

Water limitation affects weed competitive ability for light. A demonstration using a model-based approach combined with an automated watering platform

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Delphine Moreau, Hugues Busset, Annick Matejicek, Marion Prudent, Nathalie Colbach. Water limitation affects weed competitive ability for light. A demonstration using a model-based approach combined with an automated watering platform. Weed Research, 2022, 62 (6), pp.381-392. 10.1111/wre.12554. hal-03910768

HAL Id: hal-03910768 https://hal.inrae.fr/hal-03910768

Submitted on 14 Jun 2023

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     Running head (40 characters): Response of weed species to water stress
15
16
17
     Keywords:
     Climate change, competition, trait, specific leaf area, height to biomass ratio, root to total
18
     biomass ratio, leaf to aboveground biomass ratio, Abutilon theophrasti, Alopecurus
19
     myosuroides, Amaranthus hybridus
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     Total word count: 8143
22
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25 **1. Summary**

Because of climate change, investigating how morphological traits involved in competition for 26 light (main resource for which crops and weeds compete in conventional cropping systems and 27 temperate and tropical climate) respond to water limitation is crucial to better understand crop-28 29 weed dynamics in the field. Our objective was to develop an innovative approach to quantify the response of weed species to water limitation, and test it with three species. This approach 30 31 combined (1) key morphological traits involved in competition for light (taken from a mechanistic crop-weed model) as criteria to analyze response to water limitation and (2) a 32 33 pot/greenhouse platform allowing automated precision-watering and daily quantification of soil water availability in each pot. Response to water limitation differed among species/stages. For 34 all species/stages, increased plant height per unit of aboveground biomass and production of 35 36 smaller/thicker leaves were the most noteworthy responses. Plants with a strong increase of 37 plant height per unit of aboveground biomass in response to water limitation were able to maintain high specific leaf area, even at low soil water availability. Increase in biomass 38 39 allocation to roots (vs. aboveground parts) and to leaves (vs. stems and reproductive organs) were also observed, but not for all species/stages. Overall, these effects of water limitation on 40 41 morphological traits, suggest strong interactions between competition for light and water.

43 **2. Introduction**

44 Weeds can greatly reduce yield and harvest quality, mainly by competing with crops for 45 resources (Oerke, 2006). That is the reason (together with a high efficiency) why herbicides generally play a key role in ensuring crop production efficiency in conventional cropping 46 systems. However, reducing the excessive use of herbicides has become necessary in view of 47 harmfulness for the environment and public health (Hasanuzzaman et al., 2020). In this context, 48 promoting weed ecological regulation by shifting resource availability and use from weed to 49 crop may provide a more sustainable weed management (Petit et al., 2018). One promising 50 51 option could be to use competitive crop plants, especially for light which is the main resource 52 for which crops and weeds compete in conventional cropping systems and in temperate and 53 tropical climates (Wilson and Tilman, 1993; Perry et al., 2003). However, the frequency of 54 water stress events may increase with climate change, especially in Southern Europe 55 (http://discomap.eea.europa.eu/climate). So, one challenge is to identify crops that are 56 competitive for light, even under water stress conditions.

57 Species competitive ability is directly related to their capacity to adapt to resource availability (Navas and Violle, 2009). Thus, investigating the response of crop and weed species to water 58 59 limitation is crucial. Especially, we need to better understand how plant traits involved in competition for light respond to water stress. Some studies characterized the response of weed 60 species to water stress. However, most of them focused on the germination phase (e.g. Gardarin 61 et al., 2010; Ruhl et al., 2016; Yuan and Wen, 2018) or on global plant growth (e.g. Chahal et 62 al., 2018; de Oliveira et al., 2018). Moreover, they were conducted on a low number of weed 63 species, which contrasts with the high number of weed species that may be found in arable 64 fields (Fried et al., 2008). To our knowledge, no study has yet quantified the diversity of the 65 response of weed species to water availability with a focus on the key morphological traits 66 involved in competition for light. The reasons for this are two-fold. Carrying out experiments 67 with a range of water treatments requires individual daily pot watering which is very time-68 69 consuming when performed manually, especially on a large number of pots. Moreover, until 70 recently, the key morphological traits involved in crop-weed competition for light were not identified. However, two main recent advances should make it possible to lift these limitations. 71 72 On the one hand, the development of innovative high-throughput platforms with automatic and 73 precise individual pot watering in greenhouse allows to overcome these technical limitations 74 (e.g. Granier et al., 2006; Jeudy et al., 2016; Brichet et al., 2017). Up to now, such platforms 75 have almost entirely been used for crop species or for species used as models in genetic/genomic 76 studies. Even when the objective was to analyze crop competitive ability with weeds, only crop

plants were studied (Nguyen et al., 2018; Anandan et al., 2020). On the other hand, a recent 77 78 study using a mechanistic simulation model allowed to identify key morphological traits 79 involved in competition for light between arable crops and annual weeds (Colbach et al., 2019). Considering competition for light only (i.e. competition for water was neglected), Colbach et 80 al. (2019) identified the key traits explaining weed harmfulness, crop ability to limit weed-81 caused yield loss and/or crop potential yield. They were specific leaf area (SLA corresponding 82 to the leaf area per unit of leaf biomass), height to biomass ratio (HBR corresponding to plant 83 height per unit of aboveground biomass) and leaf to aboveground biomass ratio (LBR 84 85 corresponding to leaf biomass per unit of aboveground biomass).

Based on these recent advances, our objective was to quantify the response of weed species to 86 87 water stress in order to determine which morphological traits are influenced by water stress and 88 whether the response to water stress depends on plant species and stage. Our aim was also 89 technical: to test an innovative approach for weed characterization combining (1) morphological traits derived from a simulation study as criteria to analyze weed response to 90 91 water limitation and (2) an experimental platform allowing automatic and precise plant watering. The final aim was to determine whether the results from this approach (applied here 92 93 to a small number of weed species and stages as a proof of concept) could help in the near future 94 for the large-scale characterization of the large panel of weed species potentially present in 95 agricultural fields.

96

97 **3. Materials and methods**

98 **3.1.Experimental treatments**

A greenhouse experiment was conducted in Dijon (France) using three annual weed species: 99 Alopecurus myosuroides Huds. (monocotyledonous C3 species preferentially germinating in 100 autumn), Amaranthus hybridus L. (dicotyledonous C4 species preferentially germinating in 101 102 spring and summer) and Abutilon theophrasti Medik. (dicotyledonous C3 species preferentially germinating in summer). Five water treatments were applied, corresponding to 20%, 30%, 40%, 103 104 55% and 75% of field capacity. The five treatments were applied to the three species, except the 40% treatment on A. hybridus (due to space limitation in the greenhouse). Eight plants were 105 106 grown per species × water treatment combination. Plant species and water treatments were 107 randomly arranged. The duration of the experiment was 62 days.

108

109 **3.2.Growing conditions**

Seeds were germinated in an incubator with a 16-h photoperiod and day/night temperatures 110 adapted to each species (25/20°C for A. myosuroides and 30/25°C for A. hybridus and A. 111 theophrasti). On 14 March 2017, germinated seeds were sown in the greenhouse into 1-L pots 112 filled with a solid and inert substrate made up of 80 % of expanded clay and 20 % of attapulgite 113 (volume proportions), with a bulk density at 0.67 + 0.01 g cm⁻³. The retention curve of the 114 substrate is given in Supplementary Material Section A online. One seed was sown in each pot. 115 116 The pots were placed on fixed tables and watered automatically with a complete nutrient-rich 117 solution made up of N-P-K (10-10-10) and oligo-elements via tubes at a frequency allowing non-limiting watering. Twenty-three days later (on 6 April, when plants were between six and 118 119 nine leaves), plants were transferred into an automated platform where the five water treatments 120 were imposed. This platform (described in Jeudy et al., 2016) is based on conveyors that automatically transport pots towards a watering unit (consisting of a weighing terminal and a 121 122 high-precision pump-watering station) (Supplementary Material Section B online). In our experiment, plants were conveyed three times per day to be weighed. When actual weight was 123 lower than target weight, they were watered (Supplementary Material Section C online). There 124 was no drain because irrigation did not exceed field capacity. Over all irrigation events, 125 irrigation allowed to reach the pot target weight ± 0.13 % (calculated as the absolute difference 126 between the target and the actual weight just after the irrigation event, divided by the target 127 weight). The elapsed time between the irrigation of the first and the last pot (over 130 pots) was 128 34 ± 9 minutes. 129

130 Throughout the experiment, the same complete nutrient-rich solution was used in order to avoid 131 strong plant-nitrogen-limitation even when water supply was low. In the discussion (Section 132 5.4), the comparison of our results to the literature suggests that this methodological approach 133 allowed to minimize a potential nitrogen limitation.

134

135 Target weight was determined as:

136

137
$$\frac{[(sw_{100\%} - sw_{0\%}) \times \% \text{WaterCapacity}]}{100} + sw_{0\%} + \text{EPotW} + \text{PlantW} \qquad [Equation 1]$$

138

139 $SW_{100\%}$ and $SW_{0\%}$ are substrate weight (in g). $SW_{100\%}$ is measured at field capacity, and $SW_{0\%}$ is

140 the weight of dry substrate. *%WaterCapacity* is the target proportion of field capacity (in %,

ranging from 20 to 75 %). *EPotW* is empty pot weight (in g). *PlantW* is fresh plant weight (in

142 g). Plant weight was considered as negligible at the beginning of the experiment. It was adjusted

during the experiment for *A. theophrasti* (using plant weight data from the first sampling date;see Section 3.3).

At each weighing session, the weight of each pot was recorded before and after watering. These data were used to calculate the daily fraction of evapotranspirable soil water, currently named 'Fraction of Transpirable Soil Water' (FTSWd in %), for each species × water treatment combination. The FTSWd relates the actual plant-available soil water content to the total plantavailable soil water content for day d (Lebon et al., 2006). It was calculated as:

150

151
$$FTSWd = \frac{(sw_{100\%} - sw_{0\%}) + \sum_{d=1}^{n} (WaterInput_d) - \sum_{d=1}^{n-1} (Evapotranspiration_d)}{(sw_{100\%} - sw_{0\%})}$$
[Equation 2]

152

WaterInput_d (in g) is the daily amount of water provided by automatic watering at day d, and *Evapotranspiration_d* (in g) is the daily amount of water lost by evapotranspiration at day d.

Note that the day the pots were transferred into the platform, FTSW was close to 100% for all pots, and then differentiated water treatments started (Supplementary Material D). So, during the first days, water input was nil. Water input and evapotranspiration were the same only when the target weight was reached.

Throughout the experiment, artificial light was provided using 400 W lamps (HPS Plantastar, OSRAM, Munich, Germany), in addition to natural light. Mean photoperiod was 14.5 ± 0.7 h. Incident photosynthetically active radiation was 17.2 ± 2.1 mol m⁻² day⁻¹ (silicon sensors; Solems, Palaiseau, France). Air temperature was 21.2 ± 1.7 °C (PT100 sensors; Pyro-Contrôle, Vaulx-en-Velin, France). Air relative humidity was 57.7 ± 3.0 % (AIDC HIH-4000-001; Honeywell, Minneapolis, USA).

165

166 **3.3.Plant measurement and trait calculation**

Plant measurements were made at two sampling dates, and four plants were sampled per species 167 168 \times water treatment \times sampling date combination. At the first sampling date, all the species \times water treatment combinations were sampled. Abutilon theophrasti and A. myosuroides were at 169 170 the vegetative stage, and A. hybridus was at the flowering stage. At the second date, only one 171 species (A. theophrasti) was sampled at the flowering stage. The first sampling was performed 21 days after beginning of water treatments (i.e. 45 days after sowing), leaving enough time for 172 the water treatments to stabilize (Supplementary Material Section D online) and plants to adapt. 173 The second sampling was performed 38 days after beginning of water treatments (i.e. 62 days 174

175 after sowing), when *A. theophrasti* reached the flowering stage.

At each sampling date, the following plant growth traits were measured. Plant height and plant 176 177 leaf area (LI-3100 Area Meter; LI-COR, Lincoln, NE, USA) were measured. Leaf, stem and root biomass were independently determined after 48 h at 80°C. These plant growth traits were 178 179 used to calculate plant morphological traits that play a key role in competition for light (Colbach et al., 2019): (1) leaf to above ground biomass ratio (LBR) is the ratio of the leaf to above ground 180 181 biomass. It reflects the efficiency for producing leaves for a given aboveground plant biomass, with a higher value for leafier plants. (2) Specific leaf area (SLA) is the ratio of the leaf area to 182 the leaf biomass at the plant level. It is the efficiency for producing leaf area from a given leaf 183 184 biomass. High values indicate thin leaves, while low values mean thicker leaves. (3) Height to biomass ratio (HBR) is plant height relatively to aboveground biomass. The higher height to 185 186 biomass ratio, the taller the plants are for a given biomass. We also calculated (4) Root to total 187 plant biomass ratio (RBR), i.e. root biomass relative to total plant biomass, reflecting plant 188 efficiency for producing root biomass from total plant biomass, with a higher value for plants favoring root vs. aboveground growth. This trait is often considered to play a key role in the 189 190 interaction between competition for light and water (McCarthy and Enquist, 2007; Craine and 191 Dybzinski, 2013).

192

193 **3.4. Statistical analyses**

Soil-water availability was expressed in fraction of transpirable soil water, with FTSWmean the mean over FTSWd values from the beginning of water treatment to sampling date. FTSWmean was compared between plant species and plant stages by analysis of variance (lm function of R). As the three weed species were not sampled at both phenological stages, the relative effects of species and stage could not be dissociated. So, the global effect of both species and stage was analyzed.

200 The response of each morphological trait to FTSWmean was analyzed by covariance analysis201 (Im function of R). The following model was used:

202

203
$$plant_trait_{ps} = constant + c_s + a \times FTSWmean_{ps} + b_s \times FTSWmean_{ps} + error_{ps}$$
 [Equation 3]
204

where plant_trait_{ps} was the morphological trait measured on plant p for a given species \times stage combination s, and *constant*, *a*, *c*_s and *b*_s were regression slopes, with the latter two depending on the species \times stage combination s. This model was applied to the four morphological traits. For HBR, a log_n transformation was applied to normalize residue distribution. For SLA, a piecewise regression was performed using the segmented package in R (Muggeo, 2008), in order to account for its two-phase response pattern (Figure 1). So, for HBR, RBR and LBR, the response of species \times stage combinations to water availability was compared on one slope value, while for SLA it was compared on two slope values and breakpoint (i.e. FTSW_{mean} at which slope values change).

214





Figure 1: Theoretical example of the two-phase-response-pattern of specific leaf area (SLA) to water availability (characterized by the fraction of transpirable soil water, FTSW). It shows three parameters: breakpoint (i.e. FTSW at which slope value changes) and left and right slopes (i.e. slope of the regression when FTSW is lower and higher, respectively, than breakpoint value).

220

4. Results

222 **4.1. Dynamics of soil-water availability**

223 The five water treatments resulted in different soil-water status characterized by the mean 224 fraction of transpirable soil water (FTSW) (Figure 2). For most water treatments (defined by the target proportion of field capacity), the mean daily FTSW (averaged over the period running 225 from the beginning of water treatments to sampling date) varied with the plant species/stage 226 (Figure 2). In general, FTSW was lower for A. theophrasti at the reproductive stage. This was 227 due to the dynamics of FTSW that progressively decreased over time until stabilization 228 229 (Supplementary Material Section D online) and to the later sampling date (39 instead of 22 days after the beginning of water treatments) for A. theophrasti at the reproductive stage, resulting 230 in more days at low FTSW values. As a given water treatment mostly resulted in different 231 FTSW values depending on species/stage, plant morphology was analyzed in response to FTSW 232 233 in the following sections.



235

Figure 2: Fraction of transpirable soil water (FTSW) averaged over the period running
from the beginning of water treatments to sampling date, for each species/stage and water
treatment combination. P-values refer to variance analysis performed independently for each
water treatment (defined by the target proportion of water in pot). Within each treatment, bars
with the same letter show values that are not significantly different (least significant difference
test).

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- 243

244 **4.2.Response of plant growth to water availability**

Water availability strongly influenced all the plant growth traits, i.e. biomass, leaf area and height (Table 1A and Supplementary Material Section E online). The response of all these traits to water availability varied with the plant species/stage (Table 1A). For all species × stage combinations, plant leaf area was the most responsive variable (Table 1B).

249

Table 1: Effects of water availability and species/stage on plant growth traits. A. Partial R²
(calculated from the type III sum of square of Anova function of R) indicates the proportion of
variance explained by each factor for each trait. B. Variation factor (calculated as the ratio of
the maximum to minimum value) shows the variability for each trait and each species/stage.
FTSW for fraction of transpirable soil water. *** for P<0.001

Plant growth traits	Total biomass	Aboveground biomass	Leaf biomass	Root biomass	Leaf area	Height
A. Partial R ²						
FTSW	0.393***	0.374***	0.459***	0.448***	0.482***	0.480***
Species/stage	0.374***	0.378***	0.329***	0.351***	0.343***	0.308***
Interaction	0.212***	0.229***	0.182***	0.141***	0.150***	0.111***

Total R ²	0.980	0.982	0.970	0.940	0.974	0.899	
B. Variation factors for each species/stage							
Abutilon theophrasti vegetative	7.2	8.0	7.8	5.8	14.9	3.2	
Abutilon theophrasti reproductive	15.1	18.4	12.3	9.8	23.0	5.9	
<i>Alopecurus myosuroides</i> vegetative	4.9	4.9	4.8	4.9	12.9	2.6	
Amaranthus hybridus reproductive	4.9	5.4	5.7	4.0	11.0	5.6	

258 **4.3.Response of plant morphology to water availability**

259 The four studied morphological traits varied with soil water availability and, for all of them, the

260 response to water availability varied with the plant species/stage (Table 2A). Height to biomass

ratio HBR and specific leaf area SLA were the traits that responded the most to soil-water

- availability for each species /stage (Table 2B).
- 263

Table 2: Effects of water availability and species/stage on morphological traits. A. Partial 264 R² (calculated from the type III sum of square of Anova function of R) indicates the proportion 265 of variance explained by each factor for each trait. B. Variation factor (calculated as the ratio of 266 the maximum to minimum value) shows the variability for each trait and each species/stage. 267 FTSW for fraction of transpirable soil water. As the specific leaf area shows a two-phase 268 response pattern, the analysis was performed separately for FTSW values lower vs. higher than 269 270 breakpoint (i.e. FTSW at which slope value changes). Grey cells indicate when FTSW had nonsignificant effects on the morphological trait. HBR was log_n-transformed for statistical analysis. 271 ns for P>0.05; ** for P<0.01; *** for P<0.001. 272

Morphological traits	Leaf to	Height to	Root to total	Specific leaf area SLA			
	aboveground	biomass ratio	plant	For FTSW	For FTSW		
	biomass	HBR	biomass ratio	>breakpoint	<breakpoint< td=""></breakpoint<>		
	ratio LBR		RBR				
A. Partial R ²							
FTSW	0.05**	0.32***	0.13***	<0.01ns	0.22***		
Species/stage	0.40***	0.48***	0.38***	0.47***	0.28***		
Interaction	0.12***	0.05***	0.10**	0.10ns	0.21**		
Total R ²	0.57	0.85	0.61	0.57	0.72		
B. Variation factors for each species/stage							
Abutilon theophrasti							
vegetative	1.17	3.37	1.37	2.21			
Abutilon theophrasti							
reproductive	1.64	3.75	1.87	1.97			

Alopecurus				
myosuroides vegetative	1.33	3.69	1.71	3.42
Amaranthus hybridus				
reproductive	1.54	2.40	1.72	2.48

277 4.3.1. Height to biomass ratio

For each species \times stage combination, height to biomass ratio HBR increased exponentially with decreasing soil water availability (Figure 3). In response to water limitation, both aboveground biomass and plant height decreased, but the effect was larger for aboveground biomass than for plant height (Table 1; Supplementary Material Section E online). *A. theophrasti* was the most responsive species, followed by *A. myosuroides* and *A. hybridus* (see slope values on Figure 3). For A. *theophrasti* (the only species with data at both vegetative and reproductive stages), the phenological stage did not significantly affect the response slope.



285

Figure 3: Response of height to biomass ratio (logn-transformed) to soil-water availability (fraction of transpirable soil water) for each species × stage combination. The slope of the relationships (+/- standard error) and R² are shown (** for P<0.01 and *** for P<0.001). Slopes with the same letter show non-significant differences. Each symbol represents one plant.

290

291 *4.3.2. Specific leaf area*

292 Contrary to height to biomass ratio, specific leaf area SLA decreased with decreasing soil water

293 availability (Figure 4), meaning that plants produced smaller and thicker leaves in response to







313 Figure 4: Response of specific leaf area to soil-water availability (fraction of transpirable

314 soil water) for each species/stage combination. The slopes and breakpoint of the relationships

315 (+/- standard error) and R² (** for P<0.01 and *** for P<0.001) are shown. Values with the
316 same letter show non-significant differences. Each symbol represents one plant.

317

318 *4.3.3. Root to total plant biomass ratio*

For two out of the four combinations of species × stage (*A. theophrasti* at the vegetative stage and *A. myosuroides*), root to total plant biomass ratio RBR was independent of soil water availability (average value at 0.25 g g⁻¹ on Figure 5). Only for *A. hybridus* and *A. theophrasti* at the reproductive stage, RBR increased linearly with increasing soil water limitation. This indicated that proportionally more biomass was invested to root vs. aboveground plant parts in response to water limitation. The intensity of this RBR increase (reflected by the slopes on Figure 5) was similar for both species.

326



327

Figure 5: Response of root to total plant biomass ratio to soil-water availability (fraction of transpirable soil water) for each species/stage combination. The slopes (+/- standard error) and R² (*** for P<0.001) are shown when significant. Values with the same letter show non-significant differences. Each symbol represents one plant.</p>

332

333 4.3.4. Leaf to above ground biomass ratio

Leaf to aboveground biomass ratio LBR (i.e. ratio of leaf to aboveground biomass) was independent of soil water availability in most situations (Figure 6). Only for *A. theophrasti* at the reproductive stage, LBR increased in response to water limitation, meaning that

- 337 proportionally more biomass was invested to leaf vs. stem and reproductive organs in response
- to water limitation.
- 339



Figure 6: Response of leaf to biomass ratio to soil-water availability (fraction of
transpirable soil water) for each species/stage combination. The slope (+/- standard error)
and R² (*** for P<0.001) are shown when significant. Each symbol represents one plant.

345 *4.3.5. Correlations among morphological parameters*

To identify potential synergies/antagonisms among processes involved in plant morphological response to water limitation, correlations among parameters were analyzed. We focused on the three parameters for which we had values for the four species × stage combinations (SLA breakpoint, SLA left slope and HBR slope). Interestingly, the more height to biomass ratio increased in response to water limitation (low HBR slope), the more they were able to maintain high SLA even at low FTSW (low SLA breakpoint), and the stronger the decrease of SLA to water limitation (high SLA left slope) (**Figure 7**).

353



Figure 7: Correlation between the slope of the response of height to biomass ratio (HBR)
to soil-water availability (FTSW) and (A) the specific leaf area (SLA) breakpoint and (B)
the left slope of the response of SLA to soil-water availability. The parameter values (+/standard error) come from Figure 3 (for height to biomass ratio) and Figure 4 (for specific leaf
area). Each symbol represents a species × stage combination.

360

361 **5. Discussion**

5.1. An efficient methodology to analyse the diversity of weed response to water availability 362 363 The response of weed species to water stress has rarely been studied. Here, we used an innovative approach by (1) focusing on morphological traits previously identified in a 364 simulation study as playing a key role in light competition between arable crops and annual 365 weeds (Colbach et al., 2019), and (2) growing weed species on an experimental platform 366 allowing automatic and precise plant watering as well as quantification of soil water availability 367 368 in each individual pot. Only Jeudy et al. (2016) used such a platform, that they combined with high-throughput phenotyping, in order to compare root distribution among species including 369 370 both crop and weed species, in response to soil-nitrogen availability. The present study is the 371 first one using an automated watering platform to quantify the response of weed species to soil water availability. The possibility of this platform (1) to automatically weight and water each 372 pot individually with a high precision and several times a day, and (2) to quantify daily soil 373 water availability in each individual pot was crucial. In most previous studies, the response of 374 375 weed species to different water regimes was based on manual watering. For example, in some

studies a constant volume of water was supplied daily (Monaco et al., 2005). Moreover, in these 376 377 studies, water availability in each individual pot was not quantified. As a consequence, water treatments were most often characterized qualitatively (for example, referring to 'full' vs. 378 379 'medium' vs. 'low' water in Zhang and Wen (2009) or to the target proportion of water in pot in Chahal et al. (2018)). Yet, our study shows that a given water treatment may result in different 380 381 amounts of soil water availability (described here by the fraction of transpirable soil water, FTSW) depending on plant species/stage or sampling date. Moreover, FTSW values may 382 diverge from target proportion of water in pot. Finally, such platforms make it easier to water 383 384 pots several times a day, allowing a more constant soil water availability than with manual watering (limiting pot watering to a maximum of once per day). Based on this innovative 385 386 approach, our study provided new insights on how competition for water and light may interact 387 in determining crop-weed dynamics in the field.

388

5.2. Water availability strongly affected light competition traits

All the studied morphological traits responded to water availability. For the best-documented morphological traits, results agreed well with the literature. Decrease of specific leaf area in response to water limitation is generally observed in annual plant species (Poorter et al., 2009) including weeds (Monaco et al., 2005). This classical physiological adaptation in plants allows them to decrease transpiring leaf area, thereby reducing plant water requirement under dry conditions (Poorter et al., 2009).

Similarly, biomass allocation to roots vs. aboveground plant parts frequently increases with 396 water limitation, both in crop and in wild/weed species (Monaco et al., 2005; Acciaresi and 397 Guiamet, 2010; Eziz et al., 2017; Chahal et al., 2018). This phenomenon refers to the 'functional 398 399 equilibrium' or 'optimal partitioning' theories, considering that plants preferentially allocate biomass to the compartment that acquires the most limiting resource (Brouwer, 1962). A 400 proportionally larger investment in root biomass in case of water limitation is often considered 401 as a way for plants to increase soil exploration and access to water (McCarthy and Enquist, 402 403 2007).

In our study, the preferential biomass allocation to [leaf] vs. [stem and reproductive organs] in response to water limitation occurred for only *A. theophrasti* at the reproductive stage (not for the other species/stages). Previous studies identified that the leaf to aboveground biomass ratio either increased with or did not respond to water limitation (Lu et al., 2014; Chahal et al., 2018; de Oliveira et al., 2018). Further studies with a larger number of plant species are needed to test the importance of leaf to aboveground biomass ratio in the response of plant morphology towater stress.

411 For height to biomass ratio, the scarcity of references in the literature made the comparison 412 difficult. As this trait is known to play a key role in competition for light, it is often analyzed in response to light availability (Leishman and Westoby, 1994; Pakeman et al., 2015; Colbach et 413 414 al., 2020). We did not find any references quantifying its response to water limitation. The observed increase of height to biomass ratio in response to water limitation can simply be 415 interpreted as the result a stronger impact of water limitation on aboveground biomass than on 416 417 plant height, without necessarily conducting to a physiological/competitive advantage for 418 plants.

419 Our study allowed ranking plant traits according to the magnitude of their response to water 420 limitation. Increased plant height per unit of aboveground biomass and production of 421 smaller/thicker leaves were the most noteworthy responses to water limitation. Not only these phenomena occurred for all the studied plant species/stages, but also the magnitude of the 422 423 response of the corresponding morphological traits (height to biomass ratio and specific leaf 424 area, respectively) to water were the strongest. Conversely, the response of root to total plant 425 biomass ratio and leaf to aboveground biomass ratio to water limitation was significant for, 426 respectively, only two and one out of the four plant species \times stage combinations under study. Moreover, the order of magnitude of the response of these morphological traits was lower than 427 for height to biomass ratio and specific leaf area. When focusing on the most documented traits 428 429 in the literature, our ranking results differed from Monaco et al. (2005) showing that root to 430 total plant biomass ratio was more responsive than specific leaf area. The focus of their study on another weed species (Isatis tinctoria), other water treatments (50 vs. 100 mL of water 431 432 supply per day) and other environmental conditions (e.g. pot size, light intensity, photoperiod...) probably explains these discrepancies, as reported by Poorter et al. (2012a) and 433 Poorter et al. (2012b). 434

435

436 **5.3. Species and stage-specificities**

437 Our study provides information regarding the interspecies variability of the morphological 438 response to water limitation in a few weed species. To avoid confounding effects with plant 439 stage and sampling date (and therefore duration of water limitation), the species effect could be 440 analyzed only by comparing *A. myosuroides* and *A. theophrasti* at the vegetative stage. Only 441 the response of specific leaf area SLA differed between species, with *A. theophrasti* maintaining 442 high SLA values even when soil water availability was low, which was not the case for *A*.443 *myosuroides*.

444 The stage effect was analyzed by focusing on A. theophrasti (the only species studied at two different stages). For this species, the phenological stage affected three out of the four 445 morphological traits. From the vegetative to the reproductive stage, specific leaf area became 446 less sensitive to water limitation, while root to total plant biomass ratio and leaf to aboveground 447 biomass ratio became sensitive to water limitation. Poorter et al. (2012c) hypothesized that this 448 449 delayed change of biomass allocation in response to water limitation could be a mechanism of 450 plant preservation. Indeed, they argued that changing biomass allocation too quickly might 451 result in a suboptimal growth after restoration of the water supply, as water availability often 452 depends on rain events that are unpredictable. Thus, in accordance with this hypothesis, our 453 results suggest that changes in biomass allocation would occur in the long term. Short-term 454 response to water limitation would primarily change plant height per unit of aboveground biomass and specific leaf area. 455

To be noted that correlations among these short-term responses were identified, with plants strongly increasing height per unit of aboveground biomass in response to water limitation being able to maintain high specific leaf area, even at low soil water availability. Studies on additional plant species would be necessary to confirm this interaction between morphological changes.

461

462 **5.4.** Disentangling the effects of nitrogen from those of water limitation

In our experiment, plants were watered with a nutrient-rich solution in order to limit a possible 463 464 indirect effect of a nitrogen limitation that could be associated with a lower water supply. Nonetheless, a nitrogen limitation cannot be excluded in our study (Supplementary Material 465 466 Section F online). A parallel study was conducted on A. myosuroides in order to analyse the morphological response to nitrogen limitation, focusing on the same morphological traits 467 except root to total plant biomass ratio (Supplementary Material Section G online) (Perthame 468 469 et al., 2022). Height to biomass ratio was unresponsive to soil-nitrogen limitation in Perthame et al. (2022), which contrasts with the strong response of this trait to water in the present study. 470 471 Moreover, leaf to aboveground biomass ratio decreased in response to nitrogen limitation in 472 Perthame et al. (2022), while this trait was unresponsive to water limitation in the present study. 473 Specific leaf area was the only trait responding in the same direction (i.e. decrease) to both nitrogen and water limitation. However, the responsiveness was stronger to water than to 474 475 nitrogen. Moreover, the two-phase response pattern observed in the present study for water was

476 not identified when analysing the response to nitrogen. Thus, altogether these results strongly 477 suggest that, even if a nitrogen limitation cannot be totally excluded in our study, the water 478 effect was larger than a possible nitrogen effect. It is likely that the lower nitrogen supply in 479 treatments with a strong water limitation was sufficient to fulfil the lower nitrogen requirements 480 of water-stressed plants, thus limiting the putative confounding effects of water and nitrogen 481 deprivations.

482

483 **5.5. Practical implications**

484 In temperate and non-irrigated field conditions, water availability may vary, depending on 485 rainfall events. Given the strong influence of water on seed germination and emergence (Durr 486 et al., 2015), these events play a key role in determining which weed species emerge and when 487 in a given field during the cropping season. Situations of alternation among conditions of water 488 limitation also result in diverse crop-weed competition situations: either for water mainly, or 489 for light mainly under non-limiting water conditions (provided that nitrogen is not a limiting 490 factor). In this context, the strong effect identified in the present study of water limitation on key morphological traits involved in competition for light, suggests that water 491 492 limitation/competition could affect light competition in the field. Consequences on crop-weed 493 communities could be expected.

Water limitation affects weed dynamics in the field (Souza et al., 2020). The present study 494 provides insight regarding the underlying mechanisms, with plant morphology possibly playing 495 a key role. Moreover, by showing species- and stage-specificities in the response of 496 497 morphological traits to water availability, our study provides explanations on why the 498 consequences of water limitation on crop-weed dynamics may depend on the starting date of 499 the water limitation, its intensity and its length, as well as on the weed species (that may respond 500 differently to water limitation). Our results also suggest that promoting weed ecological 501 regulation by competition requires taking into consideration competition not only for light but also for water (Andrew et al., 2015), especially in areas with dry climates and/or subjected to 502 503 the context of climate change (Iglesias et al., 2012). However, the results from our comparative ecology study are, as such, insufficient to identify management options that would be relevant 504 505 to regulate weeds in situations of water limitation. First, we studied only three weed species. 506 Second, interactions among traits should not be neglected. For example, plant species A with a 507 high ability to increase height to biomass ratio could be assumed to have a competitive 508 advantage over a neighboring plant species B with a lower ability. However, a better tolerance 509 of photosynthesis (per unit leaf area) to water stress of species B could counterbalance the

shading effect of species A. Third, many and diverse components interact in the agroecosystem
(e.g. management techniques, pedoclimate, biophysical processes, diversity of the weed flora,
long time-step resulting from the persistence of the weed seed bank in the soil).

Using a simulation model quantifying the effects of cropping systems on weed and crop 513 dynamics is essential to cope with this complexity. In particular, process-based models are 514 useful to synthesize existing knowledge, identify knowledge gaps, explore prospective 515 scenarios in different contexts in the long term, and design new cropping systems (Colbach, 516 517 2010; Renton and Chauhan, 2017). Among the existing models simulating weed dynamics 518 (Holst et al., 2007), FLORSYS is, to our knowledge, the only individual-based model quantifying 519 the effects of cropping systems on the dynamics of multispecies weed floras, in interaction with 520 pedoclimate (Colbach et al., 2014; Colbach et al., 2021). This model currently only considers 521 crop-weed competition for light (Munier-Jolain et al., 2013) and nitrogen (Moreau et al., 2021), 522 assuming that water resources are sufficient to sustain both crop and weed requirements. So, using the approach developed in the present study, the next step will consist of investigating the 523 524 large diversity of response to water limitation among the numerous weed and crop species in 525 arable fields. Such information will be useful to (1) introduce competition for water into 526 FLORSYS, and (2) perform simulations with the new model version in order to identify innovative weed management strategies that are both sustainable (e.g. low reliance on herbicide 527 528 use) and robust to climatic hazards (e.g. the weed management strategies are efficient even in 529 case of climatic hazards).

530

531 6. Conclusions

Weed response to water availability was analyzed using an original approach (1) focusing on 532 key species traits involved in competition for light that were derived from a mechanistic 533 534 simulation model and (2) using an experimental platform allowing automatic and precise watering. Focusing on three weed species, this study allowed ranking traits according to their 535 responsiveness to water limitation and characterizing species- and stage-specificities in 536 537 responses. The strong effect of water limitation on key morphological traits involved in competition for light suggests that water competition could strongly affect light competition in 538 539 the field, with consequences on crop-weed communities. In the near future, the methodological 540 approach presented here will be applied to the high-throughput characterization of the large 541 diversity of weed species. Knowledge gained from this approach will feed both comparative ecology approaches on weed species and mechanistic simulation models that will be used to 542 543 better understand crop-weeds dynamics in different scenarios of water limitation.

545 7. Acknowledgments

This work was supported by INRAE, the CoSAC project funded by the ANR (ANR-15-CE18-546 547 0007), the Casdar RAID project funded by the French Ministry in charge of Agriculture and Food (Ministère de l'Agriculture et de l'Alimentation, avec la contribution financière du 548 compte d'affectation spéciale 'Développement agricole et rural'), the European Union's 549 Horizon 2020 Research and innovation program under grant agreement N. 7272171 (ReMIX 550 project), the COPRAA project funded by the French Ministries in charge of Ecology and 551 552 Agriculture, and the PPR SPECIFICS project (ANR-20-PCPA-0008) funded by the "Growing and Protecting crops Differently" French Priority Research Program (PPR-CPA), part of the 553 national investment plan operated by the French National Research Agency (ANR). The authors 554 are grateful to Camille Perrot, Ilana Choulet and the staff of the 4PMI platform of INRAE Dijon 555 for technical assistance, Bruno Chauvel for helpful discussions when designing the experiment, 556 and Pauline Souche--Suchovsky for critical reading of the manuscript. 557

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