al. 2014; Loreto and Fineschi
33
34 124 2015; Tattini et al. 2015). We hypothesize that both enhanced production and profound re-adjustment
35
36 125 in the isoprenoid and phenylpropanoid pool (i.e. metabolic plasticity) may occur in hygrophilic and
37
38 126 fast-growing plants such as *M. oleifera* when facing drought conditions. To test this hypothesis and
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40 127 explore physiological and biochemical mechanisms linked to drought resistance in hygrophilic plants,
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42 128 we exposed *M. oleifera* plants to a water stress treatment of increasing severity.

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45 46 130 **2. Materials & Methods**

47 48 49 131 2.1. Plant material and experimental conditions

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52 132 Two-month-old seedlings of *Moringa oleifera* Lam. were planted in 50 L pots with a sand/peat
53
54 133 substrate (9/1, v/v), and were grown outside in Florence, Italy (43° 49' N, 11° 37'). The experiment
55
56 134 was conducted during summer 2014, under minimum/maximum temperatures of $17.7 \pm 2.4/30.8 \pm$

1 135 3.2°C (mean \pm standard deviation, S.D.) and midday irradiance (measured over the 200-3000 nm
2
3 136 range of solar wavebands) of $780 \pm 85 \text{ W m}^{-2}$ (mean \pm S.D.). Saplings were irrigated to pot capacity
4
5 137 before the onset of water stress treatment, that was applied to plants on average ~ 190 cm tall and
6
7 138 with stem diameter of ~ 2.0 cm. Water stress was imposed by withholding water for 30 days (WS,
8
9 139 water-stressed plants), whereas control plants (C) were irrigated daily to pot capacity. A total of thirty
10
11 140 plants were grown under these two experimental conditions (12 assigned to well-watered treatment
12
13 141 and 18 assigned to water-stressed treatment). Plants were assigned on the basis of preliminary leaf
14
15 142 gas exchange measurements to exclude significant differences in photosynthesis (A_N) and stomatal
16
17 143 conductance (g_s) among plants ($t < 0.05$, data not shown). The fraction of transpirable soil water
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19 144 (FTSW) and g_s were used as water stress indicators (Sinclair and Ludlow, 1986; Brillì et al., 2013).
20
21 145 Measurements were conducted in water-stressed plants at increasing stress severity. The three stress
22
23 146 levels corresponded to decreasing FTSW from 100% (in control plants, FTSW₁₀₀), to 60% (FTSW₆₀),
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25 147 40% (FTSW₄₀) and 25% (FTSW₂₅), corresponding to 10, 20 and 30 days after withholding water,
26
27 148 respectively. Control plants were also sampled at the same days as water-stressed plants, to make sure
28
29 149 that control conditions were maintained across the experimental period. The physiological lower limit
30
31 150 of available soil water, corresponding to the FTSW endpoint, was calculated prolonging water stress,
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33 151 until stomatal conductance approached zero, on some additional plants.

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36 152 As *M. oleifera* has bipinnate compound leaves, water relations, gas exchange, chlorophyll
37
38 153 fluorescence, isoprene and *n*-hexanal measurements were conducted on the two medial leaflets in the
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40 154 secondary pinna (hereafter denoted as leaf), on four replicate plants per treatment, at each sampling
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42 155 date. The adjacent leaf was collected for biochemical measurements, between 12:00-14:00 h.

43 44 45 156 2.2. Growth, water relations, gas exchange and chlorophyll fluorescence

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48 157 Biomass was measured at the end of the water stress period (30 d) on ten replicate plants per
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50 158 treatment. Plants were divided into shoots and roots and oven dried at 70 °C until a constant weight
51
52 159 was reached (after about 72 h). Biomass allocation was calculated on a dry mass (DM) basis, using as
53
54 160 parameters the ratio of shoot dry mass to total dry mass (BAS) and the ratio of root dry mass to total
55
56 161 dry mass (BAR). Predawn measurements of relative water content (RWC), water (ψ_w) and osmotic

162 (ψ_{π}) potentials were made on well-watered and water-stressed leaves (2 leaves for each selected
163 replicate).

164 Gas exchange was measured on intact leaves using a LI-6400 portable photosynthesis system
165 (Li-Cor, Lincoln, NE, USA). Measurements were performed at a photosynthetic photon flux density
166 (PPFD) of 1000 $\mu\text{mol photons m}^{-2} \text{ s}^{-1}$, a CO_2 concentration of 400 $\mu\text{mol mol}^{-1}$ and ambient
167 temperature. This system was utilized also to measure leaf temperature. Photosynthesis and g_s were
168 calculated using the LI-6400 software. Chlorophyll (Chl) fluorescence was measured using a
169 modulated PAM-2000 fluorometer (Heinz Walz, Effeltrich, Germany). Minimum fluorescence (F_0) was
170 measured with a 0.8 $\mu\text{mol m}^{-2} \text{ s}^{-1}$ measuring light beam on leaves that were dark-adapted for 20
171 minutes. Maximum fluorescence in the dark-adapted state (F_m) was determined using a saturating
172 pulse (0.5 s) of red light (8000 $\mu\text{mol m}^{-2} \text{ s}^{-1}$), thus allowing calculation of $F_v/F_m = (F_m - F_0)/F_m$. Actinic
173 red continuous light (1000 $\mu\text{mol m}^{-2} \text{ s}^{-1}$) was then switched on, and steady-state fluorescence was
174 recorded (F_s). Saturating pulses were then applied to record the maximum fluorescence under actinic
175 light (F'_m). These data were used to calculate non-photochemical quenching ($\text{NPQ} = (F_m - F'_m)/F'_m$)
176 (Schreiber et al., 1986), actual quantum yield of PSII ($\Phi_{\text{PSII}} = (F'_m - F_s)/F'_m$) (Genty et al. 1989), and
177 electron transport rate ($\text{ETR} = 0.5 \cdot \Phi_{\text{PSII}} \cdot \text{PAR} \cdot 0.84$), where 0.5 and 0.84 are coefficients indicating
178 an equal distribution of photons between PSI and PSII and leaf absorptance, respectively.

179 2.3. Isoprene, abscisic acid and photosynthetic pigments

180 To measure isoprene emission, the outlet of the cuvette was disconnected from the LI-6400
181 system and the flow was diverted into a silcosteel cartridge packed with 200 mg of Tenax (Agilent,
182 Cernusco sul Naviglio, Italy). A volume of 4.5 dm^3 of air was pumped through the trap at a rate of 200
183 $\text{cm}^3 \text{ min}^{-1}$. The cartridge was analysed using a Perkin Elmer Clarus 580 gas chromatograph coupled
184 with a Clarus 560 Mass-Selective-Detector and a thermal desorber TurboMatrix (Perkin Elmer Inc.,
185 Waltham, MA, USA). The desorbed compounds were separated in a 30-m Elite-5-MS capillary column.
186 The column oven temperature was kept at 40 $^{\circ}\text{C}$ for the first 5 min, then increased by 5 $^{\circ}\text{C min}^{-1}$ to 250
187 $^{\circ}\text{C}$, and maintained at 250 $^{\circ}\text{C}$ for 2 min. Helium was used as carrier gas. Compounds were identified
188 using the NIST library provided with the GC/MS Turbomass software. Quantification of isoprene was

1
2 189 conducted using authentic standards of isoprene (Rivoira, Milan, Italy) to prepare a calibration curve
3
4 190 as well as to compare the peak retention time and the peak fragmentation of isoprene found in the
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6 191 samples.

7
8 192 Abscisic acid, both in its free (free-ABA) and conjugated form (ABA glucoside ester, ABA-GE),
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10 193 was extracted and quantified as reported in Tattini et al. (2017). In detail, 200 mg of lyophilized leaf
11
12 194 tissue were ground in liquid nitrogen and combined with 50 ng of d⁶-ABA and d⁵-ABA-GE (National
13
14 195 Research Council of Canada), then extracted with 3 × 1 cm³ pH 2.5 CH₃OH/H₂O (50:50; v:v), at 4 °C for
15
16 196 30 minutes. The supernatant, after defatting with 3 × 3 cm³ of *n*-hexane, was purified using Sep-Pak
17
18 197 C18 cartridges (Waters, Milford, MA, USA) and eluted with 1 cm³ of ethylacetate. The eluate, dried
19
20 198 under nitrogen, and rinsed with 500 µL pH 2.5 CH₃OH/H₂O (50:50), was injected (3 µL aliquots) in a
21
22 199 LC-DAD-MS/MS system, consisting of a Shimadzu Nexera HPLC and a Shimadzu LCMS-8030
23
24 200 quadrupole mass spectrometer, operating in electrospray ionization (ESI) mode (Kyoto, Japan). The
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26 201 eluting solvents consisted of H₂O (added with 0.1 % of HCOOH, solvent A) and CH₃CN/CH₃OH (1:1, v:v,
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28 202 added with 0.1 % of HCOOH, solvent B). The analysis was performed in negative ion mode, using a 3 ×
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30 203 100 mm Poroshell 120 SB C18 column (2.7 µm, 100 × 4.6 mm, Agilent Technologies) and eluting a 18
31
32 204 min-run from 95% solvent A to 100% solvent B at a flow rate of 0.3 cm³ min⁻¹. Quantification was
33
34 205 conducted in multiple reaction mode (MRM), as reported by López-Carbonell et al. (2009).

35
36 206 Chlorophyll *a* and *b*, and individual carotenoids were identified and quantified as reported by
37
38 207 Beckett et al. (2012). Briefly, lyophilized leaf tissue (0.2 g) was extracted with 3 × 5 cm³ acetone
39
40 208 (added with 0.5 g cm⁻³ of CaCO₃) and injected (15 µL) in a Flexar high performance liquid
41
42 209 chromatography (HPLC) system equipped with a quaternary 200Q/410 pump and a LC 200 diode
43
44 210 array detector (DAD) (all from Perkin Elmer Bradford, CT, USA). Photosynthetic pigments were
45
46 211 separated in a 250 × 4.6 mm Agilent Zorbax SB-C18 (5 µm) column operating at 30°C, eluted for 18
47
48 212 min with a linear gradient solvent system, at a flow rate of 1 cm³ min⁻¹, from 100% CH₃CN/MeOH
49
50 213 (95/5 with 0.05% of triethylamine) to 100% MeOH/ethyl acetate (6.8/3.2). Violaxanthin cycle
51
52 214 pigments [violaxanthin (Vio), antheraxanthin (Ant), zeaxanthin (Zea), collectively named (VAZ)],
53
54 215 neoxanthin (Neo), lutein (Lut), β-carotene (β-car), chlorophyll *a* and chlorophyll *b* were identified
55
56 216 using visible spectral characteristics and retention times. Carotenoids and chlorophylls were

1
2 217 calibrated using authentic standards from Extrasynthese (Lyon-Nord, Genay, France) and from Sigma
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4 218 Aldrich (Milan, Italy), respectively. The de-epoxidation state of VAZ (DES) was calculated as:

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6 219
$$\text{DES} = (0.5A + Z)/(V + A + Z)$$

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9 220 2.4. Flavonoids

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11 221 Individual flavonoids were extracted and quantified as previously reported in Tattini et al.
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13 222 (2015). Briefly, lyophilized leaf tissue (0.2 g) was extracted with $3 \times 5 \text{ cm}^3$ of 75% EtOH/H₂O adjusted
14
15 223 to pH 2.5 with formic acid, and the supernatant partitioned with $4 \times 5 \text{ cm}^3$ *n*-hexane, reduced to
16
17 224 dryness and finally rinsed with 2 cm^3 CH₃OH/H₂O (8:2, v:v). Aliquots of 10 μL were injected into the
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19 225 Perkin Elmer liquid chromatography system reported above, and compounds separated in a 150×4.6
20
21 226 mm Sun Fire column (5 μm) (Waters Italia, Milan, Italy) operating at 30 °C and eluted at a flow rate of
22
23 227 $1 \text{ cm}^3 \text{ min}^{-1}$. The mobile phases were (A) H₂O pH 4.3 (CH₃COONH₄/CH₃COOH)/CH₃CN (90/10, v/v) and
24
25 228 (B) CH₃CN/H₂O pH 4.3 (CH₃COONH₄ /CH₃COOH) (90/10, v/v). Flavonoids were separated using a
26
27 229 linear gradient elution from A to B over a 46 min-run. Flavonoids were identified by comparison of
28
29 230 their retention times and UV spectral characteristics with those of authentic standards (Extrasynthese,
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31 231 Lyon-Nord, Genay, France) and quantified at 350 nm using five-point calibration curves of authentic
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33 232 standards.
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37 233 2.5. Lipid peroxidation indicator (*n*-hexanal)

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40 234 *N*-hexanal is one of the lipid peroxide-derived carbonyl compounds (oxylipin carbonyls) that
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42 235 reveals abiotic stress-induced damage of plants, and in particular of cellular membranes (Mano 2012).
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44 236 Analysis of *n*-hexanal was done using the same procedure as for isoprene (see above). Quantification
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46 237 of *n*-hexanal was conducted using an authentic standard (Sigma Aldrich, Milan, Italy) to prepare a
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48 238 calibration curve, as well as comparing the peak retention time and the peak fragmentation in all
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50 239 samples.
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53 240 2.6. Experimental design and statistics
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1
2 241 The experiment was performed using a completely randomized design. Biomass was measured
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4 242 on ten replicates for both well-watered and water-stressed plants at the end of the experiment.
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6 243 Physiological and biochemical measurements were conducted on four replicate plants, both in well-
7
8 244 watered plants and in plants exposed to water stress of increasing severity. Data were analysed using
9
10 245 repeated-measures ANOVA, with water treatment as between-subjects effect and sampling date as
11
12 246 within-subjects effect (SPSS v.20; IBM, Chicago IL, USA). Significant differences among means were
13
14 247 estimated at the 5% ($P < 0.05$) level, using Tukey's test.

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17 249 **3. Results**

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21 251 3.1. Water stress effects on water relations, photosynthesis and biomass production

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24
25 252 Predawn leaf water potential (ψ_w , Fig. 1A) declined in water-stressed plants compared to
26
27 253 control plants, though differences became significant only at FTSW₄₀ and FTSW₂₅. It is noteworthy
28
29 254 that at the end of the water stress cycle, when g_s of FTSW₂₅ plants was on average about 15% of the
30
31 255 control values, predawn ψ_w was still rather high (i.e., -0.60 MPa). Significant differences in leaf bulk
32
33 256 osmotic potential (ψ_π , Fig. 1B) were recorded only at the end of the experiment (FTSW₂₅), whereas
34
35 257 RWC did not significantly vary between water-stressed and control plants (Fig. 1C).

36
37 258 As FTSW declined, A_N , g_s , and C_i (Fig. 2; Tab. SM1) were progressively reduced. A strong
38
39 259 reduction of A_N (-30%, Fig. 2A) and g_s (-43%, Fig. 2B) was observed already under mild water stress
40
41 260 (FTSW₆₀). Under a more severe water stress (FTSW₂₅), A_N and g_s declined by 71% and 85%
42
43 261 respectively, compared to control leaves (FTSW₁₀₀) (Tab. SM1). Similarly, C_i (Fig. 2C) was reduced by
44
45 262 about 55% in FTSW₂₅ plants relative to FTSW₁₀₀ plants. The maximum quantum yield of PSII (F_v/F_m ,
46
47 263 Fig. 2D) did not vary between control and water-stressed plants, irrespective of the severity of the
48
49 264 stress. In contrast, the actual efficiency of PSII photochemistry (Φ_{PSII}), significantly declined already at
50
51 265 FTSW₆₀ and was further impaired at FTSW₄₀ and FTSW₂₅, relative to FTSW₁₀₀ plants (Fig. 2E). Water
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53 266 stress reductions in Φ_{PSII} were paralleled by corresponding increases in the non-photochemical
54
55 267 quenching of fluorescence (NPQ, Fig. 2F and SM1).

1
2 268 Plant biomass was significantly reduced in FTSW₂₅ compared to FTSW₁₀₀ plants (Fig. 3A). At
3
4 269 the end of the experiment, the root to shoot ratio was also significantly higher in FTSW₂₅ than in
5
6 270 FTSW₁₀₀ plants, whereas the shoot to total dry mass ratio was significantly reduced in water-stressed
7
8 271 plants (Fig. 3B).

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10 272 3.2. Water stress effects on isoprene, non-volatile isoprenoids, pigments, flavonoids, and
11
12 273 membrane lipid peroxidation.

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14
15 274 Isoprene emission increased significantly in FTSW₁₀₀ leaves during the experiment (Fig. 4A),
16
17 275 likely because of the prolonged exposure to elevated temperatures during the summer season.
18
19 276 Isoprene emission strongly and significantly increased in response to water stress. This increment was
20
21 277 particularly relevant at FTSW₂₅ (+86% compared to FTSW₄₀). The carbon lost as isoprene (C_{iso} %), also
22
23 278 increased largely in FTSW₂₅ plants, due to the simultaneous increase of isoprene emission and
24
25 279 reduction of A_n (Fig. 4B). The surging emission of isoprene was positively correlated to both the
26
27 280 decline of C_i (Fig. 5A) and the increase of the ETR/ A_n ratio (Fig. 5B) in water-stressed leaves.

28
29 281 The content of free-ABA and ABA-GE increased in water-stressed compared to control leaves
30
31 282 (Fig. 6A and 6B), and the effect was particularly strong in FTSW₂₅ plants where free-ABA and ABA-GE
32
33 283 contents were about seven and two folds higher than in FTSW₁₀₀, respectively. A strong negative linear
34
35 284 relationship ($R^2 = 0.915$) was found between foliar free-ABA levels and g_s (inset of Fig. 6A). Whereas,
36
37 285 free and conjugated ABA contents were both positively related to isoprene emission rates (inset of Fig.
38
39 286 6B).

40
41 287 Total chlorophyll (Chl_{tot}) declined significantly in FTSW₄₀ (-14%) and FTSW₂₅ (-23%) leaves in
42
43 288 comparison to FTSW₁₀₀ leaves (Fig. 7A). In contrast, total carotenoid (Car_{tot}) content did not vary
44
45 289 between control and water-stressed leaves, and increased during the experiment irrespective of water
46
47 290 treatments (Fig. 7B). However, water stress markedly altered the composition of the carotenoid pool.
48
49 291 The content of Lut (Fig. 7C) increased, whereas the content of β -car (Fig. 7D) declined significantly
50
51 292 under severe water deficit (FTSW₂₅). Among xanthophylls, Vio (Fig. 7E) declined and Zea (Fig. 7F)
52
53 293 increased significantly in water-stressed plants. Vio reduction was particularly strong at FTSW₆₀ and
54
55 294 FTSW₄₀, and partially recovered under severe stress conditions (FTSW₂₅). The contents of Ant (Fig. 7G)

1
2 295 and Neo (Fig. 7H) were not affected by water stress. However, Neo increased during the experimental
3
4 296 period in both well-watered and water-stressed plants. The content of violaxanthin-cycle pigments
5
6 297 (VAZ) relative to Chl_{tot} increased significantly as water stress progressed, and the effect was
7
8 298 particularly high (+35%) at FTSW₂₅ compared to FTSW₁₀₀ after 30 days of water stress (Fig. 7I). In
9
10 299 addition, DES increased in water-stressed compared to control leaves, but the difference was already
11
12 300 noticeable under mild water stress conditions (FTSW₆₀) (Fig.7J).

13
14 301 Water stress also considerably altered the content and composition of the flavonoid pool (Fig.
15
16 302 8 A-C). Quercetin-3-*O*-glycoside and its derivatives were the most responsive compounds to water
17
18 303 stress, as their content significantly and consistently increased with the intensity of the stress (Fig.
19
20 304 8A). In addition, the content of Kaempferol-3-*O*-glycoside derivatives also significantly increased in
21
22 305 response to stress, but the difference between water-stressed and control leaves remained constant as
23
24 306 water stress progressed (Fig. 9B). In contrast, the content of Apigenin-7-*O*-glycoside and its
25
26 307 derivatives significantly decreased in FTSW₄₀ and FTSW₂₅ leaves (Fig. 8C).

27
28 308 Compared to control leaves, the emission of *n*-hexanal did not significantly vary at both
29
30 309 FTSW₆₀ and FTSW₄₀, whereas it significantly increased in FTSW₂₅ plants (Fig. 9).

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32 310

33 34 311 **4. Discussion**

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36
37 312 4.1 Understanding the impact of water stress on the physiology of the isohydric plant *M.*
38
39 313 *oleifera*

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41
42 314 *M. oleifera* is a fast-growing species able to produce large quantities of biomass (Sánchez et al. 2006).
43
44 315 However, whether *M. oleifera* is able to acclimate and produce at satisfactory rates in arid conditions is
45
46 316 yet not known. Our study offers novel insights on the physiological and biochemical strategies adopted
47
48 317 by this species to cope with extended periods of soil water stress.

49
50 318 Our results show that *M. oleifera* possesses an effective avoidance mechanism (i.e. isohydry,
51
52 319 Nardini et al., 2014; Tardieu and Simmoneau, 1998) when subjected to water stress. This involved a
53
54 320 rapid reduction of g_s in stressed leaves, that possibly contributed to the maintenance of ψ_w and RWC
55
56 321 even in conditions of severe water stress (FTSW₂₅), when only a moderate (8%) reduction of the

1
2 322 osmotic component ψ_{π} became significant (Fig. 1 and 2B). The response of g_s of *M. oleifera* to soil
3
4 323 drying (Tab. SM1) is remarkably different from that observed in other fast-growing trees species such
5
6 324 as *Eucalyptus citriodora* (Brilli et al. 2013; Mahmood et al. 2015) and *Populus spp* (Marron et al. 2002;
7
8 325 Yin et al. 2005; Brilli et al. 2007; Centritto et al. 2011) that showed no or very little decline in g_s under
9
10 326 moderate water stress conditions. Isohydry is a crucial adaptive trait for the survival of deciduous
11
12 327 woody plants exposed to high evaporative demand and low soil water availability, as an early and tight
13
14 328 control of stomatal aperture may prevent xylem embolism (Franks et al. 2007; Yi et al. 2017). While
15
16 329 stomatal closure increased intrinsic water use efficiency (iWUE, determined as the ratio of A_n to g_s)
17
18 330 during water stress progression, it also constrained photosynthesis due to increased diffusional
19
20 331 limitations to CO_2 entry, with consequent reduction of C_i (Fig. 2C) (Lawlor and Cornic 2002; Centritto
21
22 332 et al. 2011; Lauteri et al. 2014; Fini et al. 2016). The observed drop in photosynthesis under water
23
24 333 stress caused a biomass reduction (Fig. 2 and 3), probably inducing a redistribution of the assimilated
25
26 334 carbon between shoots and roots (Peuke et al. 2006). These results suggest a high degree of plasticity
27
28 335 of *M. oleifera* in biomass allocation in response to water stress (Fig. 3).

30 336 Water stress did not cause permanent damages to the photosynthetic apparatus. In fact,
31
32 337 maximal PSII photochemical efficiency (F_v/F_m) did not decline even under severe water stress
33
34 338 (FTSW₂₅), suggesting stability of photochemical reactions and structures (Fig. 2) (Flexas et al., 2006).
35
36 339 However, PSII quantum yield in the light (Φ_{PSII}) was reduced as compared to FTSW₁₀₀ leaves. While
37
38 340 this mirrored A_n reduction at mild (FTSW₆₀) and moderate (FTSW₄₀) stress level, Φ_{PSII} did not drop
39
40 341 further in severely water-stressed leaves (Havaux 1992; Lu and Zhang, 1999) revealing a likely
41
42 342 increase of photorespiratory electron transport, or alternative electron sinks (see discussion below
43
44 343 about ETR driving isoprene emission). Furthermore, changes in NPQ and Φ_{PSII} were strongly
45
46 344 correlated throughout the experiment ($\Phi_{PSII} = -0,13 \text{ NPQ} + 0.62$, $R^2 = 0.844$, linear relation shown in
47
48 345 Fig. SM1). Large excess of light energy not used by photosynthesis, as revealed by the fluorescence
49
50 346 parameter NPQ (Fig. 2F), may directly photoreduce O_2 , thus causing large ROS generation in water-
51
52 347 stressed leaves, with consequent damage to PSII. To explain why this was not observed in this
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54 348 experiment, we hypothesize a potential contribution of isoprenoids and phenylpropanoids as
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56 349 antioxidant compounds, as discussed below.

1
2 350 4.2. Exploring the significance of enhanced isoprene emission during water stress and its
3
4 351 relationship with foliar ABA

5
6 352 Our study revealed that *M. oleifera* is an isoprene emitting species (Fig. 4). Isoprene emission is
7
8 353 typical of hygrophytes that are fast-growing in temperate areas of the world (Loreto et al. 2014; Loreto
9
10 354 and Fineschi 2015), where isoprene serves important defensive (antioxidant and thermo-protective)
11
12 355 properties (Loreto and Schnitzler 2010; Velikova et al. 2011; Pollastri et al. 2014). We also show that
13
14 356 water stress promoted I_e , particularly when the stress became severe. Isoprene biosynthesis is
15
16 357 generally resistant to water stress (Brilli et al. 2007; Centritto et al. 2011; Brilli et al. 2013), and the
17
18 358 emission of isoprene is enhanced when isoprene-emitters recover from water stress (Sharkey and
19
20 359 Loreto, 1993; Fortunati et al., 2008). Stimulation of isoprene biosynthesis “during” water stress
21
22 360 episodes is less reported (Haworth et al. 2017; Marino et al. 2017). *M. oleifera* is a typical isoprene
23
24 361 emitting species, since it is a fast-growing species with high photosynthetic rates which thrives wild in
25
26 362 secondary tropical deciduous forests of the sub-Himalayan area (Loreto and Fineschi, 2015). Our data
27
28 363 suggest that declines in internal CO_2 concentration (C_i) and the increasing electron flux generated by
29
30 364 Photosystem II not used for carbon assimilation (ETR/A_N) are two important physiological drivers of
31
32 365 isoprene biosynthesis under water stress conditions (Fig. 5) (Guidolotti et al. 2011; Harrison et al.
33
34 366 2013; Morfopoulos et al. 2014; Marino et al. 2017). Reduced photosynthesis due to CO_2 starvation may
35
36 367 indeed increase the fraction of ETR available for alternative biosyntheses, including isoprenoids. In
37
38 368 addition, the increase in leaf temperature induced by stomatal closure under water stress (from $31.2 \pm$
39
40 369 0.7 °C in FTSW₁₀₀ leaves to 34.4 ± 0.6 °C in FTSW₂₅ leaves, mean \pm S.D.) might have contributed to
41
42 370 further enhance the rate of isoprene emission (Singsaas and Sharkey, 1998; Fares et al. 2011; Brilli et
43
44 371 al. 2013; Arab et al. 2016). Indeed, the activity of isoprene synthase is known to be stimulated by high
45
46 372 temperatures (Monson et al. 1992; Li et al. 2011). Increasing isoprene synthase activity may also help
47
48 373 explain the increase in I_e and $C_{iso}\%$ observed in well-watered leaves, along rising summer
49
50 374 temperatures during the course of our study (Rasulov et al. 2015).

51
52 375 We hypothesize that the rising investment of newly assimilated carbon for isoprene
53
54 376 biosynthesis helped leaves tolerate water stress because: a) isoprene protects the photosynthetic
55
56 377 apparatus from heat and oxidative damage by preserving the integrity of thylakoid membranes (Siwko

1
2 378 et al. 2007; Velikova et al. 2011, 2014, 2015) or by scavenging singlet oxygen (1O_2), a highly reactive
3
4 379 ROS in chloroplasts (Velikova et al. 2004; Zeinali et al. 2016); b) isoprene makes faster and smoother
5
6 380 the electron transport flow (Pollastri et al. 2014), especially under water stress conditions (Marino et
7
8 381 al. 2017). We found that NPQ did not vary between FTSW₄₀ and FTSW₂₅ leaves. Lower NPQ values in
9
10 382 isoprene emitters compared to non-emitters were reported both in stressful (Behnke et al. 2007,
11
12 383 2010) and physiological conditions (Pollastri et al. 2014). We, therefore, hypothesize a relationship
13
14 384 between the reduction of NPQ and the increase I_e along with the severity of water stress. A
15
16 385 downregulation of chloroplastic ATP-synthase and the consequent reduction in the flexible heat
17
18 386 dissipation component (qE) of NPQ (Demmig-Adams and Adams 2006) was reported in isoprene
19
20 387 emitting species by Velikova et al. (2014).

21
22 388 The observed strong linear relationships between I_e and foliar contents of free-ABA and ABA-
23
24 389 GE (Fig. 6B), suggest that increased isoprene formation in water stressed plants indicates enhanced
25
26 390 carbon flow through the MEP pathway, leading to higher foliar biosynthesis of abscisic acid (Fig. 6A)
27
28 391 (Marino et al. 2017). A relationship between isoprene and foliar ABA was first reported by Barta and
29
30 392 Loreto (2006) in well-watered *Populus alba* and by Tattini et al. (2014) in drought stressed transgenic
31
32 393 tobacco plants. Our results also show a strong linear correlation between free-ABA and g_s (Fig. 6A),
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34 394 despite limited variations of water relations in *M. oleifera* leaves. It is unclear whether isoprene is
35
36 395 simply of proxy of carbon flux through the MEP pathway, or has a regulatory role. Sustained isoprene
37
38 396 emission in water-stressed plants may reduce the accumulation of dimethylallyl pyrophosphate
39
40 397 (DMAPP) in the chloroplast, and may prevent DMAPP-induced feedback inhibition of the entire MEP
41
42 398 pathway (Banerjee et al. 2013). Taken together our results suggest that: a) increased isoprene
43
44 399 formation indicates and perhaps regulates free-ABA synthesis in stressed leaves, and b) free-ABA has a
45
46 400 major role in the regulation of stomatal closure compared to hydraulic signals (Chaves et al. 2016;
47
48 401 McAdam et al. 2016a). These results are in line with recent studies showing that, in strict isohydric
49
50 402 plants such as *M. oleifera*, high levels of free-ABA could be responsible for stomatal closure and could
51
52 403 promote a higher root to shoot ratio/carbon allocation (Nolan et al. 2017; McAdam et al. 2016b).

53
54
55 404 4.3. Plasticity of secondary metabolism in *M. oleifera* during water stress progression

1
2 405 We observed several changes in carotenoids and phenylpropanoids in response to increasing
3
4 406 water stress, that can be interpreted as a photoprotective trait to limit water stress induced damage.
5
6 407 The content of total carotenoids on a leaf mass basis also increased over the course of the experiment
7
8 408 in both well-watered and water-stressed leaves. While this shows a general upregulation of the MEP
9
10 409 pathway (see previous section) over the season, we argue that the investment in carotenoids was
11
12 410 much stronger in water-stressed leaves mirroring the depression in carbon assimilation. The blend of
13
14 411 carotenoids also changed along stress progression, perhaps favouring compounds active in stress
15
16 412 protection (Fig. 7). The increase in lutein in severely water-stressed plants might have enhanced the
17
18 413 capacity of leaves to quench $^3\text{Chl}^*$, that was likely generated during stress exposure (Dall'Osto et al.
19
20 414 2006; Jahns and Holzwarth 2012). In addition, compared to photosynthesis, Chl_{tot} content was less
21
22 415 affected by severe water stress, indicating a successful mechanism of protection. We also note that a
23
24 416 large switch in the composition of xanthophylls occurred in water-stressed plants. The increase in Zea
25
26 417 content was accompanied by a parallel decrease in Vio content under mild and moderate water stress,
27
28 418 showing the classic mechanism of de-epoxidation that is a major element of photoprotection in plants
29
30 419 (Demming-Adams and Adams 2006). However, when plants experienced the most severe water stress
31
32 420 the content in Zea and in Vio both increased. We suggest that the large increase in Zea biosynthesis
33
34 421 might have been originated from hydroxylation of β -car (Davison et al. 2002; Du et al. 2010). This is
35
36 422 consistent with the reduction of β -car concentration observed in leaves at FTWS₂₅. β -car might have
37
38 423 been also used as a chemical quencher of $^1\text{O}_2$ (Ramel et al. 2012), thus explaining the relatively
39
40 424 stronger decline of β -car ($-0.18 \mu\text{mol g}^{-1} \text{DW}$) as compared to the increase in Zea ($+0.07 \mu\text{mol g}^{-1} \text{DW}$)
41
42 425 when the stress became severe. The content of VAZ relative to Chl_{tot} was on average $> 70 \text{ mmol mol}^{-1}$
43
44 426 in both well-watered and water-stressed plants throughout the whole experiment, as commonly
45
46 427 observed in leaves long acclimated to full solar irradiance (Fini et al. 2014; Esteban et al. 2015). This
47
48 428 implies that only a fraction of the VAZ pool was bound to antenna systems and, hence, involved in NPQ
49
50 429 (Fig. 7I and J). In addition, the VAZ to Chl_{tot} ratio increased linearly during the water stress cycle. This
51
52 430 increasing 'unbound' VAZ pool might have served specific antioxidant functions in water-stressed
53
54 431 leaves, increasing membrane thermo-stability hence limiting lipid peroxidation (Havaux et al. 2007;
55
56 432 Esteban et al. 2015). This is an action similar to that suggested for isoprene (Velikova et al. 2011), and

1
2 433 cooperation between volatile and non-volatile isoprenoids was surmised by Beckett et al. (2012).
3
4 434 Indeed, the rate of *n*-hexanal emission, a marker of lipid peroxidation (Mano et al. 2012; Beckett et al.
5
6 435 2012), was only affected when a severe water stress was imposed (FTWS₂₅, Fig. 9), and was not
7
8 436 accompanied by irreversible degradation of membrane-bound photosynthetic machineries, namely
9
10 437 PSII photochemistry (as shown earlier).

11
12 438 The biosynthesis of antioxidant flavonoids, here constituted mainly by quercetin derivatives,
13
14 439 was stimulated in water-stressed leaves of *M. oleifera* (Fig. 8), similarly to what has been observed in
15
16 440 other plants (Tattini et al. 2004; Velikova et al. 2016; Ahrar et al. 2017). These high levels of foliar
17
18 441 flavonoids, commonly found in leaves grown under full sunlight, are not compatible with their
19
20 442 exclusive distribution in epidermal cells (Jaakola et al. 2004; Tattini et al. 2005; Agati et al. 2009; Majer
21
22 443 et al. 2014). Therefore, we suggest that water stress induced the accumulation of quercetin derivatives
23
24 444 mainly in mesophyll cells (Tattini et al. 2015), likely conferring increasing protection against enhanced
25
26 445 ROS generation (Agati and Tattini 2010; Agati et al. 2012; Nakabayashi et al. 2014), while reducing the
27
28 446 risk of permanent photodamage to PSII, by additionally acting as UV-B filters in the chloroplast
29
30 447 (Mierziak et al. 2014; Zavafer et al. 2017). The finding that water stress induced profound changes in
31
32 448 the composition of the flavonoid pool, with major increases in the biosynthesis of 'effective
33
34 449 antioxidant' quercetin derivatives (on average +46%), further supports our hypothesis. In contrast,
35
36 450 the content of less effective antioxidant' flavonoids either increased little (kaempferol glycosides,
37
38 451 +15%) or largely declined (apigenin glycosides -35%) in response to water stress. This significant
39
40 452 changes in the composition of flavonoids may also have contributed to reduce lipid peroxidation, as
41
42 453 previously discussed.

43
44 454

45 455 **Conclusions**

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48
49 456 Despite being originated in hygrophilic habitats, *M. oleifera* possesses multiple biochemical
50
51 457 and physiological mechanisms that allow this species to successfully tolerate water stress episodes.
52
53 458 These mechanisms include a strict isohydric behavior in response to water deprivation that is typical
54
55 459 of hygrophytes. The fast stomatal closure driven by high contents of foliar-ABA, however, caused an
56
57 460 early and strong depression in carbon assimilation with negative consequences for biomass

1
2 461 production. More interestingly, this study revealed that *M. oleifera* is an isoprene emitting species.
3
4 462 Increasing isoprene emission during progressive water stress was a valuable indicator for the general
5
6 463 activation of the MEP-pathway. The simultaneous increment of volatile and non-volatile isoprenoids
7
8 464 and of flavonoids, is suggested to be the key mechanism that allows *M. oleifera* to limit lipid
9
10 465 peroxidation and prevent severe photoinhibitory processes under water stress. This may allow a
11
12 466 prompt recovery of photosynthesis and growth rates when water is newly available to the roots. While
13
14 467 the observed high plasticity of stomatal conductance and secondary metabolites production may take
15
16 468 its toll on primary productivity of *M. oleifera*, it possibly also facilitates the establishment of this plant
17
18 469 to xeric environments. The extent to which the trade-off between primary and secondary metabolism
19
20 470 affects the resistance and whole-plant performance of a fast-growing plant such as *M. oleifera*, remains
21
22 471 to be determined in presence of recurrent periods of water stress.
23

24 472

25 473 **Authors' contributions**

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27
28 474 CB, FL, FF and MT planned the experiment. CB conducted the study, collected samples,
29
30 475 analyzed the data, and prepared the draft. AG, LG and DR helped in performing physiological and
31
32 476 chemical analyses. CB and MT interpreted the results and drafted the manuscript. FL, AF and MC
33
34 477 reviewed the manuscript.
35

36 478

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1
2 790 **Figure and table legends**

3
4 791 Figure 1. Predawn leaf water (Ψ_w , A) and osmotic (Ψ_π , B) potentials, and relative water content
5 792 (RWC, C) in FTSW₁₀₀ (F₁₀₀) plants (open bars) and in FTSW₆₀ (F₆₀), FTSW₄₀ (F₄₀) and FTSW₂₅ (F₂₅)
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7 793 water-stressed plants (grey bars) of *Moringa oleifera*, corresponding to 10, 20 and 30 days after
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9 794 withholding water, respectively. Data (means \pm SD, n = 4) were subjected to repeated measures with
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11 795 ANOVA, and bars not accompanied by the same letter significantly differ at the 5% level, using Tukey's
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13 796 test.

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17 797 Figure 2. Photosynthesis (A_N , A), stomatal conductance (g_s , B), intercellular CO₂ concentration
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19 798 (C_i , C), maximum (F_v/F_m , D) and actual (Φ_{PSII} , E) efficiency of PSII photochemistry and non-
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21 799 photochemical quenching (NPQ, F) in FTSW₁₀₀ (F₁₀₀) plants (open bars) and in FTSW₆₀ (F₆₀), FTSW₄₀
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23 800 (F₄₀) and FTSW₂₅ (F₂₅) water-stressed plants (grey bars) of *Moringa oleifera*. Data (means \pm SD, n = 4)
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25 801 were subjected to repeated measures with ANOVA, and bars not accompanied by the same letter
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27 802 significantly differ at the 5% level, using Tukey's test.

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31 803 Figure 3. Total biomass (A) and biomass allocation (B) in FTSW₁₀₀ (F₁₀₀) plants (open bars) and
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33 804 in FTSW₂₅ (F₂₅) water-stressed plants (grey bars) of *Moringa oleifera*. The percentage of biomass
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35 805 allocation (BA) was calculated considering the ratio of shoot dry mass to total dry mass (BAS) and the
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37 806 ratio of root dry mass to total dry mass (BAR). Data (means \pm SD, n = 10) were subjected to repeated
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39 807 measures with ANOVA, and bars not accompanied by the same letter significantly differ at the 5%
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41 808 level, using Tukey's test.

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44 809 Figure 4. Rates of isoprene emission (A) and carbon lost as isoprene (C_{iso} , B) in FTSW₁₀₀ (F₁₀₀)
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46 810 plants (open bars) and in FTSW₆₀ (F₆₀), FTSW₄₀ (F₄₀) and FTSW₂₅ (F₂₅) water-stressed plants (grey
47
48 811 bars) of *Moringa oleifera*. Data (means \pm SD, n = 4) were subjected to repeated measures with ANOVA,
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50 812 and bars not accompanied by the same letter significantly differ at the 5% level, using Tukey's test.

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52
53 813 Figure 5. Linear relationships between isoprene emission rate and (A) internal CO₂
54
55 814 concentration (C_i) or (B) the ratio of electron transport rate to photosynthesis (ETR/AN) in *Moringa*
56
57 815 *oleifera* plants. Measurements were made at FTSW₆₀ (10 d, open symbols), FTSW₄₀ (20 d, grey

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2 816 symbols), and FTSW₂₅ (30 d, closed symbols) both in well-watered plants (FTSW₁₀₀) (triangles) and
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4 817 water-stressed (circles) plants. Coefficient of determination (R^2) of each relationship are reported; ***
5
6 818 indicate $P < 0.0001$.

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9 819 Figure 6. Contents of free-ABA (A) and ABA-GE (B) in FTSW₁₀₀ (F₁₀₀) plants (open bars) and in
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11 820 FTSW₆₀ (F₆₀), FTSW₄₀ (F₄₀) and FTSW₂₅ (F₂₅) water-stressed plants (grey bars) of *Moringa oleifera*.
12
13 821 Data (means \pm SD, n = 4) were subjected to repeated measures with ANOVA, and bars not
14
15 822 accompanied by the same letter significantly differ at the 5% level, using Tukey's test. Inset in Figure
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17 823 6A shows the inverse relationship between foliar free-ABA content and stomatal conductance (g_s).
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19 824 Inset in Figure 6B shows the linear relationships between isoprene emission rates ($\text{nmol m}^{-2} \text{s}^{-1}$) and
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21 825 free-ABA and its glucoside ester (ABA-GE) contents in FTSW₁₀₀ plants (circles) and in water-stressed
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23 826 (triangles) plants at FTSW₆₀ (white symbols), FTSW₄₀ (grey symbols), and FTSW₂₅ (dark symbols),
24
25 827 respectively. Coefficient of determination (R^2) of each relationship are reported; *** indicate $P < 0.0001$.

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27
28 828 Figure 7. Effects of water stress on the contents of photosynthetic pigments (A-I), on the ratio
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30 829 of violaxanthin cycle pigment content to total chlorophyll content (VAZ Chl_{tot}⁻¹, I) and on the de-
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32 830 epoxidation state of VAZ [DES = (0.5A + Z) (V + A + Z)⁻¹, J] in FTSW₁₀₀ (F₁₀₀) plants (open bars) and in
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34 831 FTSW₆₀ (F₆₀), FTSW₄₀ (F₄₀) and FTSW₂₅ (F₂₅) water-stressed plants (grey bars) of *Moringa oleifera*.
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36 832 Data (means \pm SD, n = 4) were subjected to repeated measures with ANOVA, and bars not
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38 833 accompanied by the same letter significantly differ at the 5% level, using Tukey's test.

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41 834 Figure 8. Contents of quercetin (A), kaempferol (B) and apigenin (C) derivatives in FTSW₁₀₀
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43 835 (F₁₀₀) plants (open bars) and in FTSW₆₀ (F₆₀), FTSW₄₀ (F₄₀) and FTSW₂₅ (F₂₅) water-stressed plants
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45 836 (grey bars) of *Moringa oleifera*. Data (means \pm SD, n = 4) were subjected to repeated measures with
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47 837 ANOVA, and bars not accompanied by the same letter significantly differ at the 5% level, using Tukey's
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49 838 test.

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51
52 839 Figure 9. Rates of *n*-hexanal emission in FTSW₁₀₀ (F₁₀₀) plants (open bars) and in FTSW₆₀ (F₆₀),
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54 840 FTSW₄₀ (F₄₀) and FTSW₂₅ (F₂₅) water-stressed plants (grey bars) of *Moringa oleifera*. Data (means \pm

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2 841 SD, n = 4) were subjected to repeated measures with ANOVA, and bars not accompanied by the same
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4 842 letter significantly differ at the 5% level, using Tukey's test.

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6 843 Figure SM 1. The increase in non-photochemical quenching (NPQ) correlated negatively with
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8 844 the actual efficiency of PSII photochemistry (Φ_{PSII}). Measurements were made at FTSW₆₀ (10 d, open
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10 845 symbols), FTSW₄₀ (20 d, grey symbols), and FTSW₂₅ (30 d, closed symbols) both in well-watered
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12 846 control (FTSW₁₀₀) plants (circles) and water-stressed (triangles) plants of *Moringa oleifera*. Coefficient
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14 847 of determination (R^2) of the relationship is reported; *** indicate $P < 0.0001$.

15
16
17 848 Table SM 1. Results for photosynthesis (A_N) and stomatal conductance (g_s) (means \pm SD, n = 4)
18
19 849 in water-stressed plants of *Moringa oleifera* at different fraction of transpirable of soil water (FTSW)
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21 850 and days after the onset of water stress treatment.

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For Peer Review

1 **Metabolic plasticity in the hygrophyte *Moringa oleifera* exposed to water stress**

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Abstract

Over the past decades, introduction of many fast-growing hygrophilic, and economically valuable plants into xeric environments has occurred. However, production and even survival of these species may be threatened by harsh climatic conditions unless an effective physiological and metabolic plasticity is available. *Moringa oleifera* Lam., a multi-purpose tree originating from humid sub-tropical regions of India, is widely cultivated in many arid countries because of its multiple uses. We tested whether *M. oleifera* can adjust primary and secondary metabolism to efficiently cope with increasing water stress. It is shown that *M. oleifera* possesses an effective isohydric behavior. Water stress induced a quick and strong stomatal closure, driven by abscisic acid (ABA) accumulation, and leading to photosynthesis inhibition with consequent negative effects on biomass production. However, photochemistry was not impaired and maximal fluorescence and saturating photosynthesis remained unaffected in stressed leaves. We report for the first time that *M. oleifera* produces isoprene, and show that isoprene emission increased three-fold during stress progression. It is proposed that higher isoprene biosynthesis helps leaves cope with water stress through its antioxidant or membrane stabilizing action, and also indicates a general MEP (methylerythritol 4-phosphate) pathway activation that further helps protect photosynthesis under water stress. Increased concentrations of antioxidant flavonoids were also observed in water stressed leaves, and probably cooperate in limiting irreversible effects of the stress in *M. oleifera* leaves. The observed metabolic and phenotypic plasticity may facilitate the establishment of *M. oleifera* in xeric environments, sustaining the economic and environmental value of this plant.

Key words:

Abscisic acid, flavonoids, isoprene, isohydry, MEP (methylerythritol 4-phosphate) pathway, violaxanthin-cycle pigments, water stress.

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1. Introduction

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There is increasing interest in understanding how plants cope with the severe challenges imposed by climate change. Recurrent droughts and heat waves will likely be amplified in the near future, particularly in mid-latitude and subtropical dry regions (Dai 2013). 'Drought tolerant' plants that are adapted to arid environments (Kozlowsky and Pallardy 2002) invest a large portion of assimilated carbon to increase leaf density and thickness and on the biosynthesis of carbon-based secondary compounds, rather than promoting new growth (Niinemets 2001; Rivas-Ubach et al. 2012). Adverse climate conditions may threaten the survival of fast-growing hygrophilic species which are largely cultivated in xeric environments for ecological restoration and profitable biomass production. This is the case of *Moringa oleifera* Lam., a fast-growing tree native to sub-Himalayan northwest India (Pandey et al. 2011), where mean annual precipitations exceed 1,100 mm (Singh and Mal 2014), mainly concentrated during the monsoon season. *Moringa oleifera* is a multipurpose tree crop utilized for human food and livestock forage because of its high vitamin content (Anwar et al. 2007; Verma et al. 2009; Fuglie 2011; Nouman et al. 2014). This species is also used for many medicinal purposes and is considered a life-saving resource (Fahey 2005; Kasolo et al. 2010; Mbikay 2012; El Sohaimy et al. 2015), while its oleic acid-rich seeds can be used to produce biodiesel (Rashid et al. 2008; Da Silva et al. 2010). Because of these multiple applications, *M. oleifera* has been called a "miracle tree" and its cultivation range has rapidly expanded into sub-tropical dry regions across Africa, South America and Asia, characterized by recurrent droughts combined with both high air temperatures and solar irradiance (Leone et al. 2015). However, if climatic constraints become harsher and more frequent under the influence of climate change, they may threaten the survival and profitable production of *M. oleifera*, which apparently does not possess any conservative functional trait of adaptation to drought (Valladares et al. 2007).

Other adaptive traits related to secondary metabolism could play a determinant role in the process of plant acclimation to harsh environments, which are not explored in *M. oleifera*. There is overwhelming evidence that secondary metabolites derived from both the methylerythritol 4-

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2 81 phosphate (MEP) and the phenylpropanoid pathway play a key role in the acclimation of 'mesic'
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4 82 species to low water availability (Tattini et al. 2015; Velikova et al. 2016; Zalindas et al. 2017). For
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6 83 instance, the emission of isoprene is more widespread in hygrophytes than in xerophytes (Loreto et al.
7
8 84 2014), and isoprene is believed to ameliorate the response of fast-growing species to drought stress
9
10 85 episodes (Loreto and Fineschi 2015; for reviews, see also: Sharkey et al. 2007; Fini et al. 2017).
11
12 86 Isoprene preserves the integrity of thylakoid membranes (Velikova et al. 2011) and scavenges reactive
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14 87 oxygen species (ROS), particularly singlet oxygen (1O_2) (Velikova et al. 2004; Vickers et al. 2009a;
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16 88 Zeinali et al. 2016), which are generated at considerable rates in drought-stressed leaves. The benefits
17
18 89 of isoprene biosynthesis on chloroplast membrane-associated processes may improve the use of
19
20 90 radiant energy for carbon fixation under stressful conditions (Pollastri et al. 2014; Vanzo et al. 2015),
21
22 91 thus reducing the risk of photo-oxidative damage (Vickers et al. 2009b).

23
24 92 In drought-stressed leaves, the enhancement of carbon flow through the MEP pathway also
25
26 93 promotes the synthesis of isoprene and non-volatile isoprenoids such as carotenoids and abscisic acid
27
28 94 (ABA) (Tattini et al. 2014; Marino et al. 2017). Carotenoids are known to protect photosynthesis under
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30 95 drought stress (Beckett et al. 2012; Tattini et al. 2015). The photoprotective functions of carotenoids
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32 96 include: quenching of triple state chlorophyll ($^3Chl^*$); thermal dissipation of excess energy through de-
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34 97 epoxidation of xanthophylls (nonphotochemical quenching, NPQ) (Brunetti et al. 2015); and an
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36 98 antioxidant function of zeaxanthin (Zea) in the chloroplasts by strengthening thylakoid membranes
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38 99 under heat stress (Havaux et al. 2007; Dall'Osto et al. 2010; Esteban et al. 2015). Notably, biosynthesis
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40 100 of Zea throughout β -hydroxylation of β -carotene (β -car) may also enhance drought resistance
41
42 101 (Davison et al. 2002; Du et al. 2010), possibly because Zea interacts with light harvesting complex b
43
44 102 (LHCb), thus reducing the production of 1O_2 and sustaining NPQ in high light conditions (Johnson et al.
45
46 103 2007; Dall'Osto et al. 2010). In turn, β -car (like isoprene, see Velikova et al. 2004) is an effective
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48 104 chemical quencher of 1O_2 (Ramel et al. 2012).

49
50 105 A relationship between isoprene and foliar ABA has been repeatedly observed (Barta and
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52 106 Loreto 2006; Tattini et al. 2014; Marino et al. 2017). ABA plays a major role in the regulation of
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54 107 stomatal movements in plants capable of maintaining leaf water potential and relative water content

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2 108 unchanged under drought stress conditions (isohydric behavior) (Brodribb and McAdam 2013;
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4 109 McAdam and Brodiribb, 2014; Coupel-Ledru et al. 2017).

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6 110 Phenylpropanoid metabolism is another complex “metabolic grid” highly modulated by
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8 111 environmental constraints (Laurson et al. 2015). Strong evidence has been provided that enhanced
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10 112 biosynthesis of dihydroxy B-ring-substituted flavonoids is induced under drought, when the use of
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12 113 light for photosynthesis is reduced (Tattini et al. 2004; Treutter 2006; Agati et al. 2012; Ma et al.
13
14 114 2014). Alterations in ROS homeostasis and/or in the electron transport chain are main drivers for
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16 115 flavonoid biosynthesis (Taylor and Grotewold 2005; Fini et al. 2012; Fini et al. 2014). Flavonoids
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18 116 accumulate to a large extent in the mesophyll cells of sun-exposed leaves and may complement the
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20 117 functions of primary antioxidants in plants, both acting at different places and at different **times**
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22 118 (Brunetti et al. 2015; Tattini et al. 2015). In fact, flavonoids are found in sub-cellular compartments,
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24 119 such as **the** nucleus, vacuole and outer chloroplast membrane, where other antioxidants do not
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26 120 effectively operate (Agati et al. 2007; Ferreres et al. 2011).

27
28 121 **In plants exposed to drought, the modulation of secondary metabolism may be mostly**
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30 122 **intended to reduce excess of ROS by increasing the production of metabolites with antioxidant**
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32 123 **properties, including isoprenoids and phenylpropanoids (Nakabayashi et al. 2014; Loreto and Fineschi**
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34 124 **2015; Tattini et al. 2015). We hypothesize that both enhanced production and profound re-adjustment**
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36 125 **in the isoprenoid and phenylpropanoid pool (i.e. metabolic plasticity) may occur in hygrophilic and**
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38 126 **fast-growing plants such as *M. oleifera* when facing drought conditions. To test this hypothesis and**
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40 127 **explore physiological and biochemical mechanisms linked to drought resistance in hygrophilic plants,**
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42 128 **we exposed *M. oleifera* plants to a water stress treatment of increasing severity.**

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45 46 130 **2. Materials & Methods**

47 48 49 131 2.1. Plant material and experimental conditions

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52 132 Two-month-old seedlings of *Moringa oleifera* Lam. were planted in 50 L pots with a sand/peat
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54 133 substrate (9/1, v/v), and were grown outside in Florence, Italy (43° 49' N, 11° 37'). The experiment
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56 134 was conducted during summer 2014, under minimum/maximum temperatures of $17.7 \pm 2.4/30.8 \pm$

1 135 3.2°C (mean \pm standard deviation, S.D.) and midday irradiance (measured over the 200-3000 nm
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3 136 range of solar wavebands) of $780 \pm 85 \text{ W m}^{-2}$ (mean \pm S.D). Saplings were irrigated to pot capacity
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5 137 before the onset of water stress treatment, that was applied to plants on average ~ 190 cm tall and
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7 138 with stem diameter of ~ 2.0 cm. Water stress was imposed by withholding water for 30 days (WS,
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9 139 water-stressed plants), whereas control plants (C) were irrigated daily to pot capacity. A total of thirty
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11 140 plants were grown under these two experimental conditions (12 assigned to well-watered treatment
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13 and 18 assigned to water-stressed treatment). Plants were assigned on the basis of preliminary leaf
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15 141 gas exchange measurements to exclude significant differences in photosynthesis (A_N) and stomatal
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17 142 conductance (g_s) among plants ($t < 0.05$, data not shown). The fraction of transpirable soil water
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19 143 (FTSW) and g_s were used as water stress indicators (Sinclair and Ludlow, 1986; Brill et al., 2013).
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21 144 Measurements were conducted in water-stressed plants at increasing stress severity. The three stress
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23 145 levels corresponded to decreasing FTSW from 100% (in control plants, FTSW₁₀₀), to 60% (FTSW₆₀),
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25 146 40% (FTSW₄₀) and 25% (FTSW₂₅), corresponding to 10, 20 and 30 days after withholding water,
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27 147 respectively. Control plants were also sampled at the same days as water-stressed plants, to make sure
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29 148 that control conditions were maintained across the experimental period. The physiological lower limit
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31 149 of available soil water, corresponding to the FTSW endpoint, was calculated prolonging water stress,
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33 150 until stomatal conductance approached zero, on some additional plants.
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36 152 As *M. oleifera* has bipinnate compound leaves, water relations, gas exchange, chlorophyll
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38 153 fluorescence, isoprene and *n*-hexanal measurements were conducted on the two medial leaflets in the
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40 154 secondary pinna (hereafter denoted as leaf), on four replicate plants per treatment, at each sampling
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42 155 date. The adjacent leaf was collected for biochemical measurements, between 12:00-14:00 h.
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45 156 2.2. Growth, water relations, gas exchange and chlorophyll fluorescence

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48 157 Biomass was measured at the end of the water stress period (30 d) on ten replicate plants per
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50 158 treatment. Plants were divided into shoots and roots and oven dried at 70 °C until a constant weight
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52 159 was reached (after about 72 h). Biomass allocation was calculated on a dry mass (DM) basis, using as
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54 160 parameters the ratio of shoot dry mass to total dry mass (BAS) and the ratio of root dry mass to total
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56 161 dry mass (BAR). Predawn measurements of relative water content (RWC), water (ψ_w) and osmotic

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2 162 (ψ_{π}) potentials were made on well-watered and water-stressed leaves (2 leaves for each selected
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4 163 replicate).

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6 164 **Gas exchange was** measured on intact leaves using a LI-6400 portable photosynthesis system
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8 165 (Li-Cor, Lincoln, NE, USA). Measurements were performed at a photosynthetic photon flux density
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10 166 (PPFD) of 1000 $\mu\text{mol photons m}^{-2} \text{ s}^{-1}$, a CO_2 concentration of 400 $\mu\text{mol mol}^{-1}$ and **ambient**
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12 167 **temperature**. This system was utilized also to measure leaf temperature. Photosynthesis and g_s were
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14 168 calculated using the LI-6400 software. Chlorophyll (Chl) fluorescence was measured **using a**
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16 169 **modulated PAM-2000 fluorometer** (Heinz Walz, Effeltrich, Germany). Minimum fluorescence (F_0) was
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18 170 measured with a 0.8 $\mu\text{mol m}^{-2} \text{ s}^{-1}$ measuring light beam on leaves that were dark-adapted for **20**
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20 171 **minutes**. Maximum fluorescence in the dark-adapted state (F_m) was determined using a saturating
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22 172 pulse (0.5 s) of red light (8000 $\mu\text{mol m}^{-2} \text{ s}^{-1}$), thus allowing calculation of $F_v/F_m = (F_m - F_0)/F_m$. Actinic
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24 173 red continuous light (1000 $\mu\text{mol m}^{-2} \text{ s}^{-1}$) was then switched on, and steady-state fluorescence was
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26 174 recorded (F_s). Saturating pulses were then applied to record the maximum fluorescence under actinic
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28 175 light (F'_m). These data were used to calculate non-photochemical quenching ($\text{NPQ} = (F_m - F'_m)/F'_m$)
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30 176 (Schreiber et al., 1986), actual quantum yield of PSII ($\Phi_{\text{PSII}} = (F'_m - F_s)/F'_m$) (Genty et al. 1989), and
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32 177 electron transport rate ($\text{ETR} = 0.5 \cdot \Phi_{\text{PSII}} \cdot \text{PAR} \cdot 0.84$), where 0.5 and 0.84 are coefficients indicating
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34 178 an equal distribution of photons between PSI and PSII and leaf absorptance, respectively.

35 36 37 179 2.3. Isoprene, abscisic acid and photosynthetic pigments

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40 180 To measure isoprene emission, the outlet of the cuvette was disconnected from the LI-6400
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42 181 system and the flow was diverted into a silcosteel cartridge packed with 200 mg of Tenax (Agilent,
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44 182 Cernusco sul Naviglio, Italy). A volume of 4.5 dm^3 of air was pumped through the trap at a rate of 200
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46 183 $\text{cm}^3 \text{ min}^{-1}$. The cartridge was analysed using a Perkin Elmer Clarus 580 gas chromatograph coupled
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48 184 with a Clarus 560 Mass-Selective-Detector and a thermal desorber TurboMatrix (Perkin Elmer Inc.,
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50 185 Waltham, MA, USA). The desorbed compounds were separated in a 30-m Elite-5-MS capillary column.
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52 186 The column oven temperature was kept at 40 $^{\circ}\text{C}$ for the first 5 min, then increased by 5 $^{\circ}\text{C min}^{-1}$ to 250
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54 187 $^{\circ}\text{C}$, and maintained at 250 $^{\circ}\text{C}$ for 2 min. Helium was used as carrier gas. Compounds were identified
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56 188 using the NIST library provided with the GC/MS Turbomass software. Quantification of isoprene was

1
2 189 conducted using authentic standards of isoprene (Rivoira, Milan, Italy) to prepare a calibration curve
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4 190 as well as to compare the peak retention time and the peak fragmentation of isoprene found in the
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6 191 samples.

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8 192 Abscisic acid, both in its free (free-ABA) and conjugated form (ABA glucoside ester, ABA-GE),
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10 193 was extracted and quantified as reported in Tattini et al. (2017). In detail, 200 mg of lyophilized leaf
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12 194 tissue were ground in liquid nitrogen and combined with 50 ng of d⁶-ABA and d⁵-ABA-GE (National
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14 195 Research Council of Canada), then extracted with 3 × 1 cm³ pH 2.5 CH₃OH/H₂O (50:50; v:v), at 4 °C for
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16 196 30 minutes. The supernatant, after defatting with 3 × 3 cm³ of *n*-hexane, was purified using Sep-Pak
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18 197 C18 cartridges (Waters, Milford, MA, USA) and eluted with 1 cm³ of ethylacetate. The eluate, dried
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20 198 under nitrogen, and rinsed with 500 μL pH 2.5 CH₃OH/H₂O (50:50), was injected (3 μL aliquots) in a
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22 199 LC-DAD-MS/MS system, consisting of a Shimadzu Nexera HPLC and a Shimadzu LCMS-8030
23
24 200 quadrupole mass spectrometer, operating in electrospray ionization (ESI) mode (Kyoto, Japan). The
25
26 201 eluting solvents consisted of H₂O (added with 0.1 % of HCOOH, solvent A) and CH₃CN/CH₃OH (1:1, v:v,
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28 202 added with 0.1 % of HCOOH, solvent B). The analysis was performed in negative ion mode, using a 3 ×
29
30 203 100 mm Poroshell 120 SB C18 column (2.7 μm, 100 × 4.6 mm, Agilent Technologies) and eluting a 18
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32 204 min-run from 95% solvent A to 100% solvent B at a flow rate of 0.3 cm³ min⁻¹. Quantification was
33
34 205 conducted in multiple reaction mode (MRM), as reported by López-Carbonell et al. (2009).

35
36 206 Chlorophyll *a* and *b*, and individual carotenoids were identified and quantified as reported by
37
38 207 Beckett et al. (2012). Briefly, lyophilized leaf tissue (0.2 g) was extracted with 3 × 5 cm³ acetone
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40 208 (added with 0.5 g cm⁻³ of CaCO₃) and injected (15 μL) in a Flexar high performance liquid
41
42 209 chromatography (HPLC) system equipped with a quaternary 200Q/410 pump and a LC 200 diode
43
44 210 array detector (DAD) (all from Perkin Elmer Bradford, CT, USA). Photosynthetic pigments were
45
46 211 separated in a 250 × 4.6 mm Agilent Zorbax SB-C18 (5 μm) column operating at 30°C, eluted for 18
47
48 212 min with a linear gradient solvent system, at a flow rate of 1 cm³ min⁻¹, from 100% CH₃CN/MeOH
49
50 213 (95/5 with 0.05% of triethylamine) to 100% MeOH/ethyl acetate (6.8/3.2). Violaxanthin cycle
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52 214 pigments [violaxanthin (Vio), antheraxanthin (Ant), zeaxanthin (Zea), collectively named (VAZ)],
53
54 215 neoxanthin (Neo), lutein (Lut), β-carotene (β-car), chlorophyll *a* and chlorophyll *b* were identified
55
56 216 using visible spectral characteristics and retention times. Carotenoids and chlorophylls were

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2 217 calibrated using authentic standards from Extrasynthese (Lyon-Nord, Genay, France) and from Sigma
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4 218 Aldrich (Milan, Italy), respectively. The de-epoxidation state of VAZ (DES) was calculated as:

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6 219
$$\text{DES} = (0.5A + Z)/(V + A + Z)$$

7

8 9 220 2.4. Flavonoids

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11 221 Individual flavonoids were extracted and quantified as previously reported in Tattini et al.
12
13 222 (2015). Briefly, lyophilized leaf tissue (0.2 g) was extracted with $3 \times 5 \text{ cm}^3$ of 75% EtOH/H₂O adjusted
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15 223 to pH 2.5 with formic acid, and the supernatant partitioned with $4 \times 5 \text{ cm}^3$ *n*-hexane, reduced to
16
17 224 dryness and finally rinsed with 2 cm^3 CH₃OH/H₂O (8:2, v:v). Aliquots of 10 μL were injected into the
18
19 225 Perkin Elmer liquid chromatography system reported above, and compounds separated in a 150×4.6
20
21 226 mm Sun Fire column (5 μm) (Waters Italia, Milan, Italy) operating at 30 °C and eluted at a flow rate of
22
23 227 $1 \text{ cm}^3 \text{ min}^{-1}$. The mobile phases were (A) H₂O pH 4.3 (CH₃COONH₄/CH₃COOH)/CH₃CN (90/10, v/v) and
24
25 228 (B) CH₃CN/H₂O pH 4.3 (CH₃COONH₄ /CH₃COOH) (90/10, v/v). Flavonoids were separated using a
26
27 229 linear gradient elution from A to B over a 46 min-run. Flavonoids were identified by comparison of
28
29 230 their retention times and UV spectral characteristics with those of authentic standards (Extrasynthese,
30
31 231 Lyon-Nord, Genay, France) and quantified at 350 nm using five-point calibration curves of authentic
32
33 232 standards.
34
35

36 37 233 2.5. Lipid peroxidation indicator (*n*-hexanal)

38
39
40 234 *N*-hexanal is one of the lipid peroxide-derived carbonyl compounds (oxylipin carbonyls) that
41
42 235 reveals abiotic stress-induced damage of plants, and in particular of cellular membranes (Mano 2012).
43
44 236 Analysis of *n*-hexanal was done using the same procedure as for isoprene (see above). Quantification
45
46 237 of *n*-hexanal was conducted using an authentic standard (Sigma Aldrich, Milan, Italy) to prepare a
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48 238 calibration curve, as well as comparing the peak retention time and the peak fragmentation in all
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50 239 samples.
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52 53 240 2.6. Experimental design and statistics

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1
2 241 The experiment was performed using a completely randomized design. Biomass was measured
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4 242 on ten replicates for both well-watered and water-stressed plants at the end of the experiment.
5
6 243 Physiological and biochemical measurements were conducted on four replicate plants, both in well-
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8 244 watered plants and in plants exposed to water stress of increasing severity. Data were analysed using
9
10 245 repeated-measures ANOVA, with water treatment as between-subjects effect and sampling date as
11
12 246 within-subjects effect (SPSS v.20; IBM, Chicago IL, USA). Significant differences among means were
13
14 247 estimated at the 5% ($P < 0.05$) level, using Tukey's test.

15 248

16 249 **3. Results**

17 250

18 251 **3.1. Water stress effects on water relations, photosynthesis and biomass production**

19
20 252 Predawn leaf water potential (ψ_w , Fig. 1A) declined in water-stressed plants compared to
21
22 253 control plants, though differences became significant only at FTSW₄₀ and FTSW₂₅. It is noteworthy
23
24 254 that at the end of the water stress cycle, when g_s of FTSW₂₅ plants was on average about 15% of the
25
26 255 control values, predawn ψ_w was still rather high (i.e., -0.60 MPa). Significant differences in leaf bulk
27
28 256 osmotic potential (ψ_π , Fig. 1B) were recorded only at the end of the experiment (FTSW₂₅), whereas
29
30 257 RWC did not significantly vary between water-stressed and control plants (Fig. 1C).

31
32 258 As FTSW declined, A_N , g_s , and C_i (Fig. 2; Tab. SM1) were progressively reduced. A strong
33
34 259 reduction of A_N (-30%, Fig. 2A) and g_s (-43%, Fig. 2B) was observed already under mild water stress
35
36 260 (FTSW₆₀). Under a more severe water stress (FTSW₂₅), A_N and g_s declined by 71% and 85%
37
38 261 respectively, compared to control leaves (FTSW₁₀₀) (Tab. SM1). Similarly, C_i (Fig. 2C) was reduced by
39
40 262 about 55% in FTSW₂₅ plants relative to FTSW₁₀₀ plants. The maximum quantum yield of PSII (F_v/F_m ,
41
42 263 Fig. 2D) did not vary between control and water-stressed plants, irrespective of the severity of the
43
44 264 stress. In contrast, the actual efficiency of PSII photochemistry (Φ_{PSII}), significantly declined already at
45
46 265 FTSW₆₀ and was further impaired at FTSW₄₀ and FTSW₂₅, relative to FTSW₁₀₀ plants (Fig. 2E). Water
47
48 266 stress reductions in Φ_{PSII} were paralleled by corresponding increases in the non-photochemical
49
50 267 quenching of fluorescence (NPQ, Fig. 2F and SM1).

1
2 268 Plant biomass was significantly reduced in FTSW₂₅ compared to FTSW₁₀₀ plants (Fig. 3A). At
3
4 269 the end of the experiment, the root to shoot ratio **was** also significantly higher in FTSW₂₅ than in
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6 270 FTSW₁₀₀ plants, whereas the shoot to total dry mass ratio was significantly reduced in water-stressed
7
8 271 plants (Fig. 3B).
9

10
11 272 3.2. Water stress effects on isoprene, non-volatile isoprenoids, pigments, flavonoids, and
12
13 273 membrane lipid peroxidation.
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15
16 274 Isoprene emission increased **significantly** in FTSW₁₀₀ leaves during the experiment (Fig. 4A),
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18 275 likely because of the prolonged exposure to elevated temperatures during the summer season.
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20 276 Isoprene emission strongly and significantly increased in response to water stress. This increment was
21
22 277 particularly relevant at FTSW₂₅ (+86% compared to FTSW₄₀). The carbon lost as isoprene (C_{iso} %), also
23
24 278 increased largely in **FTSW₂₅ plants, due to the simultaneous increase of isoprene emission and**
25
26 279 **reduction of A_n** (Fig. 4B). The surging emission of isoprene was positively correlated to both the
27
28 280 decline of C_i (Fig. 5A) and the increase of the ETR/ A_n ratio (Fig. 5B) in water-stressed leaves.
29

30
31 281 The content of free-ABA and ABA-GE increased in water-stressed compared to control leaves
32
33 282 (Fig. 6A and 6B), and the effect was particularly strong in FTSW₂₅ plants where free-ABA and ABA-GE
34
35 283 contents were about seven and two folds higher than in FTSW₁₀₀, respectively. A strong negative linear
36
37 284 relationship ($R^2 = 0.915$) was found between foliar free-ABA levels and g_s (inset of Fig. 6A). Whereas,
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39 285 free and conjugated ABA contents were both positively related to isoprene emission rates (inset of Fig.
40
41 286 6B).
42

43
44 287 Total chlorophyll (Chl_{tot}) declined significantly in FTSW₄₀ (-14%) and FTSW₂₅ (-23%) leaves in
45
46 288 comparison to FTSW₁₀₀ leaves (Fig. 7A). In contrast, total carotenoid (Car_{tot}) content did not vary
47
48 289 between control and water-stressed leaves, and increased during the experiment irrespective of water
49
50 290 treatments (Fig. 7B). However, water stress markedly altered the composition of the carotenoid pool.
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52 291 The content of Lut (Fig. 7C) increased, whereas the content of β -car (Fig. 7D) declined significantly
53
54 292 under severe water deficit (FTSW₂₅). Among xanthophylls, Vio (Fig. 7E) declined and Zea (Fig. 7F)
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56 293 increased significantly in water-stressed plants. Vio reduction was particularly strong at FTSW₆₀ and
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58 294 FTSW₄₀, and partially recovered under severe stress conditions (FTSW₂₅). The contents of Ant (Fig. 7G)

1
2 295 and Neo (Fig. 7H) were not affected by water stress. However, Neo increased during the experimental
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4 296 period in both well-watered and water-stressed plants. The content of violaxanthin-cycle pigments
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6 297 (VAZ) relative to Chl_{tot} increased significantly as water stress progressed, and the effect was
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8 298 particularly high (+35%) at FTSW₂₅ compared to FTSW₁₀₀ after 30 days of water stress (Fig. 7I). In
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10 299 addition, DES increased in water-stressed compared to control leaves, but the difference was already
11
12 300 noticeable under mild water stress conditions (FTSW₆₀) (Fig. 7J).

13
14 301 Water stress also considerably altered the content and composition of the flavonoid pool (Fig.
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16 302 8 A-C). Quercetin-3-*O*-glycoside and its derivatives were the most responsive compounds to water
17
18 303 stress, as their content significantly and consistently increased with the intensity of the stress (Fig.
19
20 304 8A). In addition, the content of Kaempferol-3-*O*-glycoside derivatives also significantly increased in
21
22 305 response to stress, but the difference between water-stressed and control leaves remained constant as
23
24 306 water stress progressed (Fig. 9B). In contrast, the content of Apigenin-7-*O*-glycoside and its
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26 307 derivatives significantly decreased in FTSW₄₀ and FTSW₂₅ leaves (Fig. 8C).

27
28 308 Compared to control leaves, the emission of *n*-hexanal did not significantly vary at both
29
30 309 FTSW₆₀ and FTSW₄₀, whereas it significantly increased in FTSW₂₅ plants (Fig. 9).

31
32 310

33 34 311 4. Discussion

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36
37 312 4.1 Understanding the impact of water stress on the physiology of the isohydric plant *M.*
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39 313 *oleifera*

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41
42 314 *M. oleifera* is a fast-growing species able to produce large quantities of biomass (Sánchez et al. 2006).
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44 315 However, whether *M. oleifera* is able to acclimate and produce at satisfactory rates in arid conditions is
45
46 316 yet not known. Our study offers novel insights on the physiological and biochemical strategies adopted
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48 317 by this species to cope with extended periods of soil water stress.

49
50 318 Our results show that *M. oleifera* possesses an effective avoidance mechanism (i.e. isohydry,
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52 319 Nardini et al., 2014; Tardieu and Simmoneau, 1998) when subjected to water stress. This involved a
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54 320 rapid reduction of g_s in stressed leaves, that possibly contributed to the maintenance of ψ_w and RWC
55
56 321 even in conditions of severe water stress (FTSW₂₅), when only a moderate (8%) reduction of the

1
2 322 osmotic component ψ_{π} became significant (Fig. 1 and 2B). The response of g_s of *M. oleifera* to soil
3
4 323 drying (Tab. SM1) is remarkably different from that observed in other fast-growing trees species such
5
6 324 as *Eucalyptus citriodora* (Brilli et al. 2013; Mahmood et al. 2015) and *Populus spp* (Marron et al. 2002;
7
8 325 Yin et al. 2005; Brilli et al. 2007; Centritto et al. 2011) that showed no or very little decline in g_s under
9
10 326 moderate water stress conditions. Isohydry is a crucial adaptive trait for the survival of deciduous
11
12 327 woody plants exposed to high evaporative demand and low soil water availability, as an early and tight
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14 328 control of stomatal aperture may prevent xylem embolism (Franks et al. 2007; Yi et al. 2017). While
15
16 329 stomatal closure increased intrinsic water use efficiency (iWUE, determined as the ratio of A_n to g_s)
17
18 330 during water stress progression, it also constrained photosynthesis due to increased diffusional
19
20 331 limitations to CO_2 entry, with consequent reduction of C_i (Fig. 2C) (Lawlor and Cornic 2002; Centritto
21
22 332 et al. 2011; Lauteri et al. 2014; Fini et al. 2016). The observed drop in photosynthesis under water
23
24 333 stress caused a biomass reduction (Fig. 2 and 3), probably inducing a redistribution of the assimilated
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26 334 carbon between shoots and roots (Peuke et al. 2006). These results suggest a high degree of plasticity
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28 335 of *M. oleifera* in biomass allocation in response to water stress (Fig. 3).

30 336 Water stress did not cause permanent damages to the photosynthetic apparatus. In fact,
31
32 337 maximal PSII photochemical efficiency (F_v/F_m) did not decline even under severe water stress
33
34 338 (FTSW₂₅), suggesting stability of photochemical reactions and structures (Fig. 2) (Flexas et al., 2006).
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36 339 However, PSII quantum yield in the light (Φ_{PSII}) was reduced as compared to FTSW₁₀₀ leaves. While
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38 340 this mirrored A_n reduction at mild (FTSW₆₀) and moderate (FTSW₄₀) stress level, Φ_{PSII} did not drop
39
40 341 further in severely water-stressed leaves (Havaux 1992; Lu and Zhang, 1999) revealing a likely
41
42 342 increase of photorespiratory electron transport, or alternative electron sinks (see discussion below
43
44 343 about ETR driving isoprene emission). Furthermore, changes in NPQ and Φ_{PSII} were strongly
45
46 344 correlated throughout the experiment ($\Phi_{PSII} = -0,13 \text{ NPQ} + 0.62$, $R^2 = 0.844$, linear relation shown in
47
48 345 Fig. SM1). Large excess of light energy not used by photosynthesis, as revealed by the fluorescence
49
50 346 parameter NPQ (Fig. 2F), may directly photoreduce O_2 , thus causing large ROS generation in water-
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52 347 stressed leaves, with consequent damage to PSII. To explain why this was not observed in this
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54 348 experiment, we hypothesize a potential contribution of isoprenoids and phenylpropanoids as
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56 349 antioxidant compounds, as discussed below.

1
2 350 4.2. Exploring the significance of enhanced isoprene emission during water stress and its
3
4 351 relationship with foliar ABA

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6 352 Our study revealed that *M. oleifera* is an isoprene emitting species (Fig. 4). Isoprene emission is
7
8 353 typical of hygrophytes that are fast-growing in temperate areas of the world (Loreto et al. 2014; Loreto
9
10 354 and Fineschi 2015), where isoprene serves important defensive (antioxidant and thermo-protective)
11
12 355 properties (Loreto and Schnitzler 2010; Velikova et al. 2011; Pollastri et al. 2014). We also show that
13
14 356 water stress promoted I_e , particularly when the stress became severe. Isoprene biosynthesis is
15
16 357 generally resistant to water stress (Brilli et al. 2007; Centritto et al. 2011; Brilli et al. 2013), and the
17
18 358 emission of isoprene is enhanced when isoprene-emitters recover from water stress (Sharkey and
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20 359 Loreto, 1993; Fortunati et al., 2008). Stimulation of isoprene biosynthesis “during” water stress
21
22 360 episodes is less reported (Haworth et al. 2017; Marino et al. 2017). *M. oleifera* is a typical isoprene
23
24 361 emitting species, since it is a fast-growing species with high photosynthetic rates which thrives wild in
25
26 362 secondary tropical deciduous forests of the sub-Himalayan area (Loreto and Fineschi, 2015). Our data
27
28 363 suggest that declines in internal CO_2 concentration (C_i) and the increasing electron flux generated by
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30 364 Photosystem II not used for carbon assimilation (ETR/A_N) are two important physiological drivers of
31
32 365 isoprene biosynthesis under water stress conditions (Fig. 5) (Guidolotti et al. 2011; Harrison et al.
33
34 366 2013; Morfopoulos et al. 2014; Marino et al. 2017). **Reduced photosynthesis due to CO_2 starvation may**
35
36 367 **indeed increase the fraction of ETR available for alternative biosyntheses, including isoprenoids.** In
37
38 368 addition, the increase in leaf temperature induced by stomatal closure under water stress (from $31.2 \pm$
39
40 369 0.7 °C in FTSW₁₀₀ leaves to 34.4 ± 0.6 °C in FTSW₂₅ leaves, mean \pm S.D.) might have contributed to
41
42 370 further enhance the rate of isoprene emission (Singsaas and Sharkey, 1998; Fares et al. 2011; Brilli et
43
44 371 al. 2013; Arab et al. 2016). Indeed, **the activity of isoprene synthase is known to be stimulated by high**
45
46 372 **temperatures (Monson et al. 1992; Li et al. 2011). Increasing isoprene synthase activity may also help**
47
48 373 **explain the increase in I_e and $C_{iso}\%$ observed in well-watered leaves, along rising summer**
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50 374 **temperatures during the course of our study (Rasulov et al. 2015).**

51
52 375 We hypothesize that the rising investment of **newly assimilated** carbon for isoprene
53
54 376 biosynthesis helped leaves tolerate water stress because: a) isoprene protects the photosynthetic
55
56 377 apparatus from heat and oxidative damage by preserving the integrity of thylakoid membranes (Siwko

1
2 378 et al. 2007; Velikova et al. 2011, 2014, 2015) or by scavenging singlet oxygen (1O_2), a highly reactive
3
4 379 ROS in chloroplasts (Velikova et al. 2004; Zeinali et al. 2016); b) isoprene makes faster and smoother
5
6 380 the electron transport flow (Pollastri et al. 2014), especially under water stress conditions (Marino et
7
8 381 al. 2017). We found that NPQ did not vary between FTSW₄₀ and FTSW₂₅ leaves. Lower NPQ values in
9
10 382 isoprene emitters compared to non-emitters were reported both in stressful (Behnke et al. 2007,
11
12 383 2010) and physiological conditions (Pollastri et al. 2014). We, therefore, hypothesize a relationship
13
14 384 between the reduction of NPQ and the increase I_e along with the severity of water stress. A
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16 385 downregulation of chloroplastic ATP-synthase and the consequent reduction in the flexible heat
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18 386 dissipation component (qE) of NPQ (Demmig-Adams and Adams 2006) was reported in isoprene
19
20 387 emitting species by Velikova et al. (2014).

21
22 388 The observed strong linear relationships between I_e and foliar contents of free-ABA and ABA-
23
24 389 GE (Fig. 6B), suggest that increased isoprene formation in water stressed plants indicates enhanced
25
26 390 carbon flow through the MEP pathway, leading to higher foliar biosynthesis of abscisic acid (Fig. 6A)
27
28 391 (Marino et al. 2017). A relationship between isoprene and foliar ABA was first reported by Barta and
29
30 392 Loreto (2006) in well-watered *Populus alba* and by Tattini et al. (2014) in drought stressed transgenic
31
32 393 tobacco plants. Our results also show a strong linear correlation between free-ABA and g_s (Fig. 6A),
33
34 394 despite limited variations of water relations in *M. oleifera* leaves. It is unclear whether isoprene is
35
36 395 simply of proxy of carbon flux through the MEP pathway, or has a regulatory role. Sustained isoprene
37
38 396 emission in water-stressed plants may reduce the accumulation of dimethylallyl pyrophosphate
39
40 397 (DMAPP) in the chloroplast, and may prevent DMAPP-induced feedback inhibition of the entire MEP
41
42 398 pathway (Banerjee et al. 2013). Taken together our results suggest that: a) increased isoprene
43
44 399 formation indicates and perhaps regulates free-ABA synthesis in stressed leaves, and b) free-ABA has a
45
46 400 major role in the regulation of stomatal closure compared to hydraulic signals (Chaves et al. 2016;
47
48 401 McAdam et al. 2016a). These results are in line with recent studies showing that, in strict isohydric
49
50 402 plants such as *M. oleifera*, high levels of free-ABA could be responsible for stomatal closure and could
51
52 403 promote a higher root to shoot ratio/carbon allocation (Nolan et al. 2017; McAdam et al. 2016b).

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54
55 404 4.3. Plasticity of secondary metabolism in *M. oleifera* during water stress progression

1
2 405 We observed several changes in carotenoids and phenylpropanoids in response to increasing
3
4 406 water stress, that can be interpreted as a photoprotective trait to limit water stress induced damage.
5
6 407 The content of total carotenoids on a leaf mass basis also increased **over the course of the experiment**
7
8 408 in both well-watered and water-stressed leaves. While this shows a general upregulation of the MEP
9
10 409 pathway (see previous section) over the season, we argue that the investment **in** carotenoids was
11
12 410 much stronger in water-stressed leaves **mirroring the depression in carbon assimilation**. The blend of
13
14 411 carotenoids also changed along stress progression, perhaps favouring compounds active in stress
15
16 412 protection (Fig. 7). The increase in lutein in severely water-stressed plants might have enhanced the
17
18 413 capacity of leaves to quench $^3\text{Chl}^*$, that was likely generated during stress exposure (Dall'Osto et al.
19
20 414 2006; Jahns and Holzwarth 2012). In addition, compared to photosynthesis, Chl_{tot} content was less
21
22 415 affected by severe water stress, indicating a successful mechanism of protection. We also note that a
23
24 416 large switch in the composition of xanthophylls occurred in water-stressed plants. The increase in Zea
25
26 417 content was accompanied by a parallel decrease in Vio content under mild and moderate water stress,
27
28 418 showing the classic mechanism of de-epoxidation that is a major element of photoprotection in plants
29
30 419 (Demming-Adams and Adams 2006). **However, when plants experienced the most severe water stress**
31
32 420 **the content in Zea and in Vio both increased. We suggest that the large increase in Zea biosynthesis**
33
34 421 **might have been originated from hydroxylation of β -car (Davison et al. 2002; Du et al. 2010). This is**
35
36 422 **consistent with the reduction of β -car concentration observed in leaves at FTWS₂₅.** β -car might have
37
38 423 been also used as a chemical quencher of $^1\text{O}_2$ (Ramel et al. 2012), thus explaining the relatively
39
40 424 stronger decline of β -car ($-0.18 \mu\text{mol g}^{-1} \text{DW}$) as compared to the increase in Zea ($+0.07 \mu\text{mol g}^{-1} \text{DW}$)
41
42 425 when the stress became severe. The content of VAZ relative to Chl_{tot} was on average $> 70 \text{ mmol mol}^{-1}$
43
44 426 in both well-watered and water-stressed plants throughout the whole experiment, as commonly
45
46 427 observed in leaves long acclimated to full solar irradiance (Fini et al. 2014; Esteban et al. 2015). This
47
48 428 implies that only a fraction of the VAZ pool was bound to antenna systems and, hence, involved in NPQ
49
50 429 (Fig. 7I and J). In addition, the VAZ to Chl_{tot} ratio increased linearly during the water stress cycle. This
51
52 430 increasing 'unbound' VAZ pool might have served specific antioxidant functions in water-stressed
53
54 431 leaves, increasing membrane thermo-stability hence limiting lipid peroxidation (Havaux et al. 2007;
55
56 432 Esteban et al. 2015). This is an action similar to that suggested for isoprene (Velikova et al. 2011), and

1
2 433 cooperation between volatile and non-volatile isoprenoids was surmised by Beckett et al. (2012).
3
4 434 Indeed, the rate of *n*-hexanal emission, a marker of lipid peroxidation (Mano et al. 2012; Beckett et al.
5
6 435 2012), was only affected when a severe water stress was imposed (FTWS₂₅, Fig. 9), and was not
7
8 436 accompanied by irreversible degradation of membrane-bound photosynthetic machineries, namely
9
10 437 PSII photochemistry (as shown earlier).

11
12 438 The biosynthesis of antioxidant flavonoids, here constituted mainly by quercetin derivatives,
13
14 439 was stimulated in water-stressed leaves of *M. oleifera* (Fig. 8), similarly to what has been observed in
15
16 440 other plants (Tattini et al. 2004; Velikova et al. 2016; Ahrar et al. 2017). These high levels of foliar
17
18 441 flavonoids, commonly found in leaves grown under full sunlight, are not compatible with their
19
20 442 exclusive distribution in epidermal cells (Jaakola et al. 2004; Tattini et al. 2005; Agati et al. 2009; Majer
21
22 443 et al. 2014). Therefore, we suggest that water stress induced the accumulation of quercetin derivatives
23
24 444 mainly in mesophyll cells (Tattini et al. 2015), likely conferring increasing protection against enhanced
25
26 445 ROS generation (Agati and Tattini 2010; Agati et al. 2012; Nakabayashi et al. 2014), while reducing the
27
28 446 risk of permanent photodamage to PSII, by additionally acting as UV-B filters in the chloroplast
29
30 447 (Mierziak et al. 2014; Zavafer et al. 2017). The finding that water stress induced profound changes in
31
32 448 the composition of the flavonoid pool, with major increases in the biosynthesis of 'effective
33
34 449 antioxidant' quercetin derivatives (on average +46%), further supports our hypothesis. In contrast,
35
36 450 the content of less effective antioxidant' flavonoids either increased little (kaempferol glycosides,
37
38 451 +15%) or largely declined (apigenin glycosides -35%) in response to water stress. This significant
39
40 452 changes in the composition of flavonoids may also have contributed to reduce lipid peroxidation, as
41
42 453 previously discussed.

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44 454

45 455 **Conclusions**

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49 456 Despite being originated in hygrophilic habitats, *M. oleifera* possesses multiple biochemical
50
51 457 and physiological mechanisms that allow this species to successfully tolerate water stress episodes.
52
53 458 These mechanisms include a strict isohydric behavior in response to water deprivation that is typical
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55 459 of hygrophytes. The fast stomatal closure driven by high contents of foliar-ABA, however, caused an
56
57 460 early and strong depression in carbon assimilation with negative consequences for biomass

1
2 461 production. More interestingly, this study revealed that *M. oleifera* is an isoprene emitting species.
3
4 462 Increasing isoprene emission during progressive water stress was a valuable indicator for the general
5
6 463 activation of the MEP-pathway. The simultaneous increment of volatile and non-volatile isoprenoids
7
8 464 and of flavonoids, is suggested to be the key mechanism that allows *M. oleifera* to **limit lipid**
9
10 465 **peroxidation and prevent severe photoinhibitory processes under water stress. This may allow a**
11
12 466 **prompt recovery of photosynthesis and growth rates when water is newly available to the roots. While**
13
14 467 **the observed high plasticity of stomatal conductance and secondary metabolites production may take**
15
16 468 **its toll on primary productivity of *M. oleifera*, it possibly also facilitates the establishment of this plant**
17
18 469 **to xeric environments. The extent to which the trade-off between primary and secondary metabolism**
19
20 470 **affects the resistance and whole-plant performance of a fast-growing plant such as *M. oleifera*, remains**
21
22 471 **to be determined in presence of recurrent periods of water stress.**
23
24 472

25 26 473 **Authors' contributions**

27
28 474 CB, FL, FF and MT planned the experiment. CB conducted the study, collected samples,
29
30 475 analyzed the data, and prepared the draft. AG, LG and DR helped in performing physiological and
31
32 476 chemical analyses. CB and MT interpreted the results and drafted the manuscript. FL, AF and MC
33
34 477 reviewed the manuscript.
35

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2 790 **Figure and table legends**

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5 792 (RWC, C) in FTSW₁₀₀ (F₁₀₀) plants (open bars) and in FTSW₆₀ (F₆₀), FTSW₄₀ (F₄₀) and FTSW₂₅ (F₂₅)
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23 800 (F₄₀) and FTSW₂₅ (F₂₅) water-stressed plants (grey bars) of *Moringa oleifera*. Data (means \pm SD, n = 4)
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27 802 significantly differ at the 5% level, using Tukey's test.

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33 804 in FTSW₂₅ (F₂₅) water-stressed plants (grey bars) of *Moringa oleifera*. The percentage of biomass
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57 815 *oleifera* plants. Measurements were made at FTSW₆₀ (10 d, open symbols), FTSW₄₀ (20 d, grey

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4 817 water-stressed (circles) plants. Coefficient of determination (R^2) of each relationship are reported; ***
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28 828 Figure 7. Effects of water stress on the contents of photosynthetic pigments (A-I), on the ratio
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30 829 of violaxanthin cycle pigment content to total chlorophyll content (VAZ Chl_{tot}⁻¹, I) and on the de-
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41 834 Figure 8. Contents of quercetin (A), kaempferol (B) and apigenin (C) derivatives in FTSW₁₀₀
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43 835 (F₁₀₀) plants (open bars) and in FTSW₆₀ (F₆₀), FTSW₄₀ (F₄₀) and FTSW₂₅ (F₂₅) water-stressed plants
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54 840 FTSW₄₀ (F₄₀) and FTSW₂₅ (F₂₅) water-stressed plants (grey bars) of *Moringa oleifera*. Data (means \pm

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21 850 and days after the onset of water stress treatment.

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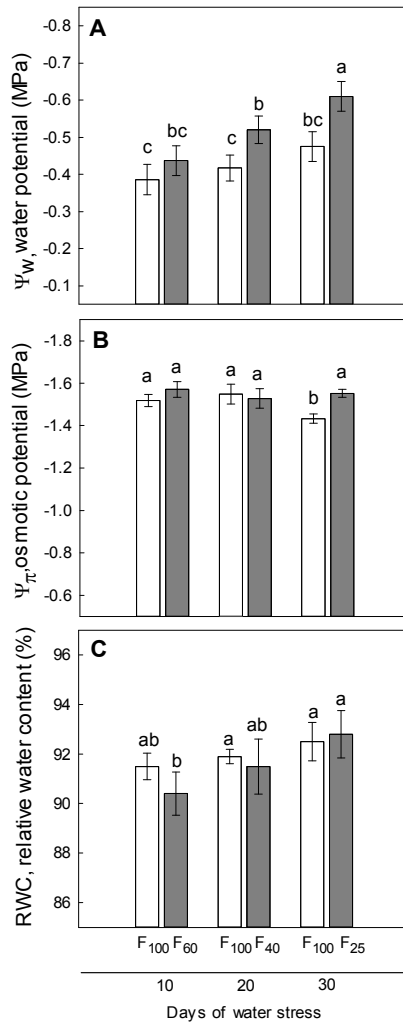
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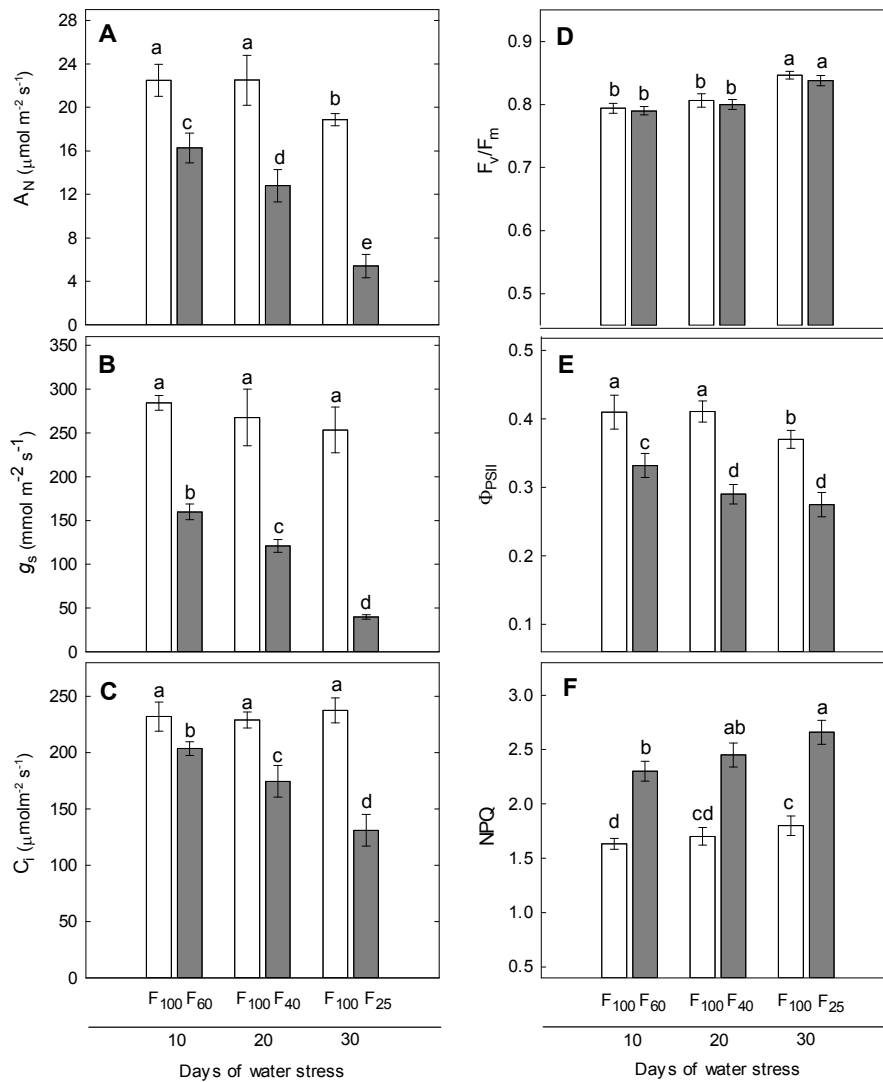
863 **Figures and Tables**



864

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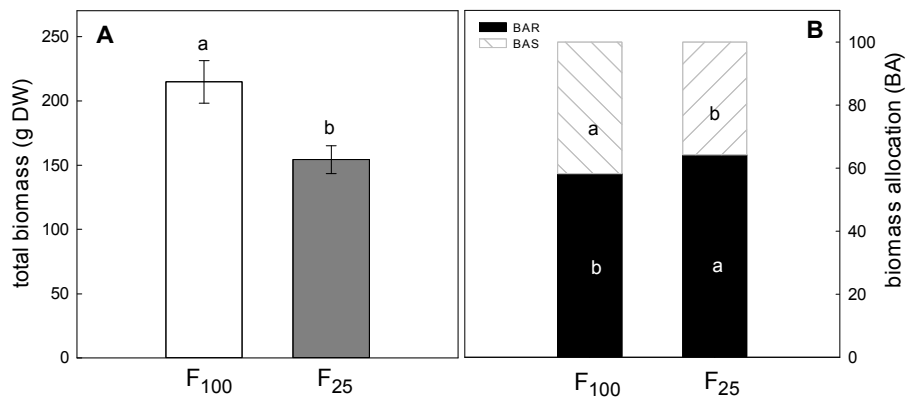
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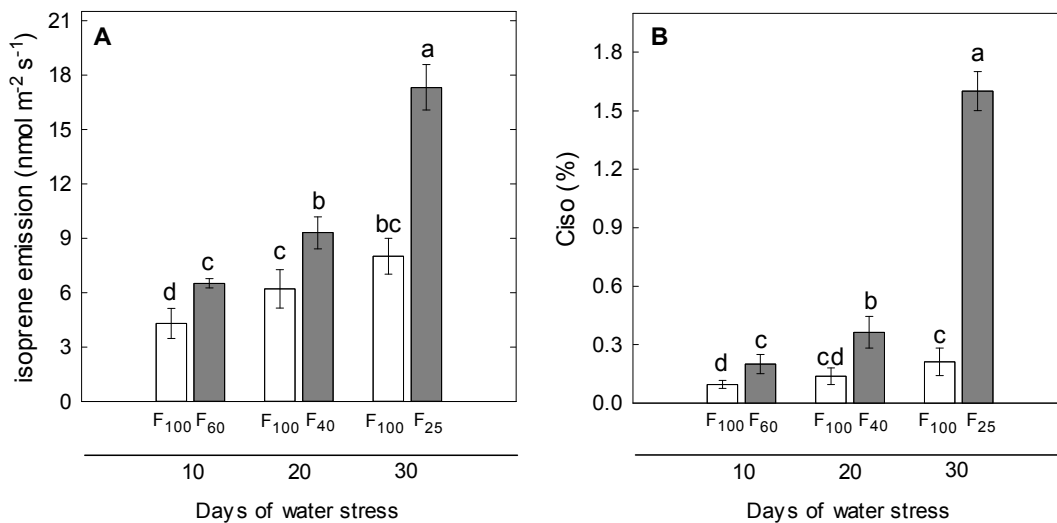
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Figure 2. Photosynthesis (A_N , A), stomatal conductance (g_s , B), intercellular CO_2 concentration (C_i , C), maximum (F_v/F_m , D) and actual (Φ_{PSII} , E) efficiency of PSII photochemistry and non-photochemical quenching (NPQ, F) in FTSW₁₀₀ (F₁₀₀) plants (open bars) and in FTSW₆₀ (F₆₀), FTSW₄₀ (F₄₀) and FTSW₂₅ (F₂₅) water-stressed plants (grey bars) of *Moringa oleifera*. Data (means \pm SD, n = 4) were subjected to repeated measures with ANOVA, and bars not accompanied by the same letter significantly differ at the 5% level, using Tukey's test.



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881 Figure 3. Total biomass (A) and biomass allocation (B) in FTSW₁₀₀ (F₁₀₀) plants (open bars) and
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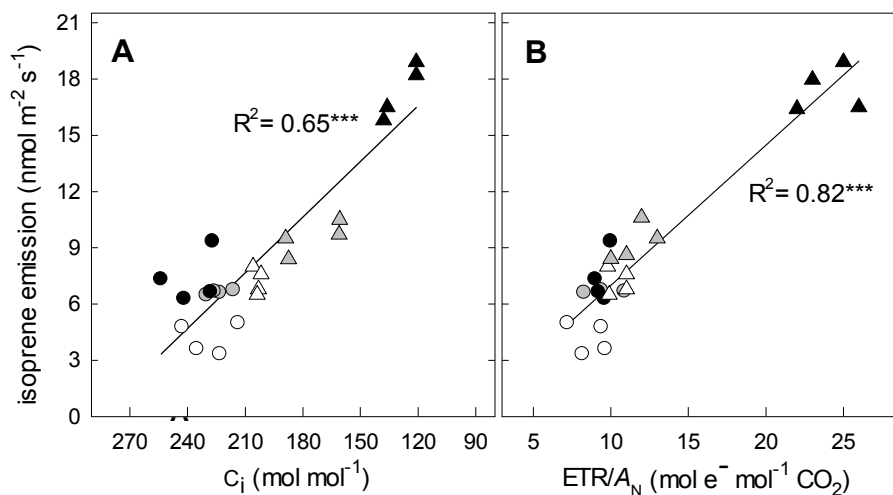
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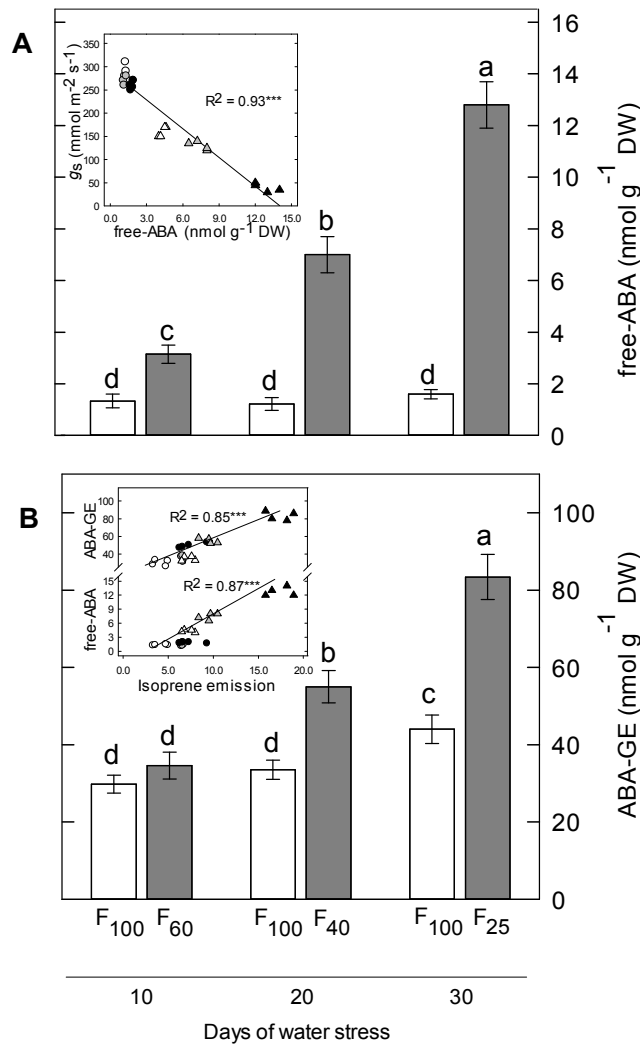
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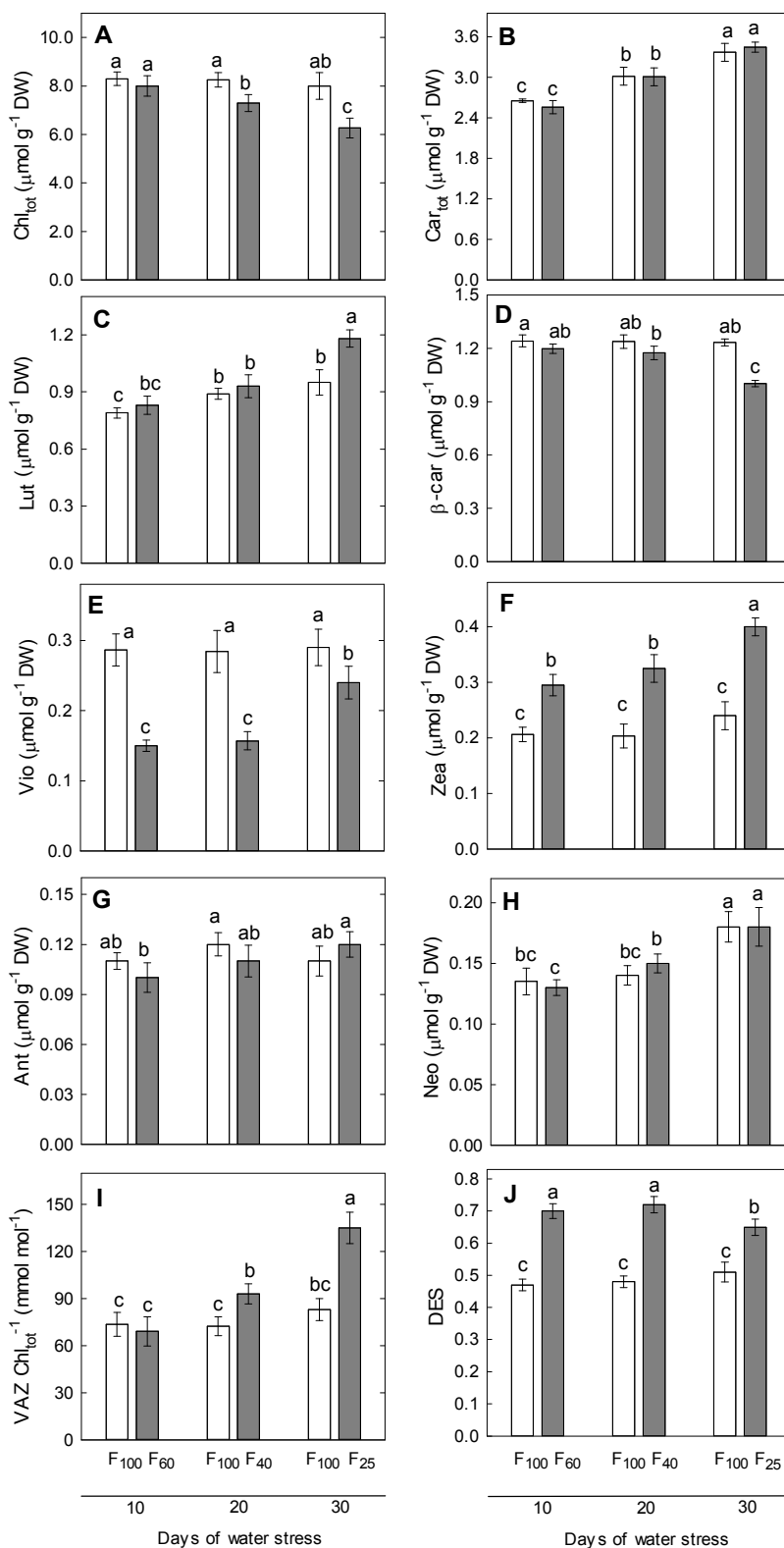


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917 Figure 5. Linear relationships between isoprene emission rate and (A) internal CO₂
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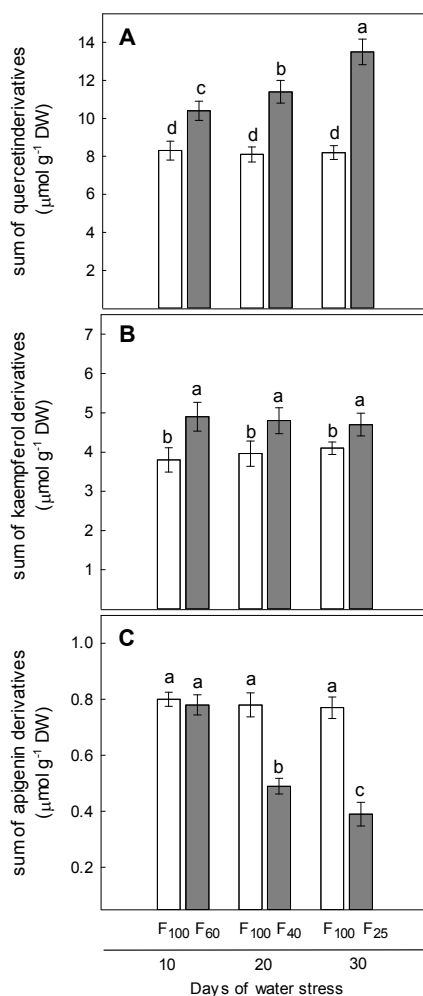
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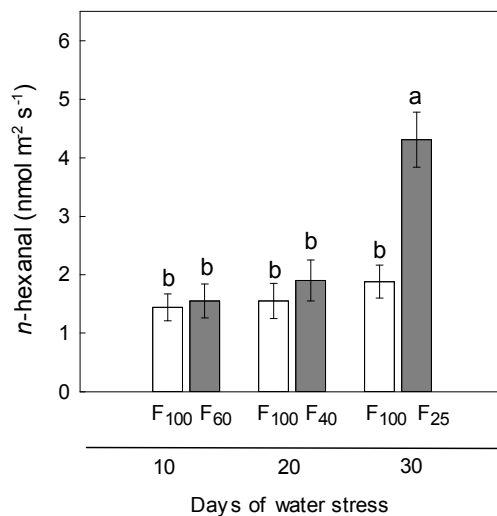
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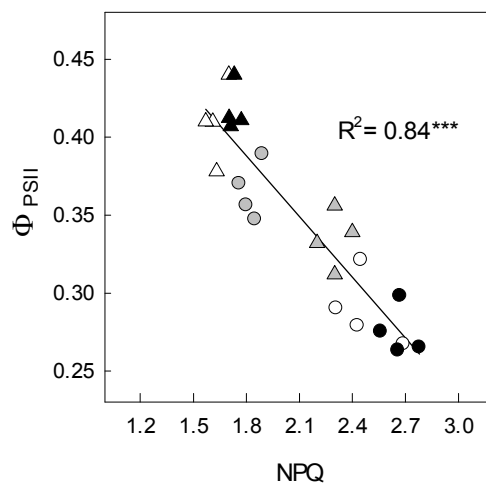
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995 control (FTSW₁₀₀) plants (circles) and water-stressed (triangles) plants of *Moringa oleifera*. Coefficient
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FTSW (%)	A_N ($\mu\text{mol m}^{-2}\text{s}^{-1}$)	g_s ($\text{mmol m}^{-2}\text{s}^{-1}$)	Days of water stress
100	22.5 \pm 1.46	284 \pm 8.51	0
60	16.3 \pm 1.36	160 \pm 9.01	10
40	12.8 \pm 1.52	121 \pm 7.39	20
25	5.4 \pm 1.08	40 \pm 2.53	30

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Table SM 1. Results for photosynthesis (A_N) and stomatal conductance (g_s) (means \pm SD, n = 4)

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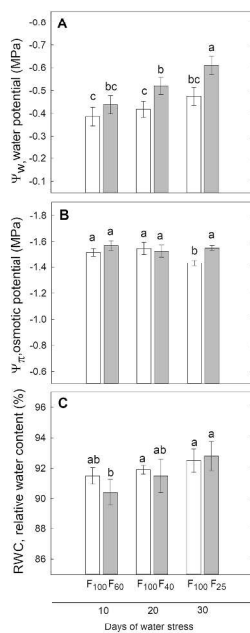


Figure 1. Predawn leaf water (Ψ_w , A) and osmotic (Ψ_π , B) potentials, and relative water content (RWC, C) in FTSW100 (F100) plants (open bars) and in FTSW60 (F60), FTSW40 (F40) and FTSW25 (F25) water-stressed plants (grey bars) of *Moringa oleifera*, corresponding to 10, 20 and 30 days after withholding water, respectively. Data (means \pm SD, n = 4) were subjected to repeated measures with ANOVA, and bars not accompanied by the same letter significantly differ at the 5% level, using Tukey's test.

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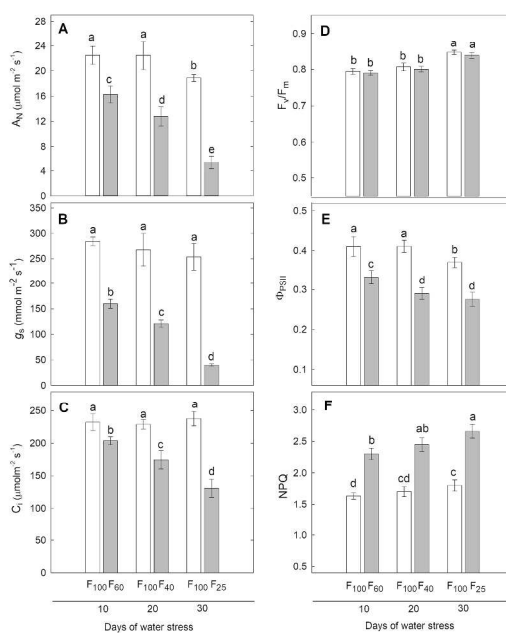


Figure 2. Photosynthesis (A_N , A), stomatal conductance (g_s , B), intercellular CO_2 concentration (C_i , C), maximum (F_v/F_m , D) and actual (Φ_{PSII} , E) efficiency of PSII photochemistry and non-photochemical quenching (NPQ, F) in FTSW100 (F100) plants (open bars) and in FTSW60 (F60), FTSW40 (F40) and FTSW25 (F25) water-stressed plants (grey bars) of *Moringa oleifera*. Data (means \pm SD, $n = 4$) were subjected to repeated measures with ANOVA, and bars not accompanied by the same letter significantly differ at the 5% level, using Tukey's test.

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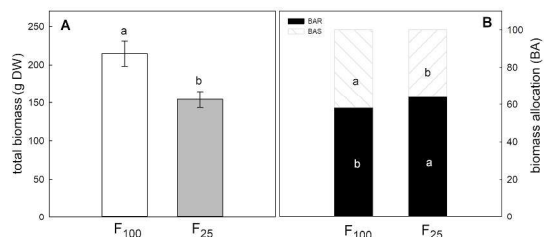


Figure 3. Total biomass (A) and biomass allocation (B) in FTSW100 (F100) plants (open bars) and in FTSW25 (F25) water-stressed plants (grey bars) of *Moringa oleifera*. The percentage of biomass allocation (BA) was calculated considering the ratio of shoot dry mass to total dry mass (BAS) and the ratio of root dry mass to total dry mass (BAR). Data (means \pm SD, n = 10) were subjected to repeated measures with ANOVA, and bars not accompanied by the same letter significantly differ at the 5% level, using Tukey's test.

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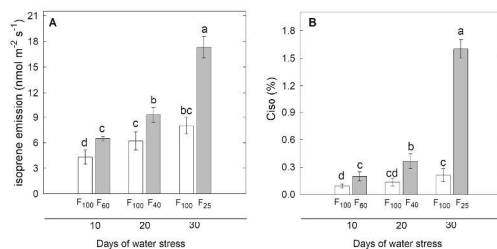


Figure 4. Rates of isoprene emission (A) and carbon lost as isoprene (Ciso, B) in FTSW100 (F100) plants (open bars) and in FTSW60 (F60), FTSW40 (F40) and FTSW25 (F25) water-stressed plants (grey bars) of *Moringa oleifera*. Data (means \pm SD, $n = 4$) were subjected to repeated measures with ANOVA, and bars not accompanied by the same letter significantly differ at the 5% level, using Tukey's test.

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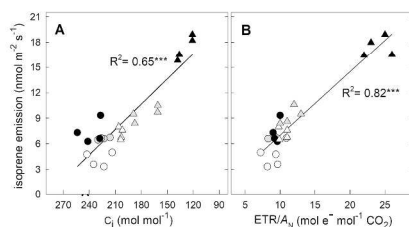


Figure 5. Linear relationships between isoprene emission rate and (A) internal CO₂ concentration (C_i) or (B) the ratio of electron transport rate to photosynthesis (ETR/AN) in *Moringa oleifera* plants. Measurements were made at FTSW60 (10 d, open symbols), FTSW40 (20 d, grey symbols), and FTSW25 (30 d, closed symbols) both in well-watered plants (FTSW100) (triangles) and water-stressed (circles) plants. Coefficient of determination (R²) of each relationship are reported; *** indicate P<0.0001.

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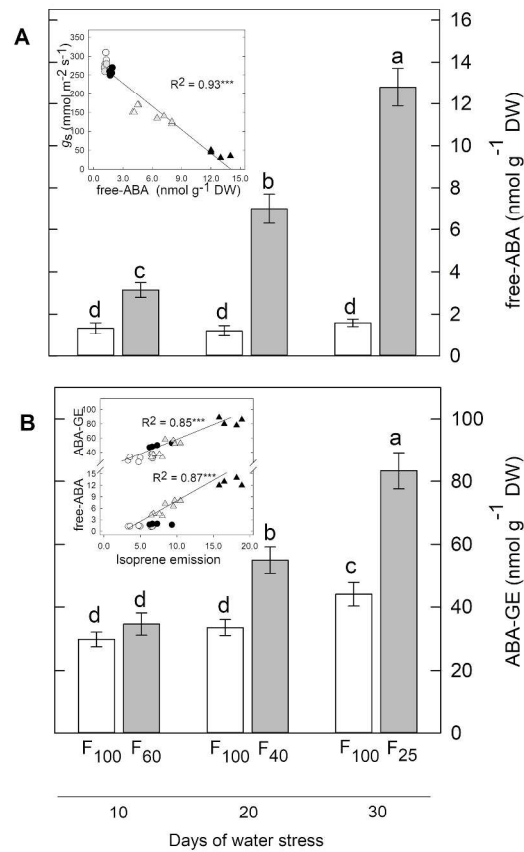


Figure 6. Contents of free-ABA (A) and ABA-GE (B) in FTSW100 (F100) plants (open bars) and in FTSW60 (F60), FTSW40 (F40) and FTSW25 (F25) water-stressed plants (grey bars) of *Moringa oleifera*. Data (means \pm SD, $n = 4$) were subjected to repeated measures with ANOVA, and bars not accompanied by the same letter significantly differ at the 5% level, using Tukey's test. Inset in Figure 6A shows the inverse relationship between foliar free-ABA content and stomatal conductance (g_s). Inset in Figure 6B shows the linear relationships between isoprene emission rates (nmol m⁻² s⁻¹) and free-ABA and its glucoside ester (ABA-GE) contents in FTSW100 plants (circles) and in water-stressed (triangles) plants at FTSW60 (white symbols), FTSW40 (grey symbols), and FTSW25 (dark symbols), respectively. Coefficient of determination (R^2) of each relationship are reported; *** indicate $P < 0.0001$.

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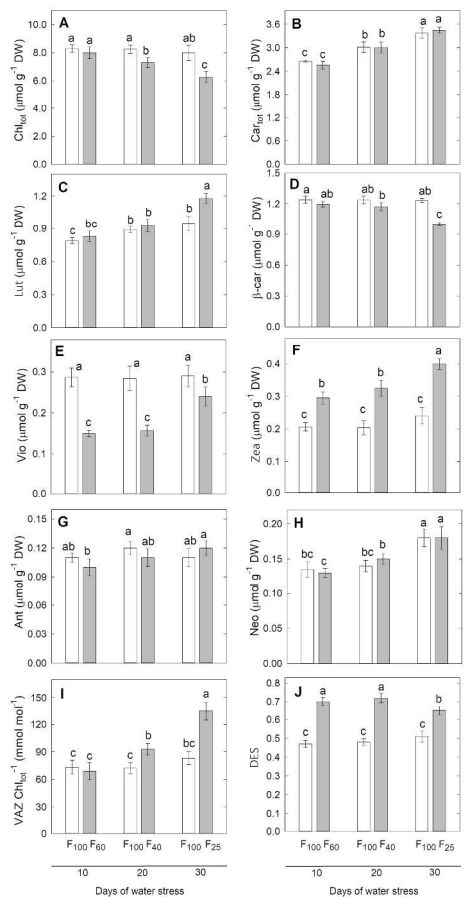


Figure 7. Effects of water stress on the contents of photosynthetic pigments (A-I), on the ratio of violaxanthin cycle pigment content to total chlorophyll content (VAZ Chltot-1, I) and on the de-epoxidation state of VAZ [DES = (0.5A + Z) (V + A + Z)-1, J] in FTSW100 (F100) plants (open bars) and in FTSW60 (F60), FTSW40 (F40) and FTSW25 (F25) water-stressed plants (grey bars) of *Moringa oleifera*. Data (means ± SD, n = 4) were subjected to repeated measures with ANOVA, and bars not accompanied by the same letter significantly differ at the 5% level, using Tukey's test.

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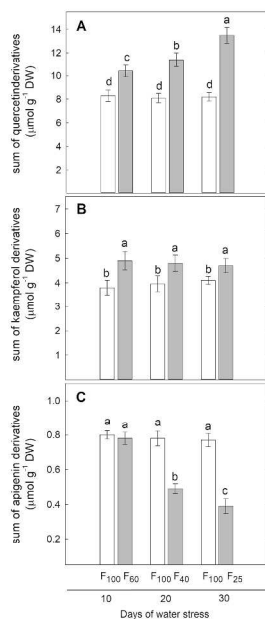


Figure 8. Contents of quercetin (A), kaempferol (B) and apigenin (C) derivatives in FTSW100 (F100) plants (open bars) and in FTSW60 (F60), FTSW40 (F40) and FTSW25 (F25) water-stressed plants (grey bars) of *Moringa oleifera*. Data (means \pm SD, n = 4) were subjected to repeated measures with ANOVA, and bars not accompanied by the same letter significantly differ at the 5% level, using Tukey's test.

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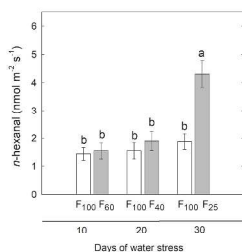


Figure 9. Rates of n-hexanal emission in FTSW100 (F100) plants (open bars) and in FTSW60 (F60), FTSW40 (F40) and FTSW25 (F25) water-stressed plants (grey bars) of *Moringa oleifera*. Data (means \pm SD, n = 4) were subjected to repeated measures with ANOVA, and bars not accompanied by the same letter significantly differ at the 5% level, using Tukey's test.

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