1	Short Title: Shared genetic control of root traits across taxa
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5 6 7	Shared genetic control of root system architecture between Zea mays and Sorghum bicolor
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28	One-sentence summary: Comparisons between root system architecture-associated genes
29	identified from maize and sorghum via GWAS enabled by high-throughput phenotyping reveal
30	conserved functional roles of syntenic orthologs.
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45 Abstract

47 Determining the genetic control of root system architecture (RSA) in plants via large-scale

48 genome-wide association study (GWAS) requires high-throughput pipelines for root

49 phenotyping. We developed CREAMD (Core Root Excavation using Compressed-air), a high-

50 throughput pipeline for the cleaning of field-grown roots, and COFE (Core Root Feature

51 Extraction), a semi-automated pipeline for the extraction of RSA traits from images. CREAMD-

52 COFE was applied to diversity panels of maize (*Zea mays*) and sorghum (*Sorghum bicolor*),

53 which consisted of 369 and 294 genotypes, respectively. Six RSA-traits were extracted from

54 images collected from >3,300 maize roots and >1,470 sorghum roots. SNP-based GWAS

55 identified 87 TAS (trait-associated SNPs) in maize, representing 77 genes and 115 TAS in

56 sorghum. An additional 62 RSA-associated maize genes were identified via eRD-GWAS.

57 Among the 139 maize RSA-associated genes (or their homologs), 22 (16%) are known to affect

58 RSA in maize or other species. In addition, 26 RSA-associated genes are co-regulated with genes

59 previously shown to affect RSA and 51 (37% of RSA-associated genes) are themselves *trans*-

60 eQTL for another RSA-associated gene. Finally, the finding that RSA-associated genes from

61 maize and sorghum included seven pairs of syntenic genes demonstrates the conservation of

62 regulation of morphology across taxa.

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65 Key Words: root system architecture; high-throughput phenotyping; syntenic orthologs

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71 Introduction

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73 The spatial arrangements of root systems, i.e., root system architecture (RSA) (Lynch, 1995), 74 plays a critical role in plant productivity and tolerance to environmental stresses. In maize (Zea 75 *mays*), the majority of the root mass is found in the top 0.3 m of soil (Amos and Walters, 2006). 76 This mass of roots has been referred to as the "root crown" (Trachsel *et al.*, 2011), the "core root" 77 (Grift et al., 2011), or the "core root system" (Hauck et al., 2015). The term core root system is 78 used hereafter for two reasons. First, the term root crown originally referred only to the above-79 ground portion of the root system (Schwarz, 1972; Bray et al., 2006). Only more recently has this 80 term been used to describe roots within the top 0.3 m of soil (Trachsel et al., 2011). Second, this 81 term is easily confused with the term "crown roots", which refers to post-embryonic shoot-borne 82 roots (Kiesselbach, 1949). 83 84 The genetic regulation of root development has been extensively studied in Arabidopsis (Dolan 85 et al., 1993; Birnbaum et al., 2003; Petricka et al., 2012; Petricka et al., 2013) resulting in an in-86 depth understanding of the relevant genes and pathways. Despite similarities in embryonic root 87 systems and some shared mechanisms of genetic regulation, there are major anatomical 88 differences between the root systems of Arabidopsis and cereal crops. The adult Arabidopsis root 89 system comprises a tap root, a basal root, hypocotyl roots, internodal shoot-borne roots, and 90 lateral roots (Zobel, 2016). By contrast, the maize root system is composed of the embryonic 91 primary root and variable numbers of seminal roots, as well as postembryonic shoot-borne and 92 lateral roots (Hochholdinger and Tuberosa, 2009). Similar to maize, sorghum (Sorghum bicolor) 93 develops shoot-borne roots; however, sorghum lacks seminal roots (Singh et al., 2010). Based on 94 these fundamental morphological differences, it is unlikely that a complete understanding of the 95 genetic regulation of RSA in these species can be elucidated from Arabidopsis. 96 97 Whereas functional studies on qualitative mutants of genes with large effect sizes have deepened 98 our understanding of the genetic control and developmental processes of the root systems of 99 cereals, a comprehensive understanding of the genes underlying quantitative variation in RSA

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has not been achieved (Hochholdinger et al., 2018).

102 Genome-wide associated study (GWAS) offers the opportunity to identify genes affecting 103 natural variation of quantitative traits via the association of markers across the genome with 104 phenotypic variation within diversity panels (Xiao et al., 2017). With the ready availability of 105 large numbers of genetic markers, phenotyping has become the bottleneck for GWAS. Several 106 root phenotyping pipelines have been developed for genetic mapping (Topp et al., 2013; Zurek et 107 al., 2015). Most of these studies were conducted on young plants grown in microcosms and 108 mesocosms (Topp, 2016). However, it has been observed in multiple species that RSA varies 109 across development and environments (Rauh et al., 2002; Magalhaes et al., 2004; Trachsel et al., 110 2013) and that roots grown under controlled conditions do not match those grown under field 111 conditions (Poorter et al., 2016). Hence, if we wish to understand the genetic control of RSA as it 112 relates to crop growth in target environments, it is necessary to phenotype roots grown under 113 agronomically relevant field conditions. However, the throughput of current pipelines for 114 analyzing RSA is insufficient to satisfy the phenotyping needs of large-scale GWAS. Hence, 115 thus far, efforts to characterize RSA have mainly focused on quantitative trait locus (OTL) 116 mapping using bi-parental populations with limited genetic diversity, population size, and 117 mapping resolution (Thomson et al., 2003; Giuliani et al., 2005; Li et al., 2005; Liang et al., 2010; 118 Cai et al., 2012; Atkinson et al., 2015; Richard et al., 2015; Guo et al., 2018). To date, few of the 119 genes underlying RSA QTL in cereal crops have been cloned (Mai et al., 2014; Hochholdinger et 120 al., 2018).

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122 Maize and sorghum are both important crops, ranked first and fifth, respectively, in global cereal 123 production (http://faostat.fao.org/). Maize and sorghum diverged from a common ancestor 124 approximately 12 mya (Swigoňová et al., 2004). Approximately 60% of annotated genes are 125 syntenically conserved between these two species and this syntenically conserved set of genes 126 accounts for >90% of all genes characterized by forward genetics in maize (Schnable and 127 Freeling, 2011; Schnable, 2015). Syntenic orthologs are more likely to retain consistent patterns 128 of gene regulation and expression across related species (Davidson et al., 2012), and may be 129 more likely to retain ancestral functional roles than non-syntenic gene copies (Dewey, 2011). 130 However, to date, the conservation of functional roles for syntenic orthologous gene pairs in 131 related species has not been widely tested.

132

- 133 We report the development of CREAMD (Core Root Excavation using Compressed-air), a high-
- 134 throughput pipeline suitable for the excavation and cleaning of field-grown roots, and COFE
- 135 (Core Root Feature Extraction), a semi-automated pipeline to extract features from images of
- 136 roots. CREAMD-COFE was used to phenotype roots from maize and sorghum diversity panels.
- 137 Comparative analyses of maize and sorghum GWAS results provided strong evidence for shared
- 138 genetic control of RSA in these two species and the conservation of functional roles for syntenic
- 139 orthologous gene pairs.
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- 141

142 **Results**

143 144

145 systems 146 Manual excavation and cleaning of field-grown roots is labor intensive (Trachsel et al., 2011; 147 Colombi et al., 2015). To simplify the phenotyping of RSA from field-grown plants and thereby 148 enable large-scale genetic studies under agronomic conditions, we developed CREAMD, a 149 pipeline for the rapid excavation and cleaning of roots. CREAMD uses compressed air to remove 150 soil from core root systems (Supplemental Text S1; Figure 1; Methods). 151 152 Following excavation and cleaning, core root systems were photographed (Figure 1; Methods). 153 COFE, a semi-automated pipeline, was used to extract traits from the resulting images 154 (Supplemental Text S1). COFE is an adaptation of the ARIA software (Pace et al., 2014), which 155 had been developed for lab-based phenotyping of immature root systems. 156 157 There are two major potential sources of error between auto-extracted trait values and ground 158 truth: 1) errors introduced via the projection of 3D traits onto a 2D image; and 2) errors in the

CREAMD-COFE enables the efficient excavation, cleaning, and phenotyping of core root

159 extraction of trait values from 2D images. To distinguish between these two potential sources of 160 error, we compared COFE-extracted trait values to trait values obtained by manually measuring 161 3D core root systems (ground truth) and to trait values manually extracted (using ImageJ) from 162 2D photos of the same core root systems. These comparisons were performed for approximately 163 5% of all collected maize and sorghum core root systems (Methods). The coefficient of determination (r^2) between COFE's auto-extraction trait values and manual measurements of 164 165 maximum width and depth from 3D core root systems are 0.54 and 0.46, respectively. By contrast, the r^2 for the same two traits between COFE's auto-extracted trait values and 166 167 measurements obtained using ImageJ from photos are 0.88 and 0.87, respectively (Supplemental 168 Figure S1; Methods). These results demonstrate that COFE can accurately extract trait values 169 from 2D images of core root systems (Figure 1) and that much of the difference between COFE-170 extracted trait values and ground truth is due to the challenge of representing 3D core root 171 systems in 2D images.

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Figure 1: Extraction of RSA traits from binary images of core root systems using COFE. A) Illustration of root cleaning and B) phenotyping of CREAMD pipeline; C) Illustration of four out of six traits extracted via COFE; D) Comparison of RSA trait values from the inbred line B73 extracted by COFE from roots collected using CREAMD or water-based root cleaning. Bars indicate mean ± SD; Student's t-test; n=15; E) Heritabilities of RSA trait values obtained from the SAM Diversity Panel via CREAMD COFE; n=3,196 roots per view.

173 The air-based root cleaning pipeline, CREAMD, increases the speed of root cleaning 6.5-fold as

174 compared to a previously described (Trachsel et al., 2011) water-based root cleaning pipeline

- 175 (Supplemental Table S1), while yielding comparably intact core root systems; trait values
- 176 obtained from 15 plants of each of four maize genotypes via CREAMD-COFE (Methods) are
- similar to those obtained via the water-based root cleaning pipeline (Figure 1; Supplemental
- 178 Figure S2). In addition to being substantially faster than the water-based root cleaning pipeline
- 179 without comprising root quality, CREAMD can be conducted at remote field sites that lack
- 180 access to water.
- 181

182 **Phenotypic variation of RSA in maize**

183 Three biological replications of 369 inbred lines from the SAM Diversity Panel (Leiboff et al., 184 2015) were grown (Methods). Core root systems from up to three competitive plants (Methods) 185 from each of the three replications were excavated and cleaned using CREAMD. Each core root 186 system was first photographed using a camera angle selected to obtain a view from a neighboring 187 plant in the row in which the plant under analysis was grown (view 1) and then again after 188 rotating the core root system by 90° (clockwise when viewing from above), resulting in view 2 189 (Methods). Trait values of core root systems of maize from the two views did not exhibit 190 statistically significant differences (Supplemental Table S2), suggesting maize plants do not 191 substantially alter their RSA in response to neighbors, at least at the planting densities employed 192 here (Methods). Even so, when viewed from above core root systems do not exhibit radial 193 symmetry (Supplemental Figure S3; Methods). Consequently, for subsequent analyses, we 194 classified the two images of each core root system as the larger and smaller on a per trait basis 195 (Supplemental Figure S4; Supplemental Table S3; Methods).

196

197 COFE was used to extract the following six types of traits from both images of each core root 198 system (Table 1; Figure 1; Supplemental Text S2; Supplemental Figure S4-6). Because we 199 extracted traits from both images of each root, a total of twelve traits were extracted. Maximum 200 and median widths (designated *smMaxWidth*, *lgMaxWidth*, *smMedWidth*, and *lgMedWidth*) 201 served as measures of the horizontal expansion of core root systems. The Adjusted Depth 202 (*smAdjDepth* and *lgAdjDepth*), which is the root depth at which the ratio of root pixels to total 203 pixels exhibits the highest heritability (Supplemental Figure S5) was used a measure of the depth 204 of the core root system. Convex hull (*smConArea* and *lgConArea*), the minimum set of points

- 205 that define a polygon containing all the pixels of a core root in an image, was used to describe 206 the overall expansion of a core root system. The penultimate trait was total root area (*smArea*)
- and *lgArea*), which is the total number of pixels of roots in a photograph.
- 208
- 209 The final extracted trait was root angle. The scientific literature does not offer a consistent
- 210 definition of root angle, particularly among, but even within, species (Vitha et al., 2000; Li et al.,
- 211 2005; Hargreaves et al., 2009; Singh et al., 2010; Courtois et al., 2013; Richard et al., 2015),
- among developmental stages (Omori and Mano, 2007; Fang et al., 2009; Trachsel et al., 2011;
- Pace et al., 2014; Zurek et al., 2015) and across environments (Topp et al., 2013; Uga et al., 2013;
- Huang et al., 2018). Due to the low heritabilities (<0.2) of two previously defined measures of
- root angle (CA) and top angle (IAngRt) (Trachsel *et al.*, 2011; Colombi *et al.*, 2015), we defined
- a root angle trait based on width profiles (*smWPA* and *lgWPA*). High values of *WPA* are
- associated with steep roots. WPA exhibits higher heritabilities (0.50 for lgWPA and 0.52 for
- 218 *smWPA*) than the two previously described root angle traits (Figure 1; Supplemental Text S2;
- 219 Supplemental Figure S6).
- 220

221 The heritabilities of the twelve traits ranged from 0.47 for *smMaxWidth* to 0.61 for *smArea* and

222 *lgArea*, with the exception of *smAdjDepth* and *lgAdjDepth*, which had the lowest heritabilities

223 (0.33 and 0.37) (Figure 1). For five of the six types of root traits (Area, ConArea, MedWdith,

- 224 *MaxWidth, Adjusted Depth*) the two views (large and small) were positively correlated.
- 225 Correlations between larger and smaller views of the collected RSA traits range from 0.92 for
- 226 MaxWidth to 0.98 for Area (Supplemental Table S4). The pair-wise Pearson correlation

227 coefficients ranged from 0.45 (between *smAdjustedDepth* and *smMedWidth*) to 0.97 (between

228 *smArea* and *smConArea*). Both views of *WPA* exhibited negative correlations with all other RSA

traits (Supplemental Table S5).

230

To determine correlations between RSA and above-ground traits, we compared the 12 RSA traits with four above-ground traits: plant height, plant ear height, flowering time (days to anthesis; DTA), and node number data from Leiboff et al. (2015). Even though the root and above-ground traits were collected in different environments, both views of five of the six types of root traits (*Area, ConArea, MedWdith, MaxWidth, Adjusted Depth*) were positively correlated with all four

- above-ground traits. Pairwise Pearson correlation coefficients ranged from 0.36 (between
- 237 *smMaxWidth* and node number) to 0.59 (between *lgArea* and *plant ear height*). Similarly, both
- 238 views of WPA exhibited negative correlations with all four above-ground traits (Supplemental
- Table S5). These correlations between RSA and above-ground traits support the hypothesis that
- 240 by selecting for the latter breeders may have inadvertently selected for the former.
- 241

242 GWAS for RSA traits

243 FarmCPU accounts for kinship and population structure in GWAS (Liu et al., 2016). An efficient 244 implementation of FarmCPU termed FarmCPUpp (Kusmec and Schnable, 2018) was used to 245 perform GWAS on the SAM Diversity Panel, which was previously genotyped with ~1.2 M 246 SNPs (Leiboff et al., 2015). RSA trait values were adjusted to account for field-based spatial 247 variation (Methods). 107 significant SNPs were associated with six types of RSA traits (each of 248 which has two views, resulting in a total of 12 traits) using an FDR cutoff of < 0.05 (Benjamini 249 and Hochberg, 1995) (Supplemental Table S6). Only 20% (20/107) of these trait-associated 250 SNPs (TASs) were associated with both views of the same trait (i.e., 10 pairs of TASs), a result 251 that is consistent with our finding that roots do not exhibit radial symmetry (Supplemental Figure 252 S3). In addition, $\sim 6\%$ (7/107) of the TASs were associated with two or more traits, a result 253 consistent with the high correlations among traits (Supplemental Table S4). For 77/87 of the 254 TASs (88%) it was possible to identify a candidate gene ("SNP-genes"), which was defined as 255 the gene nearest a TAS within a 20-kb window centered on that TAS (Supplemental Table S6). 256

A SNP located within GRMZM2G148937, Big embryo 1 (Bige1) (Figure 2), was associated with

258 the trait *smWPA*. *Bige1*, which encodes a MATE transporter, is one of only eight cloned maize

- 259 genes with a known function in root development (Hochholdinger et al., 2018). A loss-of-
- 260 function mutant of *Bige1* displays increased number of seminal roots and lateral organs during
- 261 vegetative development as compared to wild-type controls (Suzuki et al., 2015). Within the SAM
- 262 Diversity Panel, inbred lines that carry the ALT (i.e., the non-B73) allele of *Bige1* have
- significantly higher mean values of smWPA (p = 0.01) than those that carry the REF (i.e., the
- B73) allele (Figure 2). Based on published trait data (Leiboff et al., 2015), inbred lines
- homozygous for the ALT allele of *bige1* were shorter and flowered earlier than those
- 266 homozygous for the REF allele, a result consistent with those of Suzuki et al. (2015). Inbred



Figure 2: Association of *Bige1* (GRMZM2G148937) with maize *smWPA*. A) Manhattan plot of SNPbased GWAS for *smWPA*; Gene model with the position indicated of the RSA-associated SNP within the intron; B) Representitive root images of inbred lines homozygous for the ALT (non-B73) and REF (B73) alleles of the RSA-assocciated SNP within *Bige1*. Illustrated inbred lines are LH52 (ALT allele) and LH57 (REF allele). C) Distribution of trait values of inbred lines homozygous for the ALT and REF alleles. Student's t-test; stars indicate p < 0.001.

- 267 lines homozygous for the ALT allele of *bige1* also exhibit reduced ear height and plant height
- 268 (Supplemental Figure S7). In addition, based on published shoot apical meristem (SAM)
- 269 phenotypic data (Leiboff et al., 2015), inbred lines that carry the ALT allele of *Bige1* have SAMs
- 270 with larger radii. This is consistent with a previous report that SAM radius is correlated with
- 271 flowering time, ear height, and plant height (Leiboff et al., 2015).
- 272
- 273 Homologs from other species for 10 of the remaining 76 SNP-genes (13%) are known to
- influence RSA (Supplemental Table S7). For example, GRMZM2G143756, a maize homolog of
- an Arabidopsis ABCG transporter, was associated with *lgArea*. Members of a clade of five
- 276 Arabidopsis ABCG transporters are required for the synthesis of an effective suberin barrier in
- 277 roots, and seedlings of the *abcg2 abcg6* and *abcg20* triple mutant of Arabidopsis exhibit fewer
- 278 lateral root primordia and fewer lateral roots than wild-type controls (Yadav et al., 2014). In

279 potato (Solanum tuberosum), ABCG1-RNAi plants exhibit reduced suberin content in root 280 exodermis cells and tuber periderm cells. The lower suberin content leads to reduced root 281 volume (Landgraf et al., 2014). GRMZM2G013128, a maize homolog of the Arabidopsis 282 SMXL3 gene, was associated with variation in both the smMaxWidth and lgMedWidth traits. In 283 Arabidopsis, SMXL3 is highly expressed in root vasculature; double mutants of smxl3; smxl4 and 284 smx13:smx15 exhibit reduced primary root lengths as compared to wild-type controls (Wallner et 285 al., 2017). GRMZM2G013324, a maize homolog of Arabidopsis SHV3, was associated with 286 variation in the trait lgMaxWidth. SHV3 encodes a glycerophosphoryl diester phosphodiesterase 287 (GPDP)-like protein. A mutant of shv3 exhibits a defective root hair phenotype (Jones et al., 288 2006). GRMZM2G400907, a maize homolog of GTE4 in Arabidopsis, was associated with 289 variation in smMedWidth. GTE4 is a Bromodomain and Extra Terminal domain (BET) factor, 290 which functions in the maintenance of the mitotic cell cycle. An Arabidopsis mutant of gte4 291 exhibits significant shorter primary roots and defective lateral roots (Airoldi et al., 2010). 292

293 eRD-GWAS of maize RSA

294 Conventional GWAS uses SNPs as the explanatory variable. By contrast, eRD-GWAS uses gene 295 expression levels as the explanatory variables to associate genes with phenotypic variation (Lin 296 et al., 2017). Because eRD-GWAS has been shown to identify gene/trait associations that are 297 complementary to those identified via SNP-based GWAS (Lin et al., 2017), we also conducted 298 eRD-GWAS.

299

300 RNA-Seq data from 2 cm tips of germinating seedling roots are available for a subset (N=246) of

301 the SAM Diversity Panel (Kremling et al., 2018). eRD-GWAS was conducted on this subset of

302 the SAM Diversity Panel, resulting in the identification of 62 gene-trait associations

303 (Supplemental Table S8). 34% (21/62) of "eRD-genes" are associated with more than two RSA

traits, whereas 42% (26/62) are associated with the two views of the same RSA traits. For

- 305 example, GRMZM2G021410, which encodes a putative alpha/beta-hydrolase superfamily
- 306 protein, is associated with all six root traits. 12 of the 62 unique eRD-genes (19%) have
- 307 homologs in Arabidopsis or Medicago truncatula with known functions in root development
- 308 (Supplemental Table S9). For example, Arabidopsis homologs of four eRD-genes associated
- 309 with variation in the smaller view of root area (*smArea*) of maize (Figure 3) have been associated



Figure 3: Expression levels of three maize homologs of Arabidopsis root-related genes were associated with *smArea* via eRD-GWAS. A) Manhattan plot of eRD-GWAS for *smArea*. Three homologs of Arabidopsis root-related genes: *ZmSGT1*(GRMZM2G105019), *ZmSCN1* (GRMZM2G012814), and *zmWPP2* (GRMZM2G309970) were detected. Correlation coeffecients (r) of expression levels and trait values of *smArea* for the three genes are: -0.23, 0.25 and 0.22, respectively. P < 0.01 for all correlations. B-D) Representitive root images of inbred lines having extremely low and extremely high expression levels of the three candidate genes.

- 310 with root development in Arabidopsis. An Arabidopsis mutant, *sgt1b* (a homolog of
- 311 GRMZM2G105019), exhibits auxin-resistant root growth under low concentrations of auxin
- 312 (Gray et al., 2003). An RNAi mutant of *wpp2* (a homolog of GRMZM2G309970) exhibits
- 313 delayed root development, reduced root length, and fewer lateral roots as compared to wild-type
- 314 controls (Patel et al., 2004). *SCN1* (a homolog of GRMZM2G012814) encodes a RhoGTPase

- 315 GDP dissociation inhibitor (RhoGDI) that restricts the initiation of root hairs to trichoblasts
- 316 (Carol et al., 2005).

317 Network Analyses of RSA-associated Genes

318 Expression quantitative trait loci (eQTL) mapping is used to identify DNA polymorphisms

- 319 associated with variation in gene regulation (Gilad et al., 2008). 110 of the RSA-candidate genes
- 320 were expressed in at least half of the 246 genotypes used for eRD-GWAS. eQTL analyses were
- 321 conducted for each of the 66/77 qualified SNP-genes and each of the 44/62 eRD-genes that
- 322 passed this expression profile criterion (Methods). At an FDR cutoff of < 0.05, 601 eQTL were
- 323 identified for 58/66 (88%) of the SNP-genes and 39/44 (89%) of the eRD-genes (Supplemental
- 324 Table S10). 69/601 (11.5%) and 447/511 (88.5%) of these eQTL acted in *cis* and *trans*,
- 325 respectively (Supplemental Table S11; Methods). 36 of the 97 (=58+39) (37%) SNP-genes and
- 326 eRD-genes are themselves *trans*-eQTL for at least one of the other 61 genes. This level of
- 327 enrichment is statistically significant (p = 2.2e-16, Methods), and suggests the existence of a
- 328 regulatory network involving both SNP-genes and eRD-genes.
- 329

330 To further explore the existence of a regulatory network, a Gaussian Graphical Model (GGM) 331 was used to construct a GGM-based co-expression network for the 246 genotypes using the 332 RNA-Seq data from root tips that had been used in the eQTL analyses, and thereby identify 333 putative regulatory relationships among the 139 RSA-associated genes (77 SNP-genes and 62 334 eRD-genes) and the nine root-related genes (eight cloned maize root-related genes, plus rum1-335 like1, a homeolog of rum1) (Hochholdinger et al., 2018) (Methods). In total, 26 unique RSA-336 associated genes (16/77 SNP-genes and 10/62 eRD-genes) are co-expressed with one or more 337 cloned maize root-related genes. For example, 17 root candidate genes (nine SNP-genes and 338 eight eRD-genes) were included in the GGM-based co-expression network that contains *rth1*, 339 rum1, rul1, and bige1 (Figure 4). The rth1 gene encodes the SEC3 subunit (Wen et al., 2005) of 340 the exocyst complex (Hala et al., 2008) that controls the exocytotic growth of root hair tip. The 341 rum1 gene encodes an AUX/IAA protein and plays key roles in lateral and seminal root 342 formation (Woll et al., 2005; Zhang et al., 2016), whereas *rull* is a homeolog of *rum1* that 343 exhibits 92% sequence identity and shares the canonical features of AUX/IAA protein (Von 344 Behrens et al., 2011). In another module of the GGM-based co-expression network, nine root 345 candidate genes (seven SNP-genes and two eRD-genes) were co-expressed with rth3, rth5, and



Figure 4: Gaussian Graphical Model-based co-expression networks. Two clusters illustrating putative regulatory relationships among RSA-associated genes and cloned root genes. Yellow dots indicate cloned root related genes, green dots indicate genes identified via eRD GWAS, and purple dots indicate genes identified via SNP-based GWAS.

- 346 *rth6* (Figure 4). *rth5* and *rth6* play important roles in cellulose biosynthesis and are involved in
- 347 root hair elongation (Nestler et al., 2014; Li et al., 2016), whereas *rth3* is a member of COBRA
- 348 gene family that is required for root hair elongation and contributes to grain yield (Wen &
- 349 Schnable, 1994; Hochholdinger et al., 2008).
- 350

351 Comparative GWAS for RSA of maize and sorghum

352 Core root systems of up to five competitive plants were also collected and phenotyped using the 353 CREAMD-COFE pipeline for a subset (N = 294) of the Sorghum Association Panel (SAP) (Casa 354 et al., 2008), which will be designated the SAP-RSA (Supplemental Table S12). The SAP-RSA 355 was grown in Mead, NE (Methods) and phenotyped using CREAMD-COFE for the same RSA 356 traits as was done for maize. The heritabilities and pair-wise correlations of these traits in 357 sorghum were similar to maize (Supplemental Figure S8). GWAS for the SAP-RSA was 358 conducted using 205k SNPs from published GBS data (Morris et al., 2013). In total, 132 TASs 359 (comprising 115 unique TASs) were detected for the RSA traits with FDR < 0.05 (Supplemental 360 Table S13). Among the 132 sorghum TASs, 9% (12/132) were associated with multiple RSA 361 traits or two views of the same RSA trait. Whereas the minor allele frequencies (MAFs) of 362 sorghum TASs are similar to those of the maize, the effect sizes of TASs, which is an estimate of 363 the contribution of each SNP to the total genetic variance (Park et al., 2011), from sorghum are

significantly larger than those from maize (p < 0.01) (Supplemental Figure S9), presumably
reflecting the greater statistical power of the maize GWAS, resulting in a greater ability to detect
smaller effect loci.

367

368 The similarities of maize and sorghum RSAs (Yamauchi et al., 1987), in addition to the syntenic 369 relationship of their genomes (Schnable et al., 2011; Schnable et al., 2012) led us to hypothesize 370 that these species have conserved genetic control for RSA. To test this hypothesis, a comparison 371 was conducted between the unique TASs from GWAS for RSA for maize and sorghum. Syntenic 372 genes were identified within 20-kb windows centered on maize TASs and 500-kb windows 373 centered on sorghum TASs (Methods). These window sizes were selected based on average LD 374 values of 10 kb and 250 kb for the SAM diversity and SAP-RSA panels, respectively (Methods). 375 Using an FDR cut-off of <0.05 for both species, seven pairs of syntenic genes were identified 376 (Supplemental Table S14). Based on a permutation test, this is more overlap than would be 377 expected by chance (p = 1e-04, Methods). For example, GRMZM2G028521, annotated as maize 378 citrate transporter 1 (citt1), was identified via SNP-based GWAS for smArea and lgMaxWidth. 379 Its sorghum homolog Sb01g047080 was 138 kb away from the sorghum TAS associated with 380 both smArea and lgMaxWidth (Figure 5). Although some syntenic gene pairs were not associated 381 with the same RSA traits in maize and sorghum, the associated traits exhibited high correlations.

382

383 This is also more overlap than we detected in two pairs of intraspecific GWAS for above-ground 384 traits conducted as controls. First, we conducted GWAS for multiple traits using mostly 385 previously published data from two genetically distinct maize diversity panels grown in separate 386 environments. The Yan panel, which consists of 368 inbred lines, was grown in China (Li et al., 387 2013; Yang et al., 2014), whereas the SAM Diversity Panel (Leiboff et al., 2015) was grown in 388 the US (Methods). These panels do not include any shared inbred lines. Both panels were 389 phenotyped for four traits: plant height, plant ear height, flowering time, and ear length Data for 390 the Yan and SAM Diversity Panels were obtained from Yang et al. (2014) and Leiboff et al. 391 (2015), respectively (except for EL of the SAM Diversity Panel, which is previously unpublished 392 data, see Methods). Through GWAS conducted using an FDR cutoff of <0.05 for both panels, 24 393 and 18 TASs were detected from the Yan and SAM Diversity Panels, respectively (Supplemental 394 Table S15). Using methodology similar to that described for the comparative *interspecific*



Figure 5: **Comparative GWAS between maize and sorghum for** *smArea*. A) Manhattan plots of Chromosome 1from SNP-based GWAS for *smArea* of maize (top) and sorghum (botttom) identified a pair of RSA-associated syntentic genes; homologous sequences are indicated in pink. B. Genomic positions of the syntenic gene pair from panel A. C) Inbred lines of maize (left pair; LH150 and A188) and sorghum (right pair; White Kafir and D940Y) fixed for ALT and REF alleles of the SNPs associated with *smArea*. D) Distribution of trait values of maize (left) and sorghum (right) inbred lines homozygous for the ALT and REF alleles of the SNPs associated with *smArea*. Student's t-test; stars indicate p < 0.001.

- GWAS for RSA (Methods), no overlapping TASs were identified between the two maize panels,
 even using window sizes as large as 100 kb. Next, we conducted another pair of *intra*specific
- 397 GWAS on two diversity panels that consisted of the same inbred lines and that were genotyped
- 398 with the same set of SNPs, but that were grown in different environments and phenotyped by
- different groups. 97% (273/282) of the members of the Maize 282 association panel (Peiffer et
- 400 al., 2014) are a subset of the SAM panel. This subset of 273 inbred lines will be referred as the

401 "Maize273" and "SAM273" panels. Both panels were phenotyped for four traits: plant height, 402 plant ear height, flowering time, and ear length (Methods). 15 and 13 TASs were detected from 403 the Maize273 and SAM273 panels, respectively (Supplemental Table S16). Even though 404 presumably genetically identical inbred lines were analyzed with the same genotyping data, only 405 two overlapping TAS were identified. The number of shared candidates did not increase even 406 when using window sizes up to 100 kb. The absence of shared signals identified via GWAS 407 conducted within a single species and the very small number of overlapping signals within a 408 single diversity panel provides further evidence that the multiple pairs of RSA-associated 409 syntenic genes detected between the two species is significant. 410

- 411
- 412

413 Discussion

414 Accurate phenotyping is an essential component of GWAS. Phenotyping RSA, i.e., the topology 415 and distribution of roots (Lynch, 1995), is challenging due to tradeoffs between throughput and 416 intactness (Topp et al., 2016). To enable high-throughput excavation and cleaning of core root 417 systems, thereby making feasible GWAS for RSA, we developed the CREAMD pipeline, which 418 offers a 6X speed advantage in root cleaning as compared to conventional water-based methods 419 (Trachsel et al., 2011; Colombi et al., 2015), while yielding comparably intact core root systems.

420 In addition, the towable air-compressor, which is the key component of the CREAMD pipeline,

simplifies the phenotyping of RSA in multiple environments, even when a nearby water source is

422 not available. This promises to make the study of genotype-environment interactions (GXE) of

423 RSA feasible.

424

425 Another phenotyping challenge is the complicated topology and structure of RSA, particularly of 426 adult plants. Like others (Trachsel et al., 2011; Topp et al., 2013; Pace et al., 2014), we used 427 multiple 2D images in an effort to capture more of the 3D complexity of RSA. To convert these 428 images to trait values we developed the COFE software, which offers several advantages relative 429 to alternative software packages such as GiA Roots (Galkovskyi et al., 2012) and DIRT (Das et 430 al., 2015). The accuracy and flexibility of the CREAMD-COFE pipeline is supported both by 431 comparisons to ground truth data and the relatively high heritabilities observed across highly 432 diverse germplasm that exhibits highly divergent RSA phenotypes. Because the density cutoffs 433 used for the AdjDepth traits were selected to maximize heritabilities (Supplemental Figure S5) 434 and these cut-offs are likely to be affected by factors such as soil type, crop management, 435 excavation date, and weather, we recommend determining the optimal cut-offs for each 436 independent project.

437

The availability of the CREAMD-COFE pipeline enabled us to conduct high-throughput
phenotyping of RSA traits in diversity panels of adult, field-grown maize and sorghum plants.
After collecting phenotypic data, we employed two complementary GWAS approaches to
identify RSA-associated maize genes. Conventional SNP-based GWAS associate variation in
SNP genotypes across a diversity panel with phenotypic variation. By contrast, eRD-GWAS uses
expression levels of genes as the explanatory variable for GWAS (Lin et al., 2017). The

robustness of eRD-GWAS is demonstrated by the fact that even though read counts obtained
from RNA-seq data from root tips excised from germinating seedlings were used as the
explanatory variable for the RSA of adult field-grown plants, it was still possible to identify
strongly supported candidate genes. Consistent with Lin et al. (2017), few RSA-associated genes
were detected by both GWAS approaches, providing further evidence that the two approaches
are complementary.

450

451 The ability of our pipeline to detect true positives is supported by the finding that homologs of 452 16% (22/139) of the RSA-associated maize genes are known to affect RSA in other species, one 453 of the highest confirmation rates reported in crops (Xiao et al., 2017). In addition, 26 RSA-454 associated genes are co-regulated with genes previously shown to affect RSA and 37% of RSA-455 associated genes are themselves trans-eQTL for at least one other qualified RSA-associated gene 456 (again, significantly more than would be expected by chance). Finally, we detected substantially 457 more pairs of RSA-associated syntenic genes in maize and sorghum than would be expected by 458 chance. In combination, these results provide strong support for the accuracies of our gene/trait 459 associations and demonstrate that the CREAMD-COFE pipeline is sufficiently accurate for use 460 in GWAS.

461

We photographed each core root system from two directions. Initially, we were surprised that there was little overlap between the SNPs or genes associated with a given trait from the two views. However, in contrast to published reports (Colombi et al., 2015) we showed that core root systems are not radially symmetrical. As a consequence of this asymmetry, the two 2D images we captured of a given core root system typically exhibited different trait values. It is therefore not surprising that we often identified different genes as being associated with the same nominal "trait" from the smaller and larger views of the same core root system.

469

470 Seven pairs of syntenic maize and sorghum genes are associated with RSA traits, which is

471 significantly more overlap than would be expected by chance. Chen et al. (2016) used a

472 comparative GWAS approach to identify shared genetic control among maize and rice homologs

473 for biochemical composition of grain and leaves. Their analysis relied upon conservation of

biochemical pathways across taxa. It was not obvious that the regulation of morphology wouldbe shared across taxa as has been demonstrated by the current study.

476

477 There is substantial overlap among the RSA-associated genes detected in maize and sorghum 478 that were grown in different environments but phenotyped by the same group using the same 479 methodology. This is in line with the observation that overall gene expression profiles of maize 480 roots are substantially more influenced by genotype than by environmental stress factors such as 481 drought (Marcon et al., 2017). By contrast, we found no overlap among trait-associated genes 482 from the same panel of maize inbred lines that had been genotyped with the same markers, but 483 were grown in different environments and phenotyped by different groups. Although these two 484 groups were nominally measuring the same traits, the lack of overlap among trait-associated 485 genes suggests that differences in phenotyping methodologies and hence trait values may be a 486 major contributor to differences in GWAS results among experiments.

487

488 Given its fast rate of LD decay, GWAS in maize results in single-gene or near single-gene

resolution. By contrast, as a consequence of its slower rate LD decay as compared to maize

490 (Morris et al., 2013), GWAS in sorghum does not (Li et al., 2015). However, due to the syntenic

491 relationship between maize and sorghum (Schnable et al., 2011; Schnable et al., 2012), our data

492 indicate that GWAS in maize has the potential to identify candidate genes in that control

493 quantitative traits within large chromosomal windows of sorghum.

494

495 More generally, our results suggest that comparative multi-species GWAS has the potential to 496 enhance our understanding of within-species genetic architecture. Indeed, some RSA-associated 497 genes detected in maize but not sorghum may be false-negative associations (and vice-versa). 498 This is because within a given species it may not be possible to detect an association between a 499 gene and a relevant trait as a consequence of (among other factors) low minor allele frequencies, 500 small effect sizes and/or evolutionary histories (Lai et al., 2018). Hence, just as phenotypes of 501 qualitative mutants identified in one species can inform our understanding of gene function in 502 related species (Lin et al., 2012; Huang et al., 2017; Wang et al., 2018), GWAS results from one 503 species have the potential to identify candidate genes in related species that are not detectable via

504 single-species GWAS.

505

506 Conclusion

507 We report on a high-throughput phenotyping pipeline that uses compressed air to harvest and 508 clean roots, thereby overcoming current throughput limitations. We used this approach to 509 phenotype RSA in both maize and sorghum diversity panels and then conducted GWAS. The 510 finding that homologs of 16% (22/139) of the detected RSA-associated maize genes are known 511 to affect RSA in other species (one of the highest confirmation rates reported in any crop) 512 demonstrates the accuracy of our phenotyping and analysis pipeline and suggests that the RSA-513 associated genes detected in this study are worthy of further investigation and exploration for use 514 in crop improvement. Comparisons between high-confidence, RSA-associated genes identified 515 from maize and sorghum via GWAS revealed conserved functional roles of syntenic orthologs in 516 regulating quantitative variation. Our findings suggest that GWAS results from one species have 517 the potential to identify candidate genes in related species that are not detectable in that second 518 species as a consequence of, for example, low minor allele frequencies, small effect sizes, and/or 519 differing evolutionary histories.

520 521

522 Materials and Methods

523

524 Germplasm for GWAS

525 Three fully randomized replications of 380 maize (*Zea mays*) inbred lines were grown at the 526 Iowa State University's Curtiss Research Farm in Ames, IA with a planting date of May 9th 2017 527 and an inter-row spacing of 88.9 cm and an average within row plant-to-plant spacing of 25.4 cm. 528 Only phenotypic data from the 369 lines in the SAM Diversity Panel (Leiboff et al., 2015) were 529 used for GWAS.

530

- 531 For sorghum (Sorghum bicolor), a subset of the Sorghum Association Panel (SAP) (Casa et al.,
- 532 2008) was grown at the Agronomy Farm of University of Nebraska-Lincoln (UNL), Mead, NE
- 533 with a planting date of May 15th 2017 and a planting density an interrow spacing of 72 cm and an
- average within row plant-to-plant spacing of 7.7 cm. Phenotypic data from 294 accessions of the
- 535 SAP, referred to as the SAP-RSA, were used for GWAS.

536

537 <u>CREAMD - Collection and Phenotyping of Core Root Systems</u>

Core root systems, each with approximately 0.3-m³ volume (Hauck et al., 2015), of typically 538 539 three competitive plants (i.e., plants that are not the terminal plant at the beginning or end of a 540 row nor adjacent to a missing plant within a row) within a row were excavated and cleaned on 541 site using a towable commercial air compressor and an AirSpade[®] device (Supplemental Text 542 S2). Up to three maize roots were collected from up to three biological replications (i.e. plants 543 grow in three different one-row plots) for a total of up to nine roots per genotype. Roots of each 544 genotype were collected within one week of the end of flowering for that genotype. For the SAP-545 RSA, typically five competitive sorghum plants within each row were excavated and cleaned between October 12th and 18th (2017) following the same protocol. In most cases for both species 546 547 it was possible to harvest competitive plants. Harvested plants that were non-competitive (i.e., 548 adjacent to a missing plant, or that were a border plant) or that had lodged were recorded for

549 subsequent statistical modeling (see below).

550

551 Cleaned core root systems were imaged on a customized board (40.6 cm x 50.8 cm) covered with 552 blue fabric. Core root systems were positioned on the center of the imaging board and a 553 dimmable 45.7 cm-diameter light ring (Neewer Technology Co., Shenzhen, China) was placed 554 directly beneath the camera lens to provide evenly distributed lighting to reduce shadows. A 555 round orange marker ($\emptyset = 5.1$ cm) and a tag containing an ID number for each plant were placed 556 on the imaging platform next to the corresponding core root system. All images were captured 557 using an EOS 5D Mark III with an EF 24-105 mm f/4L IS USM lens (Canon, Tokyo, Japan), 558 positioned 125 cm above the imaging board surface using an adjustable mount. The camera was 559 controlled using a laptop computer (Latitude 3550, Dell, Round Rock, USA) running EOS 560 Utility 3 software (Version 3.6.30.0) to capture images. Two images from two orthogonal views 561 (North and West) of the core root system were taken based on the spray-painted identifier. 562 Images were stored using JPEG file format. 563

564 For both the maize and sorghum diversity panels, a random 5% of all collected core root systems

565 (149 maize roots and 56 sorghum roots) were chosen for ground truth measurement. The

566 maximum width and depth of core root system were manually measured for both of the two

- 567 orthogonal views (Supplemental Figure S2). In addition, ImageJ (Schneider et al., 2012) was
- 568 used to measure maximum width and depth from the images of the same sets of roots.
- 569
- 570 To determine whether core root systems exhibit radial symmetry, we collected four to six plants
- of the inbred lines B73 and Mo17 from three locations near Ames, IA on September 23rd 2018:
- 572 Curtiss Farm (GPS: GPS: 42°00'N, 93°39'W, planting date: May 31st 2018, planting density:
- 573 ~36 cm within row, 3 m between rows), Marsden Farm (GPS: 42°00'N, 93°47'W, planting date:
- 574 May 23rd 2018, planting density: ~36 cm within row, 3 m between rows) and South Woodruff
- 575 Farm (GPS: 41°58'N, 93°41'W, planting date: June 15th 2018; planting density: ~25 cm within
- 576 row, 75 cm between rows) (Supplemental Figure S3).
- 577

578 COFE - Image Analysis and feature extraction

- 579 For image analysis, we used MATLAB (The Mathworks, Natick Massachusetts, USA) to
- 580 develop an interactive software, <u>Core Root Feature Extraction (COFE)</u>. Captured images were
- analyzed via a two-phase process: pre-processing and trait extraction (Supplemental Text S1).
- 582 During pre-processing, the first visible node above the soil line of a core root system is identified
- 583 by the user. Then, the software automatically generates a binary image of the root according to
- 584 user-defined settings. During automated trait extraction the software uses a blurring and
- thresholding algorithm to prune roots that aberrantly stick out from the core root system and then
- 586 extracts traits from the core root system.
- 587

588 Comparison of CREAMD vs. Water-based Root Cleaning

589 The inbred lines B73, LH185, and PHN46 and the commercial hybrid Hoegemeyer 7089, grown

590 during the summer of 2017 at the Curtiss Farm, were used to compare CREAMD vs. a water-

- 591 based root cleaning pipeline (Trachsel et al., 2011). For each method, 15 competitive plants of
- each genotype were processed at the time of grain harvest on October 24th 2017. The cleaning of
- 593 roots with pressurized air is described in the CREAMD protocol (Methods, Notes S1). For the
- 594 water-based root cleaning, the excavated core root system was soaked in water for ~1 hour and
- then water washed as described (Trachsel et al., 2011; Colombi et al., 2015). Traits were
- 596 extracted using COFE from images of core root systems excavated and cleaned by both methods.

597

598 Comparative GWAS between Maize and Sorghum

599 For the analysis of maize RSA phenotypes, the best linear unbiased prediction (BLUP) of traits 600 extracted from COFE were calculated by treating genotype and planting row as random effects, 601 and lodging and border status as fixed effects using R package 'lme4' v1.1-21 (Bates et al., 2015) 602 (Supplemental Table S17). Broad sense heritability was calculated for all RSA traits for both 603 maize and sorghum (Cai et al. 2012). For the analysis of sorghum RSA phenotypes, means of 604 extracted trait values of all plants having the same genotype were calculated, after removing extreme values, i.e., those that were 1.5X larger than the 3rd quartile (Supplemental Table S18). 605 606 607 To conduct GWAS on the maize SAM diversity and sorghum SAP-RSA panels, we used 1.2M 608 (Leiboff et al., 2015) and 205k (Morris et al., 2013) SNPs, respectively, without filtering for 609 MAF (Bomba et al., 2017). GWAS for both species were conducted using a C++ implementation 610 of FarmCPU (Liu et al., 2016), termed FarmCPUpp (Kusmec and Schnable, 2018). Based on 611 simulation studies, for moderately complex traits, FarmCPU has been reported to have the best 612 metrics for both the detection of gene-trait associations and false-positive metrics (Miao et al., 613 2018). The first three principle components calculated using TASSEL 5.0 were used as 614 covariates to control for population structure (Bradbury et al., 2007). Linkage disequilibrium 615 (LD) values of both panels were calculated using PLINK v1.90 (Purcell et al., 2007). Based on 616 the average rates of LD in the diversity panels, 20- and 500-kb windows centered on TASs were 617 used to identify candidate genes in maize and sorghum, respectively. Maize AGPv2 genes 618 models (Schnable et al., 2009) and sorghum V1.14 genes models (Paterson et al., 2009) that 619 overlapped with the defined windows for each species using the BEDtools software (Quinlan and 620 Hall, 2010) (V2.23.0) were considered to be candidate genes. 621

622 In addition to SNP-based GWAS, eRD-GWAS (Lin et al., 2017) was conducted on a subset of

623 the SAM Diversity Panel (N=246 inbred lines) for which RNA-Seq data from seedling root

- tissue were available (Kremling et al., 2018). Genes with model frequencies over an arbitrary
- 625 cutoff of 0.05 were designated as candidate genes (eRD-genes).
- 626
- Maize and sorghum syntenic genes were identified following the methods of Zhang et al. (2017)
 using the reference genomes RefGen V2 for maize and Sbi1.4 for sorghum (Supplemental Table

S19). The permutation test was conducted by shuffling the maize-sorghum table 10,000 timesand counting the number of pairs of syntenic genes obtained from each trial (Supplemental Table

631 S19)

632

633 eQTL and co-expression network

eQTL analyses were conducted on the same 246 maize inbred lines as were used for eRD-

635 GWAS, and using the same GWAS method (i.e., FarmCPU) and SNPs as were used for the

636 maize RSA GWAS (see above), with the gene expression values as phenotypes and the SNPs as

637 explanatory variables. Only those maize RSA candidate genes expressed in at least 50% of the

638 246 lines were included in this analysis. An eQTL was defined as acting in *cis* if it was within a

639 window that extends 500 kb upstream and 500 kb downstream of the gene it regulates; eQTL

outside this 1-Mb window were defined as acting in *trans*. Ratios of *cis*- and *trans*-eQTL were

relatively stable with window sizes ranging from 50 kb–2 Mb (Supplemental Table S10). The

642 eQTL with the smallest p-value within each 50-kb window was selected for further analyses. The

643 enrichment test of RSA-associated genes and *trans*-eQTL was performed using the "fisher.test ()"

- 644 function in R.
- 645

646 Graphical Gaussian Model-based co-expression networks were constructed using the R package

647 'bnlearn' v4.4.1 (Scutari, 2010) with 5,000 bootstraps implemented with the constraint-based

648 learning algorithm max-min parents and children (mmpc).

649

650 **Comparative** *intra***specific GWAS**

Both phenotypic and genotypic data of the Yan panel were retrieved from MaizeGo

652 (<u>http://www.maizego.org/Resources.html</u>). SNP data for the Yan panel were generated by Li et

al. (2013) from RNA-Seq and MaizeSNP50 BeadChip. Phenotypic data of the Maize 282 panel

- 654 were retrieved from Panzea
- 655 (http://cbsusrv04.tc.cornell.edu/users/panzea/filegateway.aspx?category=Phenotypes).
- 656 Phenotypic data of plant height, plant ear height, and flowering time of the SAM Diversity Panel
- 657 were from Leiboff et al. (2015). Ear length data were collected from two fully randomized
- replications of 369 maize inbred lines from the SAM Diversity Panel (Leiboff et al., 2015) in
- 659 October 2016, at Iowa State University's Curtiss Research Farm (42°00'N, 93°39'W) in Ames,

660	IA, USA (Supplemental Table S20). Genotypic data for both the Maize273 and SAM273 panels		
661	is a subset of the data used for the root-GWAS of the SAM Diversity Panel. GWAS was		
662	conducted with the same protocol as in comparative GWAS between maize and sorghum (see		
663	above section), except an arbitrarily relaxed window of 100 kb, centered on the TAS was used		
664	here.		
665			
666	COFE Software availability: https://bitbucket.org/baskargroup/cofe/src/master/		
 667 668 669 670 671 672 673 674 675 	Accession Numbers The maize sequence data from this article can be found in the GenBank/EMBL data libraries under accession numbers SRP055871. The sorghum SNP data was downloaded from https://www.morrislab.org/data.		
676 677	Supplemental Data		
678	Supplemental Text S1. CREAMD-COFE protocols		
679	Supplemental Text S2. Definition of Width-Profile Angle (WPA)		
680	Supplemental Figure S1. Ground truth validation for trait values extracted from COFE. 298		
681	images from 149 maize plants were analyzed.		
682	Supplemental Figure S2. Comparisons of trait values extracted using COFE from roots of three		
683	genotypes.		
684	Supplemental Figure S3. Maize core root systems grown in three environments (Curtiss,		
685	Marsden, and South Woodruff farms) exhibit a lack of radial symmetry.		
686	Supplemental Figure S4. Classification of images taken from two angles (North and West) into		
687	larger and smaller view on a per trait basis.		
688	Supplemental Figure S5. Illustration of algorithm for determining root depth (AdjDepth) trait		
689	values		
690	Supplemental Figure S6. Width-Profile Angle (WPA) was used to measure root angle.		
691	Supplemental Figure S7. Above-ground trait values of inbred lines homozygous for the ALT		
692	and REF alleles of bige1.		
693	Supplemental Figure S8. Correlations among RSA traits for 294 sorghum inbred lines.		

- 694 Supplemental Figure S9. Minor allele frequency (MAF) and the absolute value of effect sizes
- 695 of maize and sorghum TAS.
- 696 Supplemental Table S1. Time required to process 60 core root systems via CREAMD and
- 697 water-based root cleaning.
- 698 Supplemental Table S2. RSA traits do not exhibit statistically different values between two
- 699 orthogonal views (North and West) of the maize SAM Diversity Panel.
- 700 Supplemental Table S3. Classification of trait values of root area (Area) from two angles
- 701 (North and West) into larger and smaller views on a *per trait* basis.
- 702 Supplemental Table S4. Correlation coefficients between larger and smaller views of RSA
- 703 traits in the maize SAM diversity and sorghum (SAP-RSA) panels.
- 704 **Supplemental Table S5.** Correlations among RSA traits and above-ground traits in maize.
- 705 Supplemental Table S6. Maize TAS and SNP-genes at FDR < 0.05
- 706 Supplemental Table S7. Arabidopsis homologs with known root-related functions of maize
- 707 SNP-genes
- 708 Supplemental Table S8. List of eRD-genes
- 709 Supplemental Table S9. Arabidopsis and Medicago homologs with known root-related
- 710 functions of maize eRD-genes.
- 711 Supplemental Table S10. List of cis- and trans-eQTL.
- 712 Supplemental Table S11. Percentage of cis- and trans-eQTL for qualified maize RSA-
- 713 associated genes using different window sizes
- 714 Supplemental Table S12. List of inbred lines in used in GWAS for maize (SAM Diversity
- 715 Panel) and sorghum (SAP-RSA)
- 716 **Supplemental Table S13.** Sorghum TAS at FDR < 0.05
- 717 Supplemental Table S14. Syntenic maize-sorghum gene pairs detected via comparative GWAS.
- 718 Supplemental Table S15. List of Yan panel and SAM Diversity Panel TAS for four traits (PH,
- 719 PEH, DTA, EL)
- 720 Supplemental Table S16. List of TAS for four traits (PH, PEH, DTA, EL) identified via GWAS
- conducted on the maize273 and SAM273 panels.
- 722 Supplemental Table S17. RSA trait values (BLUP) of maize SAM Diversity Panel.
- 723 Supplemental Table S18. RSA trait values of sorghum SAP-RSA panel.
- 724 Supplemental Table S19. List of syntenic genes.

Supplemental Table S20. Ear length trait values (BLUP) of maize SAM Diversity Panel.
726

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740 for assistance editing the manuscript.

- 744 Table

746 Table 1. Abbreviations of RSA traits.

Abbreviation	Trait
Area	Root Area
ConArea	Convex Hull Area
MedWidth	Median Width
MaxWidth	Maximum Width
WPA	Width-Profile Angle
AdjDepth	Adjusted Depth
Figure Legends	

755 Figure 1: Extraction of RSA traits from binary images of core root systems using COFE. A)

- 756 Illustration of root cleaning and B) phenotyping of CREAMD pipeline; C) Illustration of four out
- 757 of six traits extracted via COFE; D) Comparison of RSA trait values from the inbred line B73
- extracted by COFE from roots collected using CREAMD or water-based root cleaning. Data are
- means \pm SD; ns: not significant, Student's *t*-test; n=15; E) Heritabilities of RSA trait values
- obtained from the SAM Diversity Panel via CREAMD-COFE; n=3,196 roots per view.
- 761

762 Figure 2: Association of Bige1 (GRMZM2G148937) with maize smWPA. A) Manhattan plot

- 763 of SNP-based GWAS for *smWPA*; Gene model with the position indicated of the RSA-
- associated SNP within the intron; B) Representitive root images of inbred lines homozygous for
- the ALT (non-B73) and REF (B73) alleles of the RSA-assocciated SNP within *Bige1*. Illustrated
- ⁷⁶⁶ inbred lines are LH52 (ALT allele) and LH57 (REF allele). C) Distribution of trait values of
- inbred lines homozygous for the ALT and REF alleles. Student's *t*-test; *** p < 0.001.
- 768

769 Figure 3: Expression levels of three maize homologs of Arabidopsis root-related genes were

associated with *smArea* via eRD-GWAS. A) Manhattan plot of eRD-GWAS for *smArea*. Three

771 homologs of Arabidopsis root-related genes: ZmSGT1 (GRMZM2G105019), ZmSCN1

(GRMZM2G012814), and *zmWPP2* (GRMZM2G309970) were detected. Correlation

coeffecients (r) of expression levels and trait values of *smArea* for the three genes are: -0.23,

0.25, and 0.22, respectively. P < 0.01 for all correlations. B-D) Representitive root images of

- inbred lines having extremely low and extremely high expression levels of the three candidategenes.
- 777

Figure 4: Gaussian Graphical Model-based co-expression networks. Two clusters illustrating
putative regulatory relationships among RSA-associated genes (Panel A) and cloned root genes
(Panel B). Yellow dots indicate cloned root related genes, green dots indicate genes identified via
eRD GWAS, and purple dots indicate genes identified via SNP-based GWAS.

782

783 Figure 5: Comparative GWAS between maize and sorghum for *smArea*. A) Manhattan plots

- of Chromosome 1 from SNP-based GWAS for *smArea* of maize (top) and sorghum (botttom)
- 785 identified a pair of RSA-associated syntentic genes; homologous sequences are indicated in pink.

786	B) Genomic positions of the syntenic gene pair from panel A. C) Inbred lines of maize (left pair;
787	LH150 and A188) and sorghum (right pair; White Kafir and D940Y) fixed for ALT and REF
788	alleles of the SNPs associated with smArea. D) Distribution of trait values of maize (left) and
789	sorghum (right) inbred lines homozygous for the ALT and REF alleles of the SNPs associated
790	with <i>smArea</i> . Student's <i>t</i> -test; *** $p < 0.001$.
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