- 1 Trait components of whole plant water use efficiency are defined by unique,
- 2 environmentally responsive genetic signatures in the model C<sub>4</sub> grass *Setaria*
- 3
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- 15 One sentence summary: Regulation of plant growth can be partitioned into water
- 16 dependent and water independent processes controlled by unique genetic
- 17 components.
- 18

## 19 ABSTRACT

20 Plant growth and water use are interrelated processes influenced by the 21 genetic control of both plant morphological and biochemical characteristics. 22 Improving plant water use efficiency (WUE) to sustain growth in different 23 environments is an important breeding objective that can improve crop yields and 24 enhance agricultural sustainability. However, genetic improvements of WUE using 25 traditional methods have proven difficult due to low throughput and environmental 26 heterogeneity encountered in field settings. To overcome these limitations the study 27 presented here utilizes a high-throughput phenotyping platform to quantify plant 28 size and water use of an interspecific Setaria italica x Setaria viridis recombinant 29 inbred line population at daily intervals in both well-watered and water-limited 30 conditions. Our findings indicate that measurements of plant size and water use in 31 this system are strongly correlated; therefore, a linear modeling approach was used

32 to partition this relationship into predicted values of plant size given water use and 33 deviations from this relationship at the genotype level. The resulting traits 34 describing plant size, water use and WUE were all heritable and responsive to soil 35 water availability, allowing for a genetic dissection of the components of plant WUE 36 under different watering treatments. Linkage mapping identified major loci 37 underlying two different pleiotropic components of WUE. This study indicates that 38 alleles controlling WUE derived from both wild and domesticated accessions of the 39 model C<sub>4</sub> species *Setaria* can be utilized to predictably modulate trait values given a 40 specified precipitation regime.

41

### 42 **INTRODUCTION**

43 Improving crop productivity while simultaneously reducing agricultural 44 water input is essential to ensure the security of our global food supply and protect 45 our diminishing fresh water resources. Agriculture is by far the greatest industrial 46 consumer of fresh water, largely because productivity losses related to drought 47 stress can decrease crop yields by greater than 50% (Boyer, 1982; Hamdy et al., 48 2003). Addressing these challenges will require an integrated approach that 49 combines irrigation practices that minimize water loss and deployment of crop 50 plants with superior water use efficiency (Boutraa, 2010; Davies and Bennett, 2015; 51 Evans and Sadler, 2008; Gregory and George, 2011; Morison et al., 2008; Stanhill, 52 1986).

53 Plant water use efficiency (WUE) can be broadly defined as the ratio of 54 biomass produced to total water lost by the plant (Bacon, 2009; Blum, 2009; 55 Condon, 2004; Evans and Sadler, 2008; Monteith, 1993; Morison et al., 2008; 56 Tardieu, 2013). This complex trait is determined by many factors including 57 photosynthetic carbon assimilated per unit of water transpired (Condon et al., 2002; 58 Farquhar et al., 1989; Morison et al., 2008; Penman and Schofield, 1951; Seibt et al., 59 2008), leaf architecture (Brodribb et al., 2007; Sack and Holbrook, 2006), stomata 60 characteristics (Franks and Farguhar, 2006; Lawson and Blatt, 2014), epidermal 61 wax content (Premachandra et al., 1994), canopy and root architecture (White and

62 Snow, 2012; Martre et al., 2001), stomatal dynamics (Blatt, 2000; Hetherington and 63 Woodward, 2003; Lawson et al., 2010; Flood et al., 2011; Lawson et al., 2012), 64 hydraulic transport (Edwards et al., 2012; Holloway-Phillips and Brodribb, 2011), 65 portion of carbon lost from respiration (Escalona et al., 2012; Tomás et al., 2014) 66 and partitioning of photo-assimilate (Carmo-Silva et al., 2009; Chaves, 1991). Given 67 that plant species (Stewart et al., 1995; Winter et al., 2005; Zegada-Lizarazu and 68 lijima, 2005; Zhou et al., 2012) and ecotypes within species (Kenney et al., 2014; 69 Lopez et al., 2015; Nakhforoosh et al., 2016; Pater et al., 2017; Ryan et al., 2016; Xu 70 et al., 2009) exhibit variation in WUE it is likely that the characteristics which 71 determine this trait are under genetic control and have evolved in response to 72 different environmental conditions such as water availability (Assouline and Or, 73 2013; Brodribb et al., 2009; Huxman et al., 2004). Therefore, WUE is likely 74 influenced by both genetically encoded developmental programs and changes in 75 growth environments throughout the plant lifecycle (Fleury et al., 2010). 76 The technical challenges associated with measuring plant size and 77 transpiration in large structured genetic populations has historically limited 78 experimental efforts aimed at identifying the genetic components associated with 79 WUE. This is particularly difficult in field settings due to year-to-year climate 80 fluctuation and micro-environmental variation observed within agricultural fields. 81 The advent of controlled environment, high-throughput phenotyping instruments 82 (Chen et al., 2014; Fahlgren et al., 2015; Granier et al., 2006; Pereyra-Irujo et al., 83 2012; Reuzeau et al., 2006; Sadok et al., 2007; Tisné et al., 2013; Walter et al., 2007) 84 alleviates many of these challenges through stringent control of climatic variables 85 and automated, high-resolution measurement of plant size and evapotranspiration 86 across large breeding populations.

Evidence from studies conducted on both crop and model plants indicate that
the traits associated with WUE are heritable and largely polygenic, although
identifying the causal locus associated with differential performance has proven
difficult in crop plants due to plant size and genome complexity (Adiredjo et al.,
2014; Aparna et al., 2015; Chen et al., 2012; Coupel-Ledru et al., 2016; Honsdorf et

92 al., 2014; Parent et al., 2015; Schoppach et al., 2016; Xu et al., 2009). Utilization of 93 model plants (C<sub>3</sub> annuals Arabidopsis thaliana and Brachypodium distachyon) that 94 possess tractable genetic and experimental properties has enabled scientists to 95 identify QTL that contribute to WUE (Des Marais et al., 2016; Easlon et al., 2014; 96 Lowry et al., 2013; Mojica et al., 2016; Vasseur et al., 2014), a few of which have 97 been mapped to causal genes (Ruggiero et al., 2017). Species in the genus Setaria 98 also possess many of these desirable qualities and can be used as experimental 99 models to identify genetic components associated with WUE in a C<sub>4</sub> plant that is 100 closely related evolutionarily to C<sub>4</sub> crops like maize, sorghum and bioenergy grasses 101 (Bennetzen et al., 2012; Brutnell et al., 2010; Huang et al., 2016; Li and Brutnell, 102 2011; Zhu et al., 2017). However, in order to study the diversity of resource 103 utilization tactics present in natural and mapping populations of *Setaria* (Saha et al., 104 2016) or other C4 plant species, methods to quantify plant performance and WUE 105 in different environments must be developed. 106 The objectives of this study were to use a controlled environment high-

107 throughput phenotyping system to characterize the genetic architecture of plant 108 size, water use and WUE in an interspecific *Setaria* recombinant inbred population 109 (RIL) under two different watering regimes. Our findings indicate that plant size, 110 water use and WUE are polygenic traits which are influenced by both soil water 111 content and greater than 10 pleiotropic loci whose effect size changes differentially 112 throughout development. In addition, we identify and discuss several aspects of 113 experimental design that should be considered when performing high-throughput 114 phenomics experiments to study plant WUE.

115

## 116 MATERIALS AND METHODS

#### 117 Plant material and growth conditions

The experiment here was first described in (Feldman et al., 2017), which
focused on plant height, and the details are paraphrased here in quotes for clarity.
An interspecific *Setaria* F7 RIL population comprised of 189 genotypes (1138
individuals) was used for genetic mapping. The RIL population was generated
through a cross between the wild-type green foxtail *S. viridis* accession, A10, and the

123 domesticated S. italica foxtail millet accession, B100 (Bennetzen et al., 2012; Devos 124 et al., 1998; Wang et al., 1998). After a six-week stratification in moist long fiber 125 sphagnum moss (Luster Leaf Products Inc., USA) at 4°C, Setaria seeds were planted 126 in 10 cm diameter white pots pre-filled with  $\sim$ 470 cm<sup>3</sup> of Metro-Mix 360 soil 127 (Hummert, USA) and 0.5 g of Osmocote Classic 14-14-14 fertilizer (Everris, USA). 128 After planting, seeds were given seven days to germinate in a Conviron growth 129 chamber with long day photoperiod (16 h day/8 h night; light intensity 130 230  $\mu$ mol/m<sup>2</sup>/s) at 31°C day/21°C night before being loaded onto the Bellwether 131 Phenotyping System using a random block design replicating each genotype and 132 treatment combination in triplicate. For each replicate, individual plants of the same 133 genotype were grown side by side with one individual receiving unlimited water 134 supply while the other individual was subjected to water limitation. The growth 135 chamber location of each of these paired replicates was randomly assigned and did 136 not vary during the course of the experiment. Plants were grown on the system for 137 25 days under long day photoperiod (16 h day/8 h night; light intensity 500 138  $\mu$ mol/m<sup>2</sup>/s) with the same temperature regime used during germination. Relative 139 humidity was maintained between 40 – 80 %. Gravimetric estimation of pot weight 140 was performed 2-3 times per day and water was added to maintain soil volumetric 141 water content at either 33% full-capacity (FC) (water-limited) or 100% FC (well-142 watered) as determined by (Fahlgren et al., 2015). Prescribed soil water content 143 across both treatment blocks was achieved by 15 days after planting (DAP).

144 The volume of water transpired by individual plants at each pot weighing 145 was calculated as the difference between the measured pot weight and the weight of 146 the pre-filled pot at pot capacity (100% FC) or the difference between current pot 147 weight and the previous weight measurement if no water was added. At the 148 conclusion of each weighing, if the pot weight was below the set point, water was 149 added to the pot to return soil water content back to its target weight. This strategy 150 effectively maintains soil moisture content at a consistent level within both 151 treatment blocks. To evenly establish seedlings before the water limitation 152 treatment began, equal volumes of water (100% FC) were added to all pots for two 153 days after transfer to the system. At 10 DAP, a dry down phase was initiated (no

154 watering) to establish the water-limited treatment block (40% FC) while continuing

to maintain a soil water content of 100% FC within the well-watered treatment

- 156 block.
- 157
- 158 Image acquisition and derived measurements

159 RGB images of individual plants were acquired using a top-view and a side-160 view cameras at four different angular rotations (0°, 90° 180°, 270°) every other day 161 at the Bellwether Phenotyping Facility (Fahlgren et al., 2015). Optical zoom was 162 adjusted throughout the experiment to ensure accurate quantification of traits 163 throughout plant development. The unprocessed images and the details of the 164 configuration settings can be found at the following download site: 165 (https://plantcv.danforthcenter.org/pages/data-sets/setaria\_height.html). 166 Plant objects were extracted from images and analyzed using custom PlantCV 167 Python scripts specific to each camera (side-view or top-view), zoom level, and lifter 168 height (https://github.com/maxifeldman/Feldman Elsworth Setaria WUE 2017).

169 Scaling factors relating pixel area and distance to ground truth measurements

170 calculated by (Fahlgren et al., 2015) were used to translate pixels to relative area

- 171 (pixels/cm<sup>2</sup>) and relative distance (pixels/cm).
- 172

173 Biomass estimation

174 At the conclusion of the experiment, 176 individual plants (91 plants from 175 the 100% FC and 85 from the 40% FC) were harvested to measure aboveground 176 biomass. Gravimetric measurement of fresh weight and saturated fresh weight were taken directly upon tissue harvest after which plant tissue was placed into 177 178 polypropylene micro-perforated bags (PJP MarketPlace #361001), dried for three 179 days at 60 °C and subsequently weighed to determine dry weight biomass. 180 Multivariate linear regression was used to evaluate, select and calibrate a predictive 181 model to estimate both fresh and dry weight plant biomass.

182 Regressing plant fresh weight biomass as a function of side-view area,
183 perimeter length, height, object solidity and width indicated that each of these terms
184 is a significant predictor of fresh weight biomass after stepwise model selection

185 using Akaike's Information Criterion (AIC) (Bozdogan, 1987); multiple  $R^2 = 0.89$ ). 186 Unlike fresh weight biomass, side-view area, width, and height were the only 187 significant terms used for prediction of dry weight biomass after using the AIC 188 stepwise model selection correction procedure (multiple  $R^2 = 0.76$ ). Models 189 containing all significant terms and their interaction achieved a greater model fit, 190 but they introduced artifacts at earlier developmental time points due to model 191 over-fitting (Fig. S1). Generally, models constructed to estimate fresh weight 192 biomass in the well-watered treatment block exhibited greater explanatory power 193 than those constructed to predict dry weight biomass or those in water-limited 194 treatment blocks (Fig. 1).

195 A minimal model containing only the most significant term (side-view area) 196 in both fresh and dry weight models produced a goodness of fit similar to more 197 complex models (fresh weight  $R^2 = 0.86$ ; dry weight  $R^2 = 0.74$ ). To avoid 198 propagation of error, values that incorporated plant fresh weight biomass were 199 calculated based on adjusted side-view pixel area and translated to fresh weight 200 biomass after analysis. Cumulative biomass values calculated on a genotype within 201 treatment basis were interpolated using loess smoothing (Chambers and Hastie, 202 1992). Plant size accumulation on a per day basis was calculated as the difference 203 between the loess fit values on a given day and the estimates from the previous day.

204

205 Water loss tabulation

206 The LemnaTec instrument at the Bellwether Phenotyping Facility provided 207 measurements of water use based upon the gravimetric weight of each pot before 208 watering, the volume of water applied, and the resulting weight after watering. On 209 days when the volume of water added to a pot was greater than zero, the daily 210 volume of water added was the sum of water volume added over a single calendar 211 day. On days when water was not added (e.g. during the dry down period), the 212 water volume was calculated as the minimum gravimetric weight of the pot on the 213 day in question subtracted from the minimum weight value from the previous day. 214 The cumulative volume of water used on a specific day was the sum of all water 215 used prior to that day. By fifteen days after planting (DAP), the dry down phase for

216 the water-limited treatment group was complete and pots containing plants lost 217 substantially more water than their empty pot counterparts in the well-watered 218 treatment group (Fig. 1). This observation indicates pot water loss cannot be 219 considered a proximity measure of total plant transpiration before day 15 in this 220 experiment (Fig. 1). Examination of the ratio between fresh weight biomass 221 accumulated relative to the amount of water used and mathematical prediction of 222 the amount of water used per day over this period suggests that the amount of 223 water used between day 15 and 17 can be used as an approximation of cumulative 224 water transpired by the plant throughout this experiment up to this point (Fig. S2). 225 This data and the observation that at day 17 the plants are still relatively small (less 226 than 8% of their maximum size on average) support the rationale of starting the 227 analysis on this day (Fig. 1). Volumes of water use (daily and cumulative) on a 228 genotype within treatment basis were estimated using loess smoothing.

229

## 230 Heritability and trait variance partitioning

We used the same approach as in (Feldman et al., 2017) and the details are
paraphrased here in quotes for clarity. During this experiment, plant area was
measured every other day, so the number of replicates per treatment to calculate
broad sense heritability on any given day was limited. To alleviate this technical
shortcoming, trait values for each individual were interpolated across missing days
using loess smoothing.

Variance components corresponding to broad sense heritability and total
variance explained was estimated using a mixed linear model using the R package
lme4 (Bates et al., 2015). Broad sense heritability was calculated using two
methods. Within an individual experiment, broad sense heritability on a lineestimate basis was calculated using the following formula:

242

Equation 1:

244 245  $H^{2}_{experiment} = \sigma^{2}_{genotype} / (\sigma^{2}_{genotype} + (\sigma^{2}_{genotype X treatment} / n_{treatment}) + (\sigma^{2}_{residual} / n_{replicates}))$ 

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#### 246

210	
247	in which $n_{treatment}$ is the harmonic mean of the number of treatment blocks in which
248	each line was observed and $n_{\mbox{replicates}}$ is the harmonic mean of number of replicates of
249	each genotype in the experiment. Heritability within treatment blocks was
250	calculated by fitting a linear model with genotype as the only explanatory factor
251	within each treatment block.
252	
253	Equation 2:
254	$H^2_{treatment \ block}$ = $\sigma^2_{genotype}$ / $\sigma^2_{total \ variance}$
255	
256	The proportion of variance attributed to genotype divided by total variance
257	within each treatment block is reported as broad sense heritability within treatment
258	(equation). Total variance explained was calculated by fitting a linear model
259	including factors, genotype, treatment, plot and genotype x treatment effects across
260	all phenotypic values in all treatments. The proportion of variance that is
261	incorporated into these factors divided by the total variance in the experiment is
262	reported as total variance explained for each factor.
263	
264	QTL analysis
265	We used the same approach as in (Feldman et al., 2017) and the details are
266	repeated here in quotes for clarity. "QTL mapping was performed at each time point
267	within treatment blocks and on the numerical difference, relative difference and
268	trait ratio calculated between treatment blocks using functions encoded within the
269	R/qtl and funqtl package (Broman et al., 2003; Kwak et al., 2016). The functions
270	were called by a set of custom Python and R scripts
271	(https://github.com/maxjfeldman/foxy_qtl_pipeline). Two complimentary analysis
272	methods were utilized. First, a single QTL model genome scan using Haley-Knott
273	regression was performed to identify QTL exhibiting LOD score peaks greater than a
274	permutation based significance threshold ( $\alpha$ = 0.05, n = 1000). Next, a stepwise
275	forward/backward selection procedure was used to identify an additive, multiple

QTL model based upon maximization of penalized LOD score. Both procedures were
performed at each time point, within treatment blocks and on the numerical
difference relative difference and trait ratio calculated between phenotypic values
measured in treatment blocks at each time point. QTL associated with difference or
ratio composite traits may identify loci associated with genotype by environment
interaction (Des Marais et al., 2013).

282 The function-valued approach described by (Kwak et al., 2016), was used to 283 identify QTL associated with the average (SLOD) and maximum (MLOD) score at 284 each locus throughout the experiment. Each genotypic mean trait within treatments 285 was estimated using a logistic function, and the OTL significance threshold was 286 determined based upon permutation-based likelihood of observing the empirical 287 SLOD or MLOD test statistic. Separate, independent linkage mapping analysis 288 performed at each time point identified a larger number of QTL locations relative to 289 similar function valued analysis based on the SLOD and MLOD statistics calculated 290 at each individual marker throughout the experimental time course. 291 After refinement of OTL position estimates, the significance of fit for the full multiple 292 QTL model was assessed using type III analysis of variance (ANOVA). The 293 contribution of individual loci was assessed using drop-one-term, type III ANOVA. 294 The absolute and relative allelic effect sizes were determined by comparing the fit of 295 the full model to a sub-model with one of the terms removed. All putative protein 296 coding genes (Setaria viridis genome version 1.1) found within a 1.5-logarithm of 297 the odds (LOD) confidence interval were reported for each QTL."

298

## 299 **RESULTS**

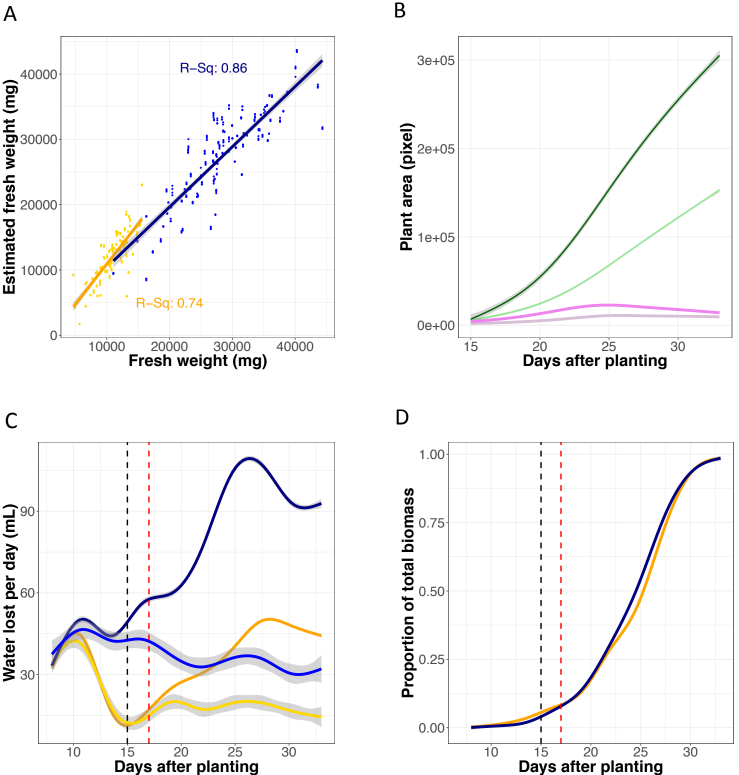
300

301 Measuring plant size and water use throughout the plant lifecycle

Recurrent measurement of plant size and water use was performed on
individuals of a *Setaria* recombinant inbred population (Devos et al., 1998; Devos et
al., 1998) grown at two soil water content levels at the Bellwether Phenotyping
Facility (Fahlgren et al., 2015). Images of each individual plant were captured every
other day from 7 to 33 days after sowing and plant objects were isolated and

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Figure 1. Plant size and water use can be accurately inferred throughout a majority of the plant life cycle. A) Significant correlations between plant fresh weight and pixel area were observed in both the wellwatered and water-limited treatment blocks. B) Plants exhibited a sigmoidal growth curve, characterized by an average maximal rate of growth between 23–26 days after planting. Green lines reflect absolute average size, whereas purple lines report on growth rate. Dark and lighter shaded lines report the wellwatered and water-limited treatment blocks respectively. C) Daily water loss can be accurately measured at 17 days after planting. Dark blue and orange lines correspond to average daily water lost from pots, whereas the lines with lighter shades of similar colors report the average water loss of empty pots. The dashed black line denotes the day at which dry down within the water-limited treatment block is complete whereas the dashed red line demarks when water use can be accurately measured. D) By 17 days after planting, plants have attained less than 8% of their total biomass.



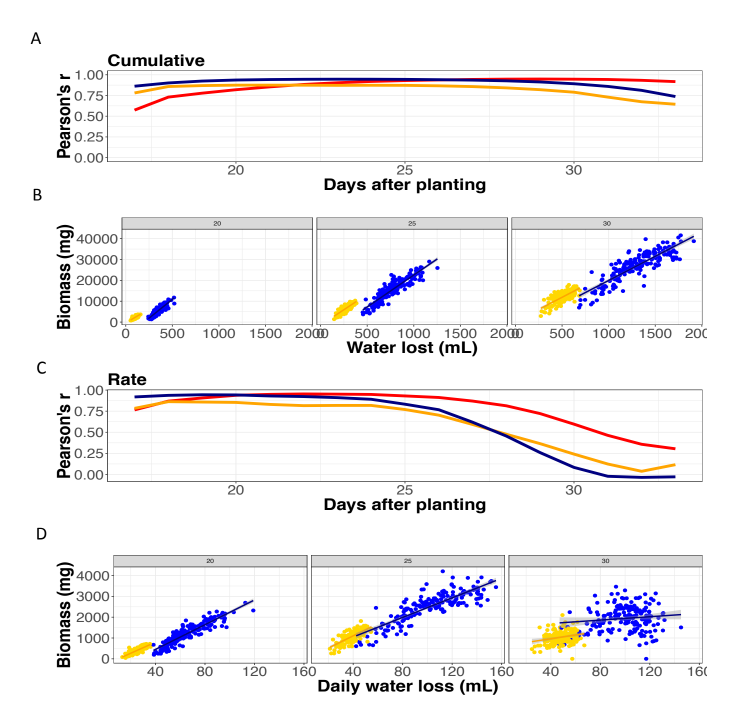
quantified using PlantCV (Fahlgren et al., 2015; Feldman et al., 2017). Weight
estimates of fresh and dry-weight aboveground biomass were calculated using a
simple linear model featuring side-view area as the only predictor (Fig. 1, Fig. S3).

310 Daily plant water use was inferred through gravimetric measurement of pot 311 weight performed two to three times each day by the LemnaTec instrument. The 312 amount of water used by individual plants was calculated as the difference between 313 the measured weight of the pot and the weight of a pre-filled pot at a fixed point that 314 is proportional to its water holding capacity (100% FC) or the difference between 315 current weight and the previous weight measurement if no water was added. At the 316 conclusion of each weighing event, if pot weight was below the set point, water was 317 added to the pot to return it to the target weight value. This strategy effectively 318 maintains soil moisture potential at a consistent level within both treatment blocks. 319 To evenly establish seedlings before the water limitation treatment, equal volumes 320 of water (100% FC) were added to all pots for two days after transfer onto the 321 system. At 10 days after sewing, a dry down phase was initiated (no watering) to 322 establish uniformity within the water-limited treatment block (40% FC) while 323 continuing to maintain a soil water content of 100% FC within the well-watered 324 treatment block.

325 Examination of water loss from empty pots relative to those containing 326 plants suggested that early in the experiment a majority of water loss was 327 exclusively due to evaporation from the soil surface and did not informatively 328 report on plant transpiration (Fig. 1) (Ge et al., 2016). Beginning the analysis at day 329 17 enabled us to minimize the artifacts of evaporation that dominated early in the 330 experiment while still capturing growth attributes over a large proportion ( $\sim 92\%$ ) 331 of the plant growth within the experiment (Fig. 1). Another potential confounding 332 issue was the use of a fixed set point for the pot weight, which neglected the 333 increasing weight of the plant when calculating the amount of water needed to 334 return the pot weight to the set point during watering jobs. This decreased the 335 volume of water present within each pot after watering by approximately 12.5% 336 (well-watered) and 17.5% (water-limited) on average by the end of the experiment 337 (Fig. S4).

## Figure 2. Plant size and water use are tightly correlated.

Pearson Correlation Coefficient both within (blue is well-watered, orange is water-limited) and between (red is across both) treatment blocks indicates strong correlation between these two characteristics, although the correlation between the rate of plant growth and daily water use decreases as plants approach maximum size. A) Correlation between cumulative plant size and water use. B) The relationship between plant size and water use at 20, 25 and 30 days after planting. C) Correlation between the rate of plant growth rate and daily water use at 20, 25 and 30 days after plant growth rate and daily water use at 20, 25 and 30 days after plant growth rate and daily water use at 20, 25 and 30 days after plant growth rate and daily water use at 20, 25 and 30 days after plant growth rate and daily water use at 20, 25 and 30 days after plant growth rate and daily water use at 20, 25 and 30 days after planting.



338 Loess smoothing was used to interpolate the values of traits on a genotype 339 level within each treatment block across all experimental time points (Chambers 340 and Hastie, 1992). Fitting of parametric models was avoided because in many cases 341 the trait values exhibited genotype specific temporal responses that could not be 342 accurately represented by a single model type across the entire population. Rate 343 statistics were calculated from these loess smoothed estimates as the difference of 344 the trait between days. Plots illustrating the mean and variance of each trait can be 345 observed in FIG. S5.

346

347

## Plant size and water use are correlated

348 Over the course of this experiment cumulative plant size and water use were 349 highly correlated. Correlation was tightest between 21 and 27 DAP in the well-350 watered treatment block (> 0.94) and quite strong between 20 and 27 DAP in the 351 water-limited treatment block (> 0.87, Fig. 2). In both treatment blocks, correlations 352 between these characters were weakest at the beginning and end of the experiment 353 but never dropped below 0.67. The correlation of the rate statistics associated with 354 these traits appeared qualitatively different. Correlation between plant growth rate 355 and the rate of water use was initially strong (> 0.79) but rapidly decreased at about 356 26 DAP as the rate of growth slowed (ultimately approaching zero) by the end of the 357 experiment (Fig. 2) while transpiration remained high.

358 We implement two numerical approaches to characterize the genetic 359 architecture of the relationship between these traits. The first method, which is 360 hereafter referred to as the water use efficiency ratio (WUE<sub>ratio</sub>), calculated the ratio 361 of biomass relative to the volume of water lost from the pot. This calculation was 362 performed on a cumulative or daily rate basis.

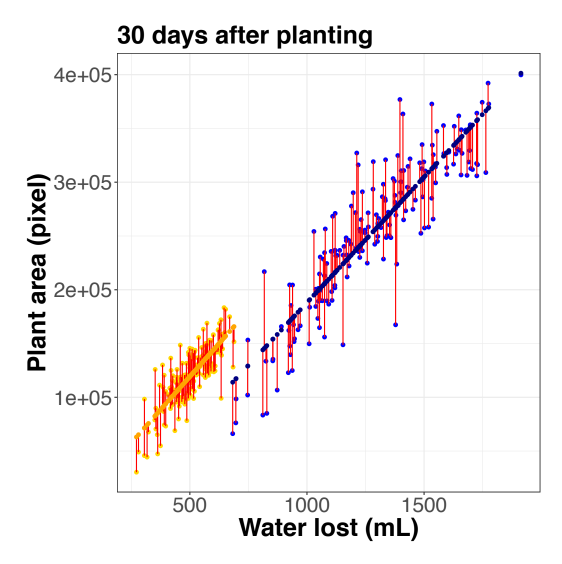
363

364 Equation 3:

365  $WUE_{ratio}$  (pixel/mL) = plant size (pixel) / plant water lost (mL)

366

367 Values of cumulative WUE<sub>ratio</sub> calculated during this experiment were 368 comparable to other experiments where plant size and water use was measured Figure 3. Modeling the relationship between plant size and water use results in two traits. This approach results in predicted value of water use given size ( $WUE_{fit}$ ) colored in dark blue and deviations from this relationship ( $WUE_{residual}$ ) plotted in red. Plot illustrates this relationship at 30 days after planting.



manually at lower throughput (25-29 grams fresh weight / Liter of water, 7-9 grams
dry weight / Liter of water). On average, the cumulative and daily rate WUE<sub>ratio</sub> was
greater in the water-limited treatment block than in well-watered conditions. In
principle, the WUE<sub>ratio</sub> should attenuate the relationship between biomass and water
use, but significant correlation was still observed between these two variables,
particularly within the rate statistic over the last week of the experiment (Fig. S6,
Fig. S7).

The high correlation between plant size and water use suggests that they were not independent traits in this experimental setup. Therefore, as a second approach, ordinary least squares linear regression was used to model the

379 relationship between plant biomass and water use. For each day of the experiment,

 $\label{eq:stability} 380 \qquad \mbox{within treatment blocks a WUE}_{model} \mbox{ was used to predict plant size}$ 

381 (dependent/response variable) based upon water loss (independent/explanatory

- variable) (Fig. 3). The residual of this model fit was evenly distributed around zero
  across the entire distribution of the predicted values suggesting minimal bias of this
  approach (Fig. S8).
- 385

386 Equation 4:

387  $WUE_{fit}$  (pixel/mL) = plant size (pixel) ~ water lost (mL) +  $WUE_{residual}$  (pixel/mL)

388

389 This approach resulted in two traits: The first was the predicted model fit 390 (WUE<sub>fit</sub>) that described the sum of squares relationship between biomass and water 391 use. The residual of this model (WUE<sub>residual</sub>) can be thought of as genotype-specific 392 deviation from this relationship combined with measurement error. As expected, 393 the correlation between the fit values derived from the WUE<sub>model</sub> was highly 394 correlated with plant size (Fig. S9). A slight correlation between cumulative plant 395 biomass and the residual of the WUE<sub>model</sub> was observed particularly later in the 396 experiment demonstrating that biomass had components that were not accounted 397 for by the linear model fit (Fig. S10). Varying the dependence structure/assignment 398 or fitting of the model using major axis regression framework (Legendre, 2014) had 399 little effect on downstream analysis.

400 Each trait (biomass, water loss, WUE<sub>ratio</sub>, WUE<sub>fit</sub> and WUE<sub>residual</sub>) exhibited 401 high average heritability over all experimental time points within and across 402 treatment blocks (0.28 - 0.77) (Fig. S11). Heritability tended to achieve its 403 maximum value in the middle of the experiment with decreased heritability 404 observed at the beginning and the end of the study. Proportionally, the treatment 405 effect of water limitation explained the largest percentage of variance within 406 biomass, water loss and the WUE<sub>fit</sub> although genotype and genotype x treatment 407 interaction also explain a substantial margin of the variance (Fig. S12). Heritability 408 of the rate traits was generally similar but on average 5% lower than the heritability 409 of the cumulative traits. In all cases, the average heritability of each trait was greater 410 within the well-watered treatment block relative to the value calculated in water-411 limited treatment block.

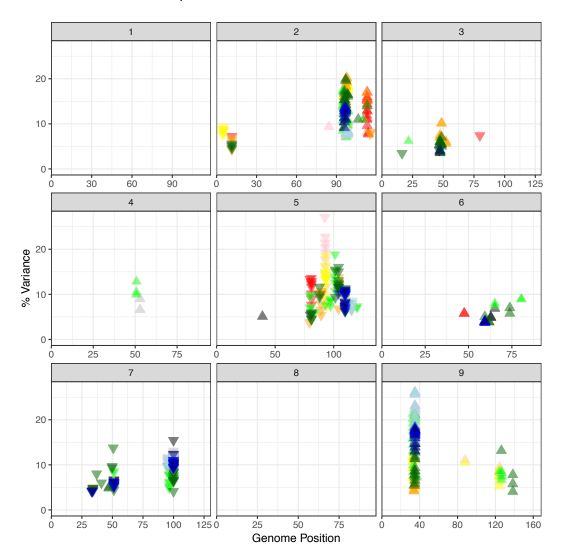
412

413 The genetic architecture of plant size and water use traits

414 For each day of the experiment, a best fit multiple QTL model was selected 415 for each trait (plant size, water use, WUE<sub>ratio</sub>, WUE<sub>fit</sub> and WUE<sub>residual</sub>) and the daily 416 rate of change of the trait within each treatment block based upon penalized LOD 417 score using a standard stepwise forward/backward selection procedure (Broman et 418 al., 2003). This approach identified 86 (cumulative Fig. 4; Table S1) and 106 (rate 419 Fig. S13; Table S1) unique SNPs associated with at least one of the five traits. Many 420 of these uniquely identified SNP positions group into clusters of tightly linked loci 421 that are likely representative of a single QTL location. These local clusters of SNPs 422 (10 cM radius) were then condensed into the most significant marker within each 423 cluster to simplify comparisons of genetic architecture between traits (Fig. S13; Fig. 424 S14). Collapsing these SNP positions yielded 23 unique QTL locations associated 425 with cumulative trait values (Fig. 5) and 27 unique rate QTL locations (Table S2). 426 Of the 23 unique QTL identified, plant biomass contributes the largest 427 proportion of QTL to this set (18) followed by WUE<sub>ratio</sub> (12), WUE<sub>fit</sub> (11), WUE<sub>residual</sub> 428 (10) and water lost (8) (Fig. 5, Fig. S16). Despite the fact that only one QTL location 429 (2@96) was common across all traits and environments, the genetic architecture 430 that contributes to each of these characteristics was clearly related. The strong

#### Figure 4. Eighty-six unique QTL locations were detected across all traits in this experiment.

Each box corresponds to an individual chromosome, where the values along the x-axis are chromosome position and values along the y-axis denote the proportion of genetic variance explained by the QTL. Each triangle represents a single QTL detected, where the color indicates the trait each QTL is associated with (green = plant size, blue = water use, orange =  $WUE_{ratio}$ , black =  $WUE_{fit}$ , red =  $WUE_{residual}$ ). The darkness of color shading is indicative of treatment block where darker represents well-watered and lighter corresponds to the water-limited block respectively. The direction of the arrow indicates the directional effect of the B100 parental allele.

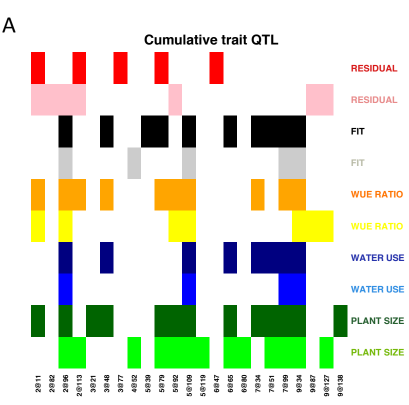


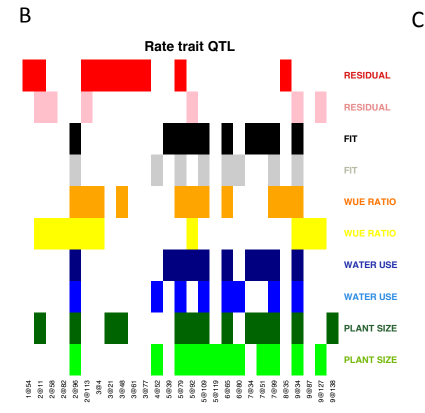
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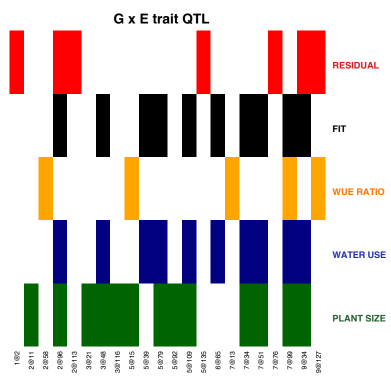
## Figure 5. The genetic components that contribute to subsets of traits largely overlap.

The QTL locations identified are plotted on the x-axis and the traits are plotted on the y-axis. Colored matrix entries denote at least one significant association within this experiment. A) The genetic architecture of cumulative traits. B) The genetic loci associated with trait rate of change. C) Genetic components associated with genotype x

environment traits.







431 correlation of plant size and water loss with the predicted value of plant size given 432 water loss (WUE<sub>ft</sub>) are clearly reflected within the genetic architecture associated 433 with these traits. Plant size, water loss and WUE<sub>fit</sub> all shared 8 QTL (2@96, 3@48, 434 5@109, 6@65, 7@34, 7@51, 7@99 and 9@34) either within the well-watered or 435 water-limited treatment block (Fig. 5, Fig. S16). Plant size, WUE<sub>ratio</sub> and deviations 436 from the relationship between plant size and water use (WUE<sub>residual</sub>) shared five QTL 437 unique to this subset (2@11, 2@113, 5@79, 5@92, and 9@127) which enable 438 divergence from the fundamental relationship between plant size and water loss 439 (Fig. 5, Fig. S16). Several QTL were identified as being uniquely associated with 440 plant size (3@21, 5@119, 6@80, 9@138), WUE<sub>residual</sub> (2@82 3@77, 6@47) and 441 WUE<sub>fit</sub> (5@39) whereas no QTL were identified as being uniquely associated with 442 water loss or WUE<sub>ratio</sub> (Fig. 5, Fig. S16).

443 The genetic architecture of all five traits appears to be influenced by water 444 availability. All traits other than water loss exhibited QTL unique to each treatment 445 block (Fig. 5, Fig. S17). Biomass, water lost, and WUE<sub>fit</sub> all shared four QTL in 446 common across environments (2@96, 5@109, 7@99 and 9@34) where as WUE<sub>ratio</sub> 447 and the WUE<sub>residual</sub> shared a single QTL (2@11) between blocks not found associated 448 with the other traits. Two QTL (3@48, 7@34) were found specifically within the 449 well-watered treatment block for all traits other than WUE<sub>residual</sub> whereas QTL 450 specific to water-limited environment identified common QTL associated with 451 biomass and WUE<sub>fit</sub> (4@52) or WUE<sub>ratio</sub> and WUE<sub>residual</sub> (9@87, 9@127).

452 The identity of QTL associated with the daily rate values suggest that the 453 genetic architectures were largely cognate with the QTL associated with the traits 454 themselves, both in identity and response to treatment. In total, 28 QTL comprised 455 the union of all unique QTL associated with both the trait value and the daily rate of 456 change calculated from the trait value. Of these QTL, 22 were common between both 457 the trait value and rate statistic associated with the trait, whereas five are only 458 found associated with the rate (1@54, 2@58, 3@4, 3@61, 8@35) and only one QTL 459 was uniquely associated with the cumulative trait values alone (6@47) (Fig. S18).

460

461 *Genotype x environment interactions* 

462 To assess the genetic architecture of genotype x environment interactions, 463 mapping was performed on numerical difference, relative difference and trait ratio 464 between the phenotypic values observed within each treatment block. In total, 148 465 unique SNP locations were identified as being significantly associated with at least 466 one of the difference trait formulations across all standard and derived plant size 467 and water use traits (Table S3). Substantial overlap between these categories of 468 genotype x interaction traits indicates that each formulation detects similar genetic 469 signals (Fig. S19) although the large number SNPs found uniquely associated with 470 the trait ratio may indicate that some of these associations may be spurious. As such, 471 these OTL (trait ratio genotype x environment OTL) were removed from further 472 analysis. The numerical difference and relative difference traits exhibited 473 association with 43 and 40 unique SNP positions, which were representative of 20 474 and 18 QTL respectively (Table S4, Fig. S20-22).

475

476 A majority of the QTL (10/15) identified as being associated with the trait 477 difference between treatment blocks were also found associated with the 478 cumulative trait in both treatment blocks (Fig. 5). The exceptions to this were QTL 479 located on 3@21, 3@48, 5@39, 7@34 and 9@127 that were identified as being 480 significantly associated with the difference between treatment blocks but only 481 identified in either well-watered (3@21, 3@48, 5@39, 7@34) or water-limited 482 conditions (9@127). Interestingly, the QTL located on 3@48, 7@34 and 9@127 483 were associated with more than one trait in a single treatment block which may 484 indicate that these QTL impart pleiotropic phenotypic effects that were dependent 485 upon soil water content (Fig. 5).

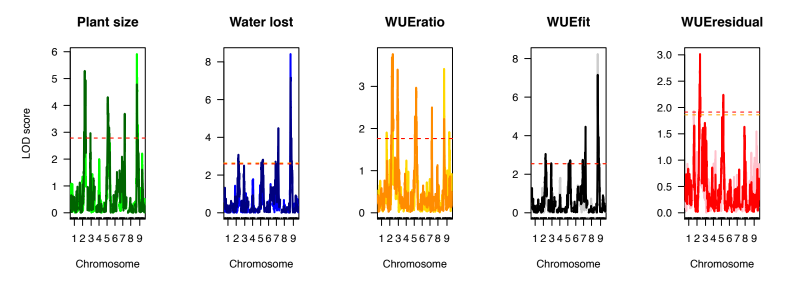
486

487 The temporal genetic architecture of plant growth and water usage

In order to account for the time dependence of the traits for the five plant
traits, we used a function-valued approach based upon average log-odds score
throughout across the experiment (SLOD) for each trait (Kwak et al., 2016). This
analysis parallels the individual time point analysis, although the reduction of

Figure 6. Significant associations identified using single marker scan functional QTL mapping.

Chromosomal position is plotted on the x-axis whereas LOD score of trait association across the genome is plotted on the y-axis. Treatment block is indicated by color intensity (darker is well-watered and lighter is water-limited). Significance thresholds (based on 1000 permutations) are plotted as dashed yellow (water-limited) and red (well-watered) lines respectively.



492 complexity (fewer, higher confidence QTL) provides an opportunity for 493 simplification and better understanding of the major loci that influence plant WUE. 494 SLOD based function-valued OTL models indicate that several major OTL 495 (2@96, 5@109, 7@99, and 9@36) influenced both plant size and water use related 496 traits, although the magnitude of statistical significance attributed to each loci 497 varied by trait and throughout plant development (Fig. 6, Fig S23). Using the SLOD 498 approach, we were able to partition combinations of QTL unique to related traits 499 (Fig. 6). For several QTL (those around 2@96 and 5@109) the positional location at 500 which maximal LOD score was observed changed noticeably in a trait and 501 environment dependent manner either due to multiple closely linked loci or noise in 502 our measurements. Because the confidence intervals of the QTL generally overlap, 503 our reporting in this section will hereafter refer to these loci by their approximate 504 chromosomal location.

505 Both plant biomass and cumulative water use exhibited almost a complete 506 overlap of QTL within the well-watered treatment block, whereas plant size given 507 water use (WUE<sub>fit</sub>) and deviation of plant size from this fundamental relationship 508 (WUE<sub>residual</sub>) each exhibit a unique genetic signature (Fig 6). As observed when trait 509 values at individual time points were treated as independent traits, a single QTL on 510 2@96 is the only genetic component that was shared across all five traits. The linear 511 modeling approach successfully partitions out QTL associated with WUE<sub>fit</sub> (2@96, 512 7@99, 9@36) from the genetic components that contribute to deviations from the 513 plant size ~ water use relationship (WUE<sub>residual</sub>; 2@96, 5@109). The QTL associated 514 with the WUE<sub>ratio</sub> (2@96, 3@52, 5@109) also likely reflects deviations from the 515 relationship between biomass given water loss associated with the WUE<sub>residual</sub>. 516 Overall, the identity of QTL associated with each trait was largely identical between 517 the two treatment blocks (Fig. 6, Fig. S23) as were the QTL associated with the 518 values of rate statistics derived from these measurements (Fig. S24, stepwise 519 method; Fig. S25, scanone method). 520

521 A temporal model of the genetic architecture that influences plant water use efficiency

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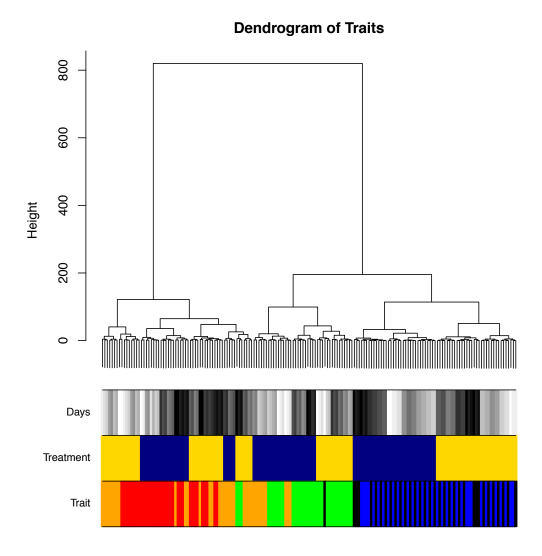
522 Our QTL results suggest at least two components of water use efficiency with 523 distinct genetic architectures. In order to compare the genetic architecture across all 524 traits, treatments and time points in a common framework, we analyzed how each 525 trait was influenced by a common set of loci. Fourteen QTL were selected based 526 upon their association with multiple traits, robust linkage with a single trait and/or 527 having differential contribution to traits across treatment blocks (Table S5) and the 528 proportional contribution of each locus to the additive genetic variance was 529 calculated using drop-one-term, type III, ANOVA performed for all experimental 530 traits, time points and treatment. Agglomerative hierarchical clustering of the 531 signed proportion of additive genetic variance explained by each locus was 532 performed to identify modules of traits and loci that define plant phenotypes. 533 Examination of scree plots of the within group sum of squares suggested that the 534 variance within traits could be attributed to approximately six groupings although a 535 majority of this variance could be captured within the largest 2-3 partitions (Fig. 536 S26). These partitions represented the major relationships between trait classes. 537 The WUE<sub>ratio</sub> and WUE<sub>residual</sub> were generally grouped separately from a larger cluster 538 of traits that included cumulative plant size, water use and WUE<sub>fit</sub> (Fig. 7). The 539 genetic architecture of plant water use and WUE<sub>fit</sub> were more related to each other 540 than they were to plant size, which formed the third group. The influence of water 541 availability on these traits was apparent from the grouping of clusters whereas the 542 effects of time were clear but distributed within the treatment blocks. The genetic 543 architecture of the WUE<sub>ratio</sub> in the well-watered treatment block at early time points 544 was more similar to the architecture of plant area than itself later in development 545 whereas plant area in the water-limited treatment block exhibited a genetic 546 architecture similar to the WUE<sub>ratio</sub> late at the end of the experiment. Examination of the signed, proportional allelic effects within the greater fixed 547 548 QTL model indicated that QTL on 2@96, 5@109, 7@99 and 9@34 contribute

549 medium-to-large effects on a majority of the traits examined in both treatment

- blocks (Fig. 8). The B100 allele associated with QTL on 2@96 and 9@34 both
- contributed to increased plant size, water loss, WUE<sub>fit</sub> and WUE<sub>ratio</sub>. The QTL on
- 552 2@96 exhibited its greatest influence in the well-watered treatment block whereas

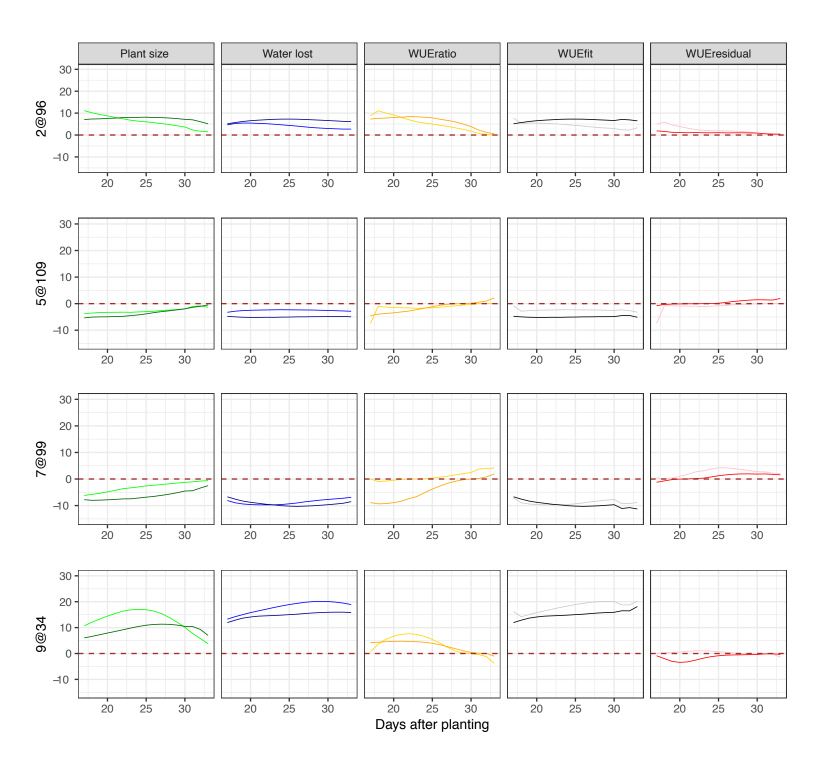
## Figure 7. Agglomerative hierarchical clustering defines the relationship between plant size, water use and derived water use efficiency traits.

The additive effect size of fourteen common QTLs was calculated across all traits, treatments and developmental time points through hierarchical clustering using Ward's method. Color bars on the bottom indicate trait (green = plant size, blue = water use, orange =  $WUE_{ratio}$ , black =  $WUE_{fit}$ , red =  $WUE_{residual}$ ), treatment block (blue = well-watered, orange = water limited), and days after planting (grey scale values where white represents the trait on day 17 and black indicates the trait on day 33).



# Figure 8. Additive relative effect size of the four major pleiotropic QTL plotted throughout the course of the experiment.

A model containing fourteen QTL was fit across traits, treatment blocks and days. The developmental time point (days after planting) is indicated by the x-axis whereas the proportional additive genetic effect size of the B100 allele is plotted along the y-axis. Columns are representative of traits (green = plant size, blue = water use, orange =  $WUE_{ratio}$ , black =  $WUE_{fit}$ , red =  $WUE_{residual}$ ) while rows correspond to individual QTL. Shading within the colors denotes treatment block (darker = well-watered, lighter = water-limited).



553 the contribution of 9@34 was greater on average in the water-limited treatment 554 block. Both OTL exhibited similar temporal patterns, showing an earlier effect on 555 plant size and  $WUE_{ratio}$  but a consistent effect across water loss. Contribution of the 556 B100 allele on 7@99 and 5@109 decrease plant size, water use and the WUE<sub>fit</sub> 557 traits; the effect of which was greater in well-watered conditions. The magnitude of 558 effects contributed by QTL on 7@99 on plant size decreased through time whereas 559 the effects on water loss and  $WUE_{fit}$  peaked after 20 days and decreases slightly 560 thereafter. The 5@109 locus behaves similarly with little temporal variation in 561 plant water use and WUE<sub>fit</sub>. A majority of the other QTL contributed minor effects 562 that became more prominent in one of the two treatment blocks or at a particular 563 developmental time points. Inheriting the B100 allele at QTL on 2@113, 3@48, 564 4@52, 6@65 and 9@127 increased the values while the B100 allele at the remaining 565 loci (2@11, 5@79, 5@95, 7@34 and 7@53) decreased the value of the traits (Fig. 566 S27).

567 A majority of the QTL exhibit unidirectional effects across both the well-568 watered and water-limited treatment blocks although the direction of the effect was 569 largely dependent on the trait (Fig. S28). The exceptions to this trend represent 570 short periods of experimental time at which the relative effect size is near zero 571 within one or both treatment blocks (Fig. 8, Fig. S27).

572 The proportional contribution of parental alleles towards increased trait 573 values varied between traits, within treatment blocks and throughout plant 574 development. For example, B100 alleles contributed to increased trait values for all 575 traits other than WUE<sub>ratio</sub> in the water-limited environment and the WUE<sub>residual</sub> 576 across both treatment blocks (Fig. 9). Alternatively, the contributions of the A10 577 alleles proportionally increased the WUE<sub>residual</sub> value early and then again late in 578 plant development relative to those inherited from the B100 parent. The influence 579 of A10 alleles on the WUE<sub>ratio</sub> was also greater their B100 counterpart under water-580 limited conditions early in plant development.

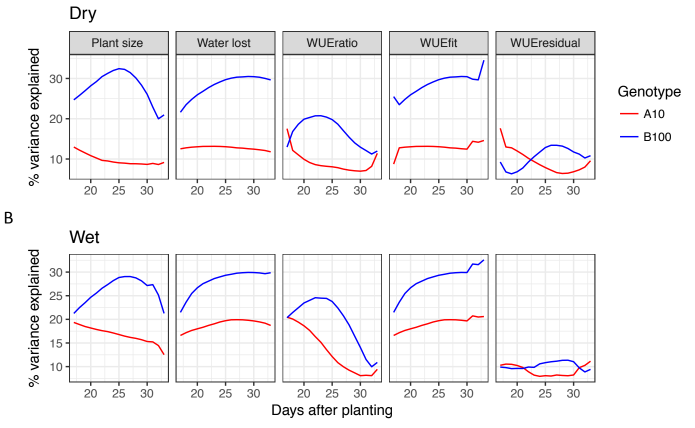
581

582 **DISCUSSION** 

## Figure 9. The proportional contribution of parental alleles to increased trait values depend upon trait, environmental water content and plant developmental stage.

Alleles derived from the B100 parent contribute a greater proportional of additive genetic variance to plant size, water use and TE model fit in both well-watered and water-limited conditions than their A10 allelic counterparts. Both the WUE ratio and TE model residual traits exhibit dynamic behavior where A10 alleles contribute either greater or close to equal proportions of additive genetic variance early and late in plant development. A) The contribution of parental alleles in the water-limited treatment block. B) The contribution of parental alleles in the water-limited treatment block.





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583 The objectives of this study were to utilize technological advances in high-584 throughput phenotyping (Chen et al., 2014; Fahlgren et al., 2015; Granier et al., 585 2006; Perevra-Irujo et al., 2012; Reuzeau et al., 2006; Sadok et al., 2007; Tisné et al., 586 2013; Walter et al., 2007) to characterize the genetic architecture of water use 587 efficiency and how this architecture responds to water-limitation in an experimental 588 C<sub>4</sub> grass model system. Although considerable efforts have been made to 589 characterize these processes in *Arabidopsis thaliana*, C<sub>3</sub> grass crops and other 590 species (Ruggiero et al., 2017) this represents the first study performed on an 591 annual C<sub>4</sub> grass RIL population. These efforts enabled us to identify genetic loci that 592 contribute to differential biomass accumulation given water use in a well-watered 593 and water-limited environment. Our findings suggest that the major genetic 594 components associated with plant size, water use and water use efficiency exhibit 595 pleiotropic behavior and that the magnitude of their allelic effects is dependent 596 upon environment and developmental stage. We used two complementary 597 approaches to define traits, and our analysis confirmed that the genetic architecture 598 was similar with both approaches. We show that the loci controlling biomass 599 accumulation can be roughly divided into two groups: those that control the amount 600 of water used to create biomass ( $WUE_{fit}$ ) and those that control how efficiently that 601 water is used (WUE<sub>residual</sub>). The results from this study indicate that alleles from 602 both domesticated foxtail millet and a species representative of its wild progenitor 603 contribute to maximal vegetative biomass yield or water use efficiency grown in 604 environments with different watering regimes. In addition, we highlight aspects of 605 our experimental design and analysis that could be improved in future studies. 606

607 The genetic architecture of plant size, water use, water use efficiency and the

608 relationship between these traits

Within the A10 x B100 *Setaria* RIL population, plant size, water use and the relationship between these two variables are unique polygenic traits whose values are all likely influenced by greater than 10 loci. Four QTL located on 2@96, 5@106, 7@99 and 9@36 exhibit strong pleiotropic influence across this suite of traits, the relative magnitude of each is dependent upon growth environment and 614 developmental time point. Despite strong correlation between plant size and water 615 use we successfully identified genetic architectures distinct to each trait. This was 616 achieved by modeling plant size as a function of water use and examining the 617 resulting values of the model fit (plant size given water use) and deviations from 618 this relationship (residual of plant size given water use). This linear modeling 619 approach has been used much less frequently in the literature (Lopez et al., 2015; 620 Nakhforoosh et al., 2016) than the more commonly used WUE<sub>ratio</sub> (Adiredjo et al., 621 2014; Honsdorf et al., 2014; Aparna et al., 2015; Fahlgren et al., 2015; Lopez et al., 622 2015). While the genetic architectures associated with the WUE<sub>ratio</sub> and WUE<sub>residual</sub> 623 in this population are closely related (Fig. 7), WUE<sub>residual</sub> exhibits substantial 624 heritability and is less correlated with plant size than the WUE<sub>ratio</sub> (Fig. S6 Fig. S10), 625 making it a more desirable metric.

626 By examining the model based components of WUE with function valued 627 single marker scan QTL analysis that accounts for multiple hypothesis testing across 628 time points (Kwak et al., 2016), we were able to partition the four major pleiotropic 629 OTL into the genetic components on 2@96, 7@99 and 9@36, which control plant 630 size given water use (WUE<sub>fit</sub>) and those on 2@96 and 5@109 that contribute to 631 deviations from this relationship (WUE<sub>residual</sub>). This result suggests that QTL 632 associated with WUE<sub>fit</sub> (7@99 and 9@36) potentially control the development of 633 transpiring plant biomass whereas the QTL associated with the WUE<sub>residual</sub> and 634  $WUE_{ratio}$  (2@96 and 5@109) influence production of non-transpiring tissues or 635 biological processes not directly related to transpiration. This conclusion is in 636 accordance with the results of other studies performed on this population which 637 demonstrate that these loci are largely pleiotropic (Mauro-Herrera and Doust, 638 2016), although the loci on 2@96 and 5@100 substantially influence plant height 639 (Feldman et al., 2017) and stem biomass, whereas those on 7 and 9 are not 640 associated with the accumulation of stem material (Banan et al., 2017). 641 Our study also identified many smaller effect QTL which influence biomass, 642 water use and WUE traits. The B100 parental allele contributes substantial positive 643 (3@48, 4@52, 6@65, 9@127) and negative (7@34, 7@53) effects on all traits, whereas QTL on 2@11, 2@113, 5@79 and 5@95 contribute either to plant 644

645 size/WUE<sub>ratio</sub>/WUE<sub>residual</sub> ratio to a greater degree than on plant size/water

646 loss/WUE<sub>fit</sub>.

647 Roughly two thirds of the OTL associated with trait plasticity as a response to 648 water availability (difference or relative difference between treatment blocks) were 649 also identified as being associated with the cumulative traits within both treatment 650 blocks. This observation indicates that in many cases, soil water content influences 651 the temporal dynamics of the allelic effects by differential progression through 652 developmental processes that share similar genetic components (Feldman et al., 653 2017). This study identifies several QTL (3@48, 7@34 and 9@127) associated with 654 genotype by environment traits which also exhibit significant influence on multiple 655 plant traits within a single treatment block. This provides relatively strong evidence 656 that these QTL have pleiotropic influence on size and water use related traits in an 657 environment specific manner. In contrast, QTL identified only by mapping on the 658 difference or relative difference of the traits between each environment are largely 659 specific to individual traits.

660 Evidence from this study supports an evolutionary genetic model where the 661 majority of QTL associated with the measured traits exhibit conditional neutrality 662 across both soil water potentials examined. Although all traits other than plant size 663 sometimes exhibit opposite directional effects across treatment blocks, the evidence 664 supporting a model of antagonistic pleiotropy is weak. When identified, QTL 665 exhibiting opposite directional effects within individual treatment blocks were 666 limited to short periods of experimental time and are characterized by negligible 667 relative effects during these time points. The contributions of alleles from both 668 parental lines contribute to increased WUE irrespective of soil water potential, 669 suggesting that neither parent was optimized for WUE. For example, alleles from the 670 A10 parent contribute a greater proportion of additive genetic variance to increased 671 WUE during early development in both well-watered and water-limited 672 environments, (particularly given the WUE<sub>residual</sub> derivation of WUE) whereas the 673 alleles derived from the B100 parent have greater affect on a majority of the 674 measured traits throughout the time course. The contribution of alleles of both 675 parents to water use efficiency is expected given earlier study performed on the

676 same platform where parental lines showed similar WUE under water-limited677 conditions (Fahlgren et al., 2015).

678

679 Considerations when measuring plant size, water use and WUE

680 As observed in many other studies (Chen et al., 2012; Fahlgren et al., 2015; 681 Ge et al., 2016; Golzarian et al., 2011; Honsdorf et al., 2014; Lopez et al., 2015; 682 Parent et al., 2015), relative plant side-view pixel area provided a robust and 683 accurate proximity measurement of plant biomass. Although incorporation of 684 additional plant architectural features can improve estimates of this relationship 685 (Parent et al., 2015), our results indicate that caution should be taken as to not over 686 fit models on ground truth data collected exclusively at the end of the experiment as 687 was performed in this study (Fig. S1).

688 Automated or manual gravimetric measurement of pot weight has proven to 689 be a reliable estimator of plant transpiration but only if the evaporative loss of 690 moisture from soil can be accounted for. Results presented in this study indicate 691 that inclusion of empty pots (or pots that contain plastic plants (Parent et al., 2015) 692 or fabric wicks (Halperin et al., 2017)) is an appropriate empirical method to 693 estimate the experimental time point at which transpiration contributes 694 meaningfully to total pot evapotranspiration (Coupel-Ledru et al., 2016; Lopez et al., 695 2015; Pereyra-Irujo et al., 2012). Estimation of evapotranspiration after this critical 696 time point has been effectively used by several other groups to identify and 697 eliminate confounding data points collected early during similar experiments 698 (Vasseur et al., 2014; Coupel-Ledru et al., 2016; Ge et al., 2016). Our findings 699 indicate that subtraction of empty pot weight (as performed by (Pereyra-Irujo et al., 700 2012; Parent et al., 2015; Coupel-Ledru et al., 2016)) may overcorrect for 701 evaporation at early experimental time points even after the point at which plant 702 transpiration contributes substantially to total pot water loss. Although not applied 703 during this experiment, utilization of plastic covering to shield pots from 704 evaporative moisture loss in combination with the approaches discussed above may 705 improve the ability to unambiguously quantify plant transpiration (Aparna et al., 706 2015; Coupel-Ledru et al., 2016; Ellsworth et al., 2017; Granier et al., 2006; Halperin 707 et al., 2017; Vasseur et al., 2014). In this study, the contribution of plant biomass to 708 overall pot weight was not accounted for during the estimation of plant water use. 709 Although the contribution of plant biomass to pot weight in most experiments 710 performed using Arabidopsis thaliana is negligible (Tisné et al., 2010), plant biomass 711 within this Setaria RIL population accounted for 12-18% of total average pot water 712 content by the end of the experiment (Fig. S4). Our inability to account for this 713 growth has the undesirable effect of systematically decreasing the soil water 714 content of larger genotypes, although in practice this small change in soil water 715 potential likely has minimal impact on transpiration dynamics of the plants. 716 Strong correlation between plant size and water use was observed in spite of 717 the fact that these traits can potentially be controlled by different physiological 718 mechanisms. A similar trend has also been described in experiments designed to 719 study water use efficiency in Arabidopsis thaliana, apple and wheat (Lopez et al.,

2015; Nakhforoosh et al., 2016; Schoppach et al., 2016; Parent et al., 2015; Vasseur
et al., 2014). The magnitude of this correlation is likely inflated in this study due to
the large differences in size between parental lines and segregants within the A10 x
B100 RIL population. Future studies aimed at investigating the genetic basis of
water use efficiency can attenuate this correlation by selecting parental lines of
similar size and flowering times that differ in their rates of transpiration within
environments of interest.

727

#### 728 **CONCLUSIONS**

729 This study leverages recent advances in high-throughput phenotyping and 730 quantitative genetics to identify the genetic loci associated with plant size, water use 731 and water use efficiency in an interspecific RIL population of the model C<sub>4</sub> grass 732 Setaria. Our findings indicate that these traits are highly heritable and largely 733 polygenic, although the effects of four major pleiotropic QTL account for a 734 substantial proportion of the variance observed within each trait. Contribution of 735 parental alleles from both the domesticated and wild progenitor lines contribute to 736 maximization of these characteristics. Overall, the underlying genetic architecture of 737 each of these processes is distinct and substantially influenced by soil water content

- as well as plant developmental stage. In addition, several aspects of our
- experimental design which could be improved to obtain a better understanding of
- 740 the genetic components that underlie plant size, water use and water use efficiency
- 741 in future high-throughput phenotyping studies.
- 742

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## 1016 **Zhu C, Yang J, Shyu C** (2017) Setaria Comes of Age: Meeting Report on the Second

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