1	Field phenomics reveals genetic variation for transpiration response to vapor
2	pressure deficit in sorghum
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20 ABSTRACT

- 21 Drought adaptation for water-limited environments relies on traits that optimize plant water
- 22 budgets. Limited transpiration (LT) reduces water demand under high vapor pressure deficit
- 23 (VPD) (i.e., dry air condition), conserving water for efficient use during the reproductive stage.
- 24 Although studies in controlled environments report genetic variation for LT, confirming its
- 25 replicability in field conditions is critical for developing water-resilient crops. Here we test the
- 26 existence of genetic variation for LT in sorghum in field trials and whether canopy temperature
- 27 (T_c) is a surrogate method to discriminate this trait. We phenotyped transpiration response to
- 28 VPD (TR-VPD) via stomatal conductance (g_s), canopy temperature (T_C) from fixed IRT sensors
- 29 (T_{Cirt}), and unoccupied aerial system thermal imagery (T_{Cimg}) in 11 genotypes. Replicability
- 30 among phenomic approaches for three genotypes revealed genetic variability for TR-VPD.
- 31 Genotypes BTx2752 and SC979 carry the LT trait, while genotype DKS54-00 has the non-LT
- 32 trait. T_C can determine differences in TR-VPD. However, the broad sense heritability (H^2) and
- 33 correlations suggest that canopy architecture and stand count hampers T_{Cirt} and T_{Cimg}
- 34 measurement. Unexpectedly, observations of g_s and VPD showed non-linear patterns for
- 35 genotypes with LT and non-LT traits. Our findings provide further insights into the genetics of
- 36 plant water dynamics.

37 Keywords: Stress tolerance, crop adaptation, plant breeding, limited transpiration, stomatal

38 conductance, canopy temperature, sorghum.

39 INTRODUCTION

40 Water scarcity is the main threat to agriculture in semi-arid regions. The increasing risk of crop

- 41 failure in such areas is exacerbated by recurrent droughts, which vary over space and time (Yuan
- 42 et al., 2019). Thereby ensuring demand for food and fiber in a world facing climate variability
- 43 and climate change is essential. Adapting with development of climate-resilient crops is a
- 44 promising strategy to cope with water scarcity (Lorite et al., 2018). However, the nature of
- 45 drought adaptation traits is complex (Blum, 2011; Cooper and Messina, 2022) since transpiration
- 46 is driven by many environmental factors. Hence, dissecting these traits requires understanding
- 47 plant response to each variable (Figure 1A). Depending on the soil water status, plants sense
- 48 water stress in leaves or roots, causing partial stomatal closure (Sampaio Filho et al., 2018).
- 49 Water stress in leaves is triggered by vapor pressure deficit (VPD) (Grantz, 1990), while water
- 50 stress in root tissue is stimulated by soil water deficit (Turner et al., 1985).
- 51 Limiting transpiration (LT) during periods of high VPD would cause partial stomatal closure and redistribute plant use of water within the growing season (Sinclair et al., 2005). 52 53 Simulation studies revealed that unlike the non-limited transpiration (non-LT) trait, the LT trait 54 reduces water uptake during the vegetative stage, saving soil moisture for grain filling (Figure 55 1B and 1C). Studies in controlled environments reported genetic variability via breakpoint (LT) 56 and linear (non-LT) transpiration response to VPD (TR-VPD). Under the hypothesis that 57 breakpoint thresholds are heritable, breeding programs could develop commercial hybrids and 58 allocate them to geographies that match the hybrid's breakpoint TR-VPD to increase water 59 productivity. However, studies were pseudo-replicated, and some studies reported the lack of reproducibility between controlled environments and field trials varied for different crops 60 61 (Gilbert et al., 2011; Guiguitant et al., 2017; Shekoofa et al., 2014). Thus, replicability of these findings requires rigorous field testing since TR-VPD phenotypic characterization to-date is 62 63 limited to controlled environments, and most genotypes were not tested in independent field 64 trials (Fletcher et al., 2007; Gilbert et al., 2011; Guiguitant et al., 2017; Schoppach and Sadok,
- 65 2013; Shekoofa et al., 2014; Yang et al., 2012).
- Phenomic approaches, such as whole plant transpiration and gas exchange, to 66 67 discriminate TR-VPD are expensive, time-consuming, and limited for few genotypes (Kirkham, 2014). To accelerate selection, breeding programs require large-scale high-throughput plant 68 phenotyping (HTPP) methods, which can effectively replicate the phenotype identified via 69 traditional methods. Canopy temperature (T_C) is a proxy to estimate plant water stress (Belko et 70 71 al., 2013; Jackson et al., 1981; Jones et al., 2009) due to its relationship with stomatal response 72 (Deery et al., 2019; Leinonen and Jones, 2004), which is grounded in the theory of energy 73 balance (Jackson et al., 1981). Studies under well-watered conditions suggest that warmer T_C and 74 low transpiration can save soil moisture (Pinter et al., 1990). Further, it was hypothesized that 75 genotypes with warmer T_C could yield better in drought environments (Pinter et al., 1990). While 76 some studies suggest T_C as a reliable approach to dissecting water use traits (Anderegg et al.,
- 77 2021; Belko et al., 2013; Mutava et al., 2011) other studies indicate that T_C can be an artifact of
- canopy architecture or micro-environmental variation (Prashar and Jones, 2014).

- 79 Sorghum is grown and adapted to water limitations. Despite this advantage, its full
- 80 potential still needs to be explored to identify traits contributing to water productivity (Borrell et
- 81 al., 2014; Vadez et al., 2014). This study aims to validate and identify the TR-VPD in field trials
- 82 in the target environment of the sorghum-producing region of the United States, predominantly
- 83 overlaying the semiarid Great Plains. We hypothesize that sorghum has a genetic variation for
- 84 TR-VPD under field conditions. Under this hypothesis, our results will be comparable to
- 85 previous studies. First, we expect low stomatal conductance (g_s) and high T_C for genotypes with
- 86 the LT trait and the opposite for genotypes with the non-LT trait (Figure 1D). Second, we expect
- 87 observations to fit breakpoint and linear TR-VPD for genotypes with LT and non-LT traits,
- 88 respectively. Our second hypothesis is that T_C is a surrogate method to discriminate TR-VPD.
- 89 Under this hypothesis, first we expect a significant negative correlation between g_s and T_C .
- 90 Second, no significant correlation between T_C and canopy architecture traits.

91 MATERIALS AND METHODS

92 Design and management of field experiments

- 93 This study tested the TR-VPD of various genotypes in Ashland Bottoms (AB), Kansas and
- 94 Greeley (GR), Colorado (Figure 2A). Based on the PRISM dataset
- 95 (https://prism.oregonstate.edu/), the maximum VPD in summer in AB is 2.5 and 3.5 in GR
- 96 respectively, corresponding to subhumid and semiarid climates (Figure 2B and 2C). Field trials
- 97 were planted in the summer seasons of 2019, 2020, and 2021 in AB and GR. The experiment
- 98 included genotypes with putative TR-VPD and commercial hybrids (Table 1). In 2019 and 2020,
- 99 the field trials comprised 10 genotypes and only 6 in 2021 (Table 1). Fewer genotypes were
- 100 planted in 2021 because the expected TR-VPD (Figure 1D) in AB19 and AB20 matched only for
- 101 genotypes DKS54-00, Tx430, BTx2752 and SC979.
- 102 Experiments were planted under a randomized complete block with four replications in
- 103 2019 and 2020. In 2021, experiments comprised two water treatments (irrigated and rainfed)
- 104 planted in a split-plot design. In these experiments, the rainfed treatment tested genotypes'
- 105 performance under drought conditions. Experiments were irrigated to isolate transpiration driven
- 106 by atmospheric VPD and avoid confounding effects due to low soil moisture (Turner et al., 1985;
- 107 Zhang et al., 2018). In AB19 and AB20, experiments were irrigated daily from full canopy cover
- to flowering time using drip lines. In GR21, the experiment was irrigated every week from
- 109 planting to hard dough stage. Irrigations at this site aimed to fill soil moisture up to 100%.

110 Atmospheric and soil variables

- 111 In each experiment, a weather station (ATMOS 41, METER Group, Inc., USA) recorded
- 112 temperature, relative humidity, solar radiation, precipitation, wind speed, and VPD at 15-minute
- 113 intervals. In 2019, soil moisture sensors (TEROS 10, METER Group, Inc., USA) were installed
- 114 at two depths (15 cm and 30 cm) in a plot of genotype 84G62. In 2021 three soil moisture
- sensors were installed on three experimental units at 30 cm. In Colorado (GR21), the seasonal
- 116 precipitation was 107 mm; in Kansas, it ranged from 338 (AB21) to 520 mm (AB19). Over three
- 117 years of assessment and across all locations, the maximum VPD ranged from 1.1 (AB) to 4.2 kPa
- (GR), and soil moisture at 30 cm ranged from 0.20 m³ m⁻³ to 0.31 m³ m⁻³. Experiments with the

- 119 lowest and highest VPD were AB19 and GR21, respectively (Figure 2B and 2C). General
- 120 characteristics of the field site, crop management, and environmental conditions for each season
- 121 are listed in Table S2.

122 Stomatal conductance phenotyping

- 123 A steady-state SC-1 leaf porometer measured (METER Group, Inc. USA) the water diffusion in
- 124 the stomatal cavity (Kirkham, 2014). Abaxial stomatal conductance (g_s) was taken on the third
- 125 fully expanded sunlit leaf on three representative plants per plot during the vegetative stage.
- 126 Information was taken between 12:00 to 16:00 hrs. Evaluations were done without cloud cover
- 127 to ensure observed transpiration was due to atmospheric VPD and not poor light quality. In
- 128 GR21, stomatal conductance was evaluated 24 hours after irrigation. Environmental conditions
- 129 (Table S2) for each evaluation were pulled from the ATMOS 41 weather station.

130 Canopy temperature phenotyping

- 131 Canopy temperature (T_C) was recorded via proximal sensing by infrared thermometer (IRT)
- 132 sensors fixed on the ground and remote sensing by thermal cameras carried by an unoccupied
- 133 aerial vehicle (UAV). A network of fixed IRT sensors wirelessly transmitted T_C measurements to
- a data logger (Dynamax Inc.). T_C collected by IRT sensors (T_{Cirt}) was recorded in three
- replications in AB19 and two replications in AB20 and GR21. Each IRT sensor has a 20-degree
- field of view and covers a circle area on the target with a diameter-to-sensor distance ratio of 1:3
- and a 0.5 °C accuracy over a temperature range of 0 °C to 50 °C. In this study, IRT sensors were
- 138 installed at around 1.5 m above the crop canopy, hence covering circle areas on the canopy with 120
- 139 a $0.5m^2$ diameter. Measurements of T_{Cirt} were analyzed from full canopy cover to pre-flowering
- 140 time to avoid any confounding effect of soil or pollen/flower temperature. For quality control,
- 141 records of T_{Cirt} (minutes) were aggregated at an hourly time step, and the time series was plotted 142 for visual inspection as a quality control.
- 143 A UAV high-throughput plant phenotyping (HTPP) system, including a quadcopter
- 144 (Matrice 100, DJI, Shenzhen, China) and thermal camera (VUE Pro R, FLIR, USA), were
- 145 integrated to collect thermal images for T_C extraction in AB19, AB21, and GR21 field trials.
- 146 Raw thermal images were collected under different heat conditions within the same day (i.e.,
- 147 morning, noon, and afternoon) to monitor T_C variation between genotypes (Table S2). Aerial
- 148 image overlap rate between two geospatially adjacent images was set to 80% both sequentially
- and laterally to ensure optimal orthomosaic photo stitching quality. All data collection flights
- 150 were operated at 35m above ground level at 3.5 m s^{-1} . After taking off, the quadcopter will hover
- at a waypoint outside the field trials for at least 2 min before real image collection, allowing the
- 152 thermal camera to self-calibrate under relatively stable ambient air condition. Multiple ground
- temperature reference panels (Figure 3B) for thermal imaging calibration were placed beside the
- 154 field trials at least 20 min. before the real data collection. Ground reference temperature values
- 155 were measured and recorded in a data logger (CR3000, Campbell Scientific, USA). A semi-
- automated image processing pipeline was used to generate field orthomosaic photos and trait
- 157 data extraction (Wang et al., 2020). Plot-level T_{Cimg} is the median of all T_C values extracted

158 within each manually-generated plot boundary. The two T_C methods were compared using the

159 RMSE (Wallach et al., 2014).

160 In-season field field phenotyping

- 161 After seedling emergence, crop establishment was scored via plant density (Figure S6). In AB19,
- 162 the plant density ranged from 19 to 25 plants m^{-2} . Even though seeding rate emergence was
- 163 similar in AB20 and GR21 (Table S2), plant density differed in both experiments. For instance,
- in AB20 and GR21, plant density ranged from 24 to 35 and 13 to 19 plants m⁻², respectively. In
- all experiments, commercial hybrids had the highest plant density; while the Tx430 genotype, an
- 166 inbred line, had the lowest plant density (Figure S6).
- 167 Flowering time was evaluated each year and determined when 50% of the two central
- rows were flowering at 50%. In AB19 and AB20, one plant from each experimental unit was
- harvested around flowering to estimate leaf area. In 2019, the leaf area per plant was calculated
- using LI-3100C (LI-COR, Inc, USA). In AB20, the leaf area and size of each leaf across the
- 171 canopy profile were estimated using photos processed with the ImageJ program (Rueden et al.,
- 172 2017). In AB20, photographs of a plant on each experimental unit were taken before flowering.
- 173 These photographs were printed to measure the leaf angle with a protractor. The angle of the
- adaxial leaf relative to the stalk was measured in the middle of the canopy. In AB19, AB21, and
- 175 GR21, the total biomass of a single plant was harvested at physiological maturity to estimate
- 176 grain weight and harvest index (HI). Similarly, in AB19 and GR21, plant height, panicle length,
- and panicle exertion were measured on three plants per experimental unit.

178 Statistical analysis

- 179 Differences between putative traits (non-LT vs. LT) and among sorghum genotypes are expected
- 180 to occur under high VPD. For this reason, we analyzed records of g_s and T_{Cirt} between 12:00 to
- 181 16:00 hours (periods with high solar irradiation). Information on T_{Cirt} was recorded from June 23
- to August 10 in AB19 and August 11, 15, 18, and 19 in AB20. Each year the variance was
- 183 analyzed with a mixed model in two stages: first, to test the size effect of the putative trait (eq.
- 184 1), and second the size effect of each genotype (eq. 2). The study conducted an additional
- 185 analysis for genotypes evaluated over all experiments (eq. 3). The models were specified as
- 186 follows:
- 187 $Y_{ijklm} = \mu + T_i + G_j + H_k + D_l + r_m + \varepsilon_{ijklm}$ (Eq. 1)
- 188 where Y_{ijklm} is the response variable in the m^{th} block in the l^{th} day in the k^{th} hour in the j^{th}
- 189 genotype with the *i*th trait, μ is the grand mean, T_i is the fixed main effect of the *i*th trait, G_i is the
- random main effect of the j^{th} genotype, H_k is the random effect of k^{th} hour, and r_m is the random effect of the m^{th} replicate.

$$Y_{jklm} = \mu + G_j + H_k + D_l + r_m + \varepsilon_{jklm}$$
(Eq. 2)

192 $Y_{ijklm} = \mu + G_i + E_j + H_k + D_l + r_m + \varepsilon_{ijklm}$ (Eq. 3)

- 193 where Y_{jklm} is the response variable in the m^{th} replicate in the l^{th} day in the k^{th} hour in the j^{th}
- 194 genotype, Y_{ijklm} is the response variable in the m^{th} block in the l^{th} day in the k^{th} hour in the j^{th}
- environment with the i^{th} genotype, G_i is the fixed main of the i^{th} genotype, E_j is the fixed effect of
- 196 the j^{th} environment, H_k is the random effect of k^{th} hour, and r_m is the random effect of the m^{th}
- 197 replicate. A pairwise comparison (Sidak test) was performed when the *F* value fell below $\alpha =$
- 198 0.05 significance threshold.

Broad sense heritability (H^2) was estimated for g_s , T_{Cirt} , and T_{Cimg} . H^2 was estimated with the lmer library (Bates et al., 2005) by fitting a random effects model which estimated variance components for each random factor on each experiment (Eq. 4) and across experiments (Eq. 5):

$$Y_{iklm} = \mu + G_i + H_k + D_l + r_m + \varepsilon_{ijkm}$$
(Eq. 4)
$$Y_{ijklm} = \mu + G_i + E_j + (GE)_{ij} + H_k + D_l + r_m + \varepsilon_{ijklm}$$
(Eq. 5)

where Y_{iklm} is the response variable in the m^{th} replicate in the l^{th} day in the k^{th} hour with the i^{th} 203 genotype, Y_{ijklm} is the response variable in the m^{th} replicate in the l^{th} day in the k^{th} hour in the j^{th} 204 environment with the i^{th} genotype, G_i is the random effect of the i^{th} genotype, E_i is the random 205 effect of the j_{th} environment, $(GE)_{ii}$ is the random two way interaction of i^{th} genotype j^{th} 206 environment, H_k is the random effect of k^{th} hour, and r_m is the random effect of the m^{th} replicate. 207 208 The H^2 for each experiment and across experiments was calculated with Eq 6 and 7, respectively. 209 The variance for environmental effects H and D were disregarded in the analysis since they do 210 not contribute to selecting the target phenotype.

$$H^{2} = \frac{\overline{\sigma}_{G}^{2}}{\overline{\sigma}_{G}^{2} + \frac{\overline{\sigma}_{E}^{2}}{n_{r}}} \quad (\text{Eq. 6})$$
211

202

$$H^{2} = \frac{\overline{\sigma}_{G}^{2}}{\overline{\sigma}_{G}^{2} + \frac{\overline{\sigma}_{G:E}^{2}}{n_{E}} + \frac{\overline{\sigma}_{E}^{2}}{n_{E} \times n_{r}}} \quad (\text{Eq.7})$$
212

213 where: σ_G^2 , $\sigma_{G:E}^2$, and σ_{ϵ}^2 represent variance components of *G*, $G \times E$, and error, respectively. 214 While, n_E is the number of environments, and n_r is the number of replicates.

Information on g_s, T_{Cirt}, and canopy–air difference was paired with corresponding VPD.
Next, g_s was analyzed via a linear (Eq. 8), segmented (Eq. 9 and Eq.10), and asymptotic (nonlinear, Eq.11) regression analysis.

$$Y = \beta_0 + \beta_1 V P D + \varepsilon \tag{Eq. 8}$$

If
$$VPD \le BP$$
: $Y = \beta_0 + \beta_1 VPD + \varepsilon$ (Eq. 9)

If
$$VPD > BP$$
: $Y = \beta_0 + \beta_1 BP + (\beta_1 + \beta_2)(VPD - BP) + \epsilon$ (Eq. 10)

219
$$Y = a + (a - b)e^{-cVPD}$$
 (Eq. 11)

220 where Y represents g_s , BP is the VPD breakpoint, and β_0 is the intercept, β_1 is the slope, and β_2 is 221 the second slope, a is the plateau (maximum attainable value) for g_s , b is the initial g_s when VPD 222 is zero, c is the inflexion point or the relative rate of increase for VPD when g_s increases. The 223 reason for testing different regressions was to identify the best model that fitted the observed 224 data. Linear regressions were performed using lm library (Bates et al., 2015), the breakpoint 225 regression was analyzed with the library segmented (Muggeo, 2016), non-linear regressions were 226 analyzed with libraries nlme, drc, and aomisc (Ritz et al., 2015). The dependence of g_s on VPD 227 was tested via 1000 permutations for linear and breakpoint regressions. To determine this 228 dependence for each genotype, the distribution of the regression coefficient for these 229 permutations was plotted and contrasted against the observed coefficients, β_1 for linear 230 regression; β_1 , β_2 , and BP for segmented regressions. The likelihood-ratio test (library lmtest),

and residual standard error (rse) indicated the accuracy of the best regression model

232 (Archontoulis and Miguez, 2015; Kuznetsova et al., 2017).

233 The effect of canopy architecture traits (i.e., leaf area and leaf size) on transpiration VPD

response were visualized using a principal component analysis (PCA) and a correlation analysis.

The PCA was estimated with the R prcomp function ("R Core Team," 2017) and conducted only

for AB20 since most canopy architecture traits were evaluated in this experiment. In thisanalysis, the leaf area represents the maximum leaf area per plant, which occurs around

- flowering time. The leaf size corresponds to the largest leaf across the canopy profile (Figure
- 239 S3). The leaf angle is at the attachment between the adaxial leaf and stalk in the central part of
- the canopy. Plant density represents the number of plants scored after seedling emergence
- 241 (Figure S6). A correlation analysis was conducted for AB19 and GR21. Grain weight and harvest
- index for experiments in AB19, AB21, and GR21 were analyzed to characterize agronomic traits
- 243 of potential donors of putative LT trait.

244 **RESULTS**

245 Classification of genotypes for their TR-VPD

- 246 To test if VPD during the experiment's growing seasons represented a typical VPD range, we
- compared the average daily maximum VPD over 22 years (2000-2021) against the daily
- 248 maximum VPD recorded on each experiment (Figure 2B and 2C). Results indicate that growing
- 249 seasons AB19 and AB20 underrepresented average VPD in Kansas; while the growing season in
- GR21 represented a regular season. In AB19 and AB20, the VPD ranged between 0.2 to 1.9 kPa
- during the evaluation period (canopy closure to pre-flowering time, Figures 2B). Otherwise in
- 252 GR21 the VPD ranged between 2.5 to 4.2 kPa.

253 To determine the existence of genetic variability for TR-VPD in field conditions, we 254 tested i) the putative non-LT versus LT classification and ii) the genotype classification in each 255 experiment (Table 1). To confirm this classification, significance was expected for the fixed 256 effect and differences among mean groups in post hoc analysis. Specifically, higher gs for the 257 putative non-LT trait and higher T_{Cirt} and T_{Cimg} for the LT trait (Figure 1D). Results in all 258 experiments indicated no significance for the trait effect for gs, T_{Cirt}, and T_{Cimg} (Figure 4, upper panels). However, as expected, the putative non-LT group exhibited slightly high gs (0.06 mol m⁻ 259 260 2 s⁻¹) and lower T_{Cirt} (-0.1 C), as indicated in Table S3.

261 The analysis of variance for each experiment where genotype was the fixed effect 262 indicated differences (p < 0.001) for g_s, T_{Cirt}, and T_{Cimg} (Figure 4). In AB19 (Figure 4A, lower 263 panel), as expected, genotypes with putative LT (BTx2752, 0.62 mol m⁻² s⁻¹) and non-LT 264 (DKS54-00, 0.77 mol $m^{-2} s^{-1}$) differed significantly. In this experiment, a genotype classified as LT (SC803) trait unexpectedly exhibited higher g_s. In AB20 (Figure 4B, lower panel), the mean 265 266 comparison indicated three groups that differed in g_s. The first group comprised a genotype with 267 the LT trait SC979, which had the lowest g_s (0.73 mol m⁻² s⁻¹). The second group included genotypes with LT (Macia, BTx2752, and BTx2363). The third group had genotypes with non-268 269 LT (DKS54-00, Tx430) and LT (Tx7000, BTx642, and SC803) traits, which had the highest g_s. 270 In GR21 (Figure 4C, lower panel), the genotype Tx430 with the non-LT trait unexpectedly exhibited the lowest g_s (Tx430, 0.72 mol m⁻² s⁻¹). As expected, the genotype DKS54-00 with the 271 272 non-LT trait had the highest (0.90 mol $m^{-2} s^{-1}$) g_s. Despite genotypes with the LT trait, BTx2752, 273 and SC979, presenting lower g_s than DKS54-00, they were not significantly different from this 274 genotype.

275 In AB19 (Figure 4D, lower panel), phenotypes identified via T_{Cirt} confirmed that six of 276 nine genotypes matched the expectation (Table 1). Genotypes with the LT trait SC979 (29.9 °C), 277 BTx2752 (29.9 °C), Macia (29.9 °C), and SC803 (30.0 °C) had higher T_{Cirt} and differed from 278 genotypes with the non-LT trait DKS54-00 (29.5 °C) and Tx430 (29.6 °C). In AB20 (Figure 4E, 279 lower panel) T_{Cirt} confirmed the classification for Tx7000 (LT), SC979 (LT), BTx642 (LT), 280 BTx2752 (LT), and Tx430 (non-LT). But, it contradicted the classification for genotypes SC803 281 (LT), Macia (LT), BTx623 (LT), and DKS54-00 (non-LT) (Figure 1D). Genotypes with the LT (BTx2752, 29.0 °C) and non-LT (Tx430, 28.1 °C) traits exhibited the highest and lowest T_{Cirt}, 282 283 respectively. In GR21, all genotypes matched the expectation. The lowest and highest canopy 284 temperature corresponded to genotypes ADV2275 and BTx2752, respectively. In all 285 experiments, T_{Cirt} was replicable only for genotypes with the putative LT trait BTx2752 and 286 SC979.

As expected, in AB21, T_{Cimg} for genotypes with the non-putative-LT trait (DKS54-00, DKS28-05, and SC1345) was lower overall than for genotypes with the putative LT trait (SC803, Macia, BTx623, BTx2752). Discrepant results were obtained for genotype SC979, for which high T_{Cimg} was expected but had low T_{Cimg} , and for genotype Tx430 for which low T_{Cimg} was expected but exhibited high T_{Cimg} . In AB21 (Figure 4H, lower panel) and GR21 (Figure 4I, lower panel), genotypes SC979 and BTx2752, classified as genotypes with the LT trait, exhibited

significantly higher TCimg than genotype DKS54-00, classified as a genotype with the non-LT
 trait. The only genotype that contradicted its non-LT classification in both experiments was

trait. The only genotype that contradicTx430, which had the highest TCimg.

- 296 Over three years of field experiments g_s and T_{Cirt} significantly differed (p < 0.001), and
- 297 the covariate effect of VPD on g_s was non-significant (Table S5). The pairwise comparison for g_s
- 298 revealed that genotypes DKS54-00 (0.81 mol $m^{-2} s^{-1}$), BTx2752 (0.78 mol $m^{-2} s^{-1}$), and SC979
- 299 (0.72 mol m⁻² s⁻¹) belonged to different groups (Table S5). Results of g_s for Tx430 (non-LT) 300 were statistically similar to genotypes BTx2752 (LT) and SC979 (LT) (Table S5). The pairwise
- 300 were statistically similar to genotypes BTx2752 (LT) and SC979 (LT) (Table S5). The pairwise 301 comparison for T_{Cirt} showed two groups. Genotype DKS54-00 with the non-LT trait had lower 302 T_{Cirt} (28.2 °C) than genotype BTx2752 with the LT trait (28.6 °C). The analysis of T_{Cirt} for four
- 303 genotypes over three years matched with the TR-VPD classification (Table S5).
- To test that different phenomic approaches capture the biology of TR-VPD, we estimated the broad sense of heritability (H^2) and expected similar H^2 for g_s, T_{Cirt}, and T_{Cing}. However, H² differed among phenomic approaches in each experiment (Figure 5). In AB19, the highest and lowest H^2 corresponded for g_s (0.5) and T_{Cing} (0.06). Estimates of H^2 Across all experiments for each method indicate a greater H^2 for g_s (0.4), followed by T_{Cing} (0.3) and T_{Cirt} (0.2).

309 Genotypic variation for stomatal response to VPD

- 310 To test that transpiration is driven by VPD and not due to soil water deficit, we compared
- 311 stomatal response in rainfed and irrigated treatments. This comparison was conducted on a day
- 312 when the VPD ranged from 3.2 to 4.0 kPa in GR21 (Figure 6A). In this comparison, we expected
- 313 a significant difference (p < 0.05) between treatments, and observations confirmed this
- 314 expectation. On average, g_s under rainfed treatments was 50% lower than in irrigated treatments.
- 315 In this VPD range, the regression slope for g_s in the irrigated treatment was close to zero (0.006
- 316 mol m⁻² s⁻¹ kPa); in the rainfed treatment, g_s declined (-0.21 mol m⁻² s⁻¹ kPa) as VPD increased.
- 317 Next, we conducted a regression analysis for each genotype. The stomatal response to VPD was
- analyzed via linear, segmented, and non-linear models. In all experiments, g_s ranged between 0.3
- to 1.2 mol m⁻² s⁻¹ and VPD between 1 to 4.5 kPa (Figure 6). The mean g_s conductance of 0.79,
- 320 0.86, 0.82, and 0.82 mol m⁻² s⁻¹ corresponded to experiments AB19, AB20, AB21, and GR21,
- 321 respectively. Unexpectedly, high VPD did not increase the mean g_s for genotypes with the
- 322 putative non-LT trait (DKS54-00 and Tx430).
- 323 Linear regression for genotypes 84G62 (NN), DKS54-00 (non-LT), BTx2752 (LT), and
- 324 SC979 (LT) indicated a positive slope but an unexpected negative one for genotype Tx430 (non-
- LT) (Figure 6B, Table 3). Segmented (breakpoint) regression for all genotypes suggested a
- 326 restriction in g_s between 1.3 to 2 kPa (Figure 6C). Genotype Tx430, which has a putative non-LT
- trait, had the lowest breakpoint and the most negative second slope. Breakpoint patterns were
- 328 similar for commercial hybrids 84G62 (NN) and DKS54-00 (putative non-LT) indicating a
- negative value for the second slope. Genotypes with the putative LT trait SC979 and BTx2752
- exhibited similar patterns, revealing positive values for the first and second slope (Figure 6C).
- 331 Non-linear regressions revealed an asymptotic curve fitting the observed patterns for all
- 332 genotypes 84G62, DKS54-00, BTx2752, and SC979, except Tx430. A non-linear regression

revealed that differences on transpiration under high VPD are represented via the maximum
plateau and the inflection point. Parameters for each regression and genotypes are indicated on
Table 3.

- 336 To test that the g_s is dependent on VPD in linear regression, we compared the observed 337 versus permuted slopes (Figure S3). For genotypes with the non-LT trait DKS54-00 and Tx430, 338 we expected the observed slope to be positive and significantly higher than permuted slopes. For 339 genotypes with the LT trait BTx2752 and SC979, which restrict gas exchange at high VPD, we 340 expected the observed slope to be non-significantly different from the permuted slopes. 341 However, observations contradicted our expectation, indicating that g_s and VPD are independent 342 for DKS54-00 ($\beta_1 = 0, p = 0.1$), while g_s depends on VPD, positive slope, for genotypes BTx2752 and SC979 ($\beta_1 = 0, p < 0.05$). Similarly, a permutation analysis for a breakpoint 343 344 regression revealed that the regressed variable (g_s) is independent of the regressor (VPD, Figure 345 S4). To determine the goodness of fit for different regression models, we used the likelihood 346 ratio test. In this analysis, we expected significance for linear regression for genotypes with the 347 non-LT, whereas significance for a breakpoint regression for genotypes with the LT trait. 348 Unexpectedly, breakpoint and non-linear regressions fitted the observed data (p < 0.001) for a 349 genotype with non-LT trait DKS54-00 (Figure 6). Otherwise, for a genotype with non-LT trait 350 Tx430, the linear and breakpoint regression revealed no significant difference (p > 0.05).
- 351 Similarly, all regressions were not significantly different for genotypes with LT trait BTx2752
- 352 and SC979 (p > 0.05, Figure 6).

353 Canopy temperature (T_c) as surrogate method to discriminate TR-VPD

354 To test if T_C is a surrogate method of g_s, and not an artifact of canopy architecture trais, a PCA 355 was conducted for AB20 and a correlation analysis for AB19 and GR21. In AB20, PC1, PC2 and PC3 represented 38.9, 23.9 and 22.1% of the total variability in the data (Figure 7). PCA1 had a 356 357 moderately positive loading for leaf area (0.4) and leaf size (0.4) and a negative loading for plant 358 density (-0.4) and T_{Cirt} (-0.4). PCA2 had a positive and negative loading with leaf size (0.5) and 359 g_s (-0.6), respectively. PCA3 is dominated by negative loading of the number of leaves (-0.7) and positive loading for T_{Cirt} (0.4). The biplot for PC1 versus PC2 and PC2 versus PC3 revealed that 360 361 T_{Cirt} and g_s are in distinct quadrants suggesting a moderate negative correlation (Figure 7SA) for 362 these phenotypes. Similarly, in GR21, the correlation between gs versus T_{Cimg} was negative and 363 significant (r = -0.9, p < 0.05). A biplot PC2 versus PC3 showed a substantial correlation between T_{Cirt} and the number of leaves. These results indicate that T_{Cirt} is a good proxy of g_s, but 364 365 canopy architecture can affect T_c. Otherwise, biplots suggested a strong correlation among 366 canopy architecture traits. For instance, leaf area negatively correlated with the number of leaves 367 (PC1 versus PC2) and leaf angle (PC2 versus PC3).

- 368To test if TR-VPD phenotype is an artifact of stand count, we conducted correlations369between g_s , T_{Cirt} , T_{Cimg} , and plant density (Figure S7). In AB20, the correlation was close to zero370for plant density and T_{Cirt} , while a negative correlation (-0.5) between plant density and g_s was371non-significant (p > 0.05). Significant negative correlations between plant density and T_{Cimg}
- 372 (Figure S7B and S7C) were obtained in AB19 (r = -0.8, p < 0.01) and GR21 (r = 0.9, p < 0.05).

- 373 Likewise, in GR21 (Figure S7C), the correlation between plant density and g_s was significant (r 374 = 0.9, p < 0.01). Overall, results suggest that stand count is the main trait that can hamper the
- 375 effectiveness of any phenomic approach when dissecting TR-VPD.

376 DISCUSSION

- 377 Testing the existence of genetic variability for the putative LT trait via different phenomic
- 378 approaches is pivotal when deciding to include this trait in a breeding pipeline. This is the first
- 379 study that phenotyped the TR-VPD in field trials in sorghum production regions of the United
- 380 States. Our findings i) help to better understand the challenges when testing the replicability of
- 381 the trait via different phenomic approaches, ii) discuss the best representation of genetic diversity
- 382 by comparing different regression models, iii) review the underlying physiological mechanism of
- 383 the putative LT trait and iv) outline future steps for crop breeding when using the LT to deliver
- 384 water efficient sorghum hybrids.

385 A phenotype can be similar or vary among phenomic approaches

- 386 Our study confirmed the TR-VPD classification via three phenomic approaches for three out of 387 eleven genotypes (Figure 4). Results for genotypes SC979 (LT), BTx2752 (LT), and DKS54-00
- 388 (non-LT) corresponded with prior references (Gholipoor et al., 2010; Shekoofa et al., 2014). A
- 389 PCA (Figure 7) and a negative correlation between g_s and T_c , although non-significant, to some
- 390 extent, suggested T_C as a surrogate phenotype to discriminate differences in transpiration (Figure
- 391 7, Figure S7). The effectiveness of $T_{\rm C}$ in determining variability for LT transpiration has been
- 392 demonstrated by comparing $T_{\rm C}$ with whole-plant transpiration in controlled environments (Belko
- 393 et al., 2013).
- 394 For other genotypes, the g_s phenotype contradicted prior classification (Figure 4). For 395 instance, unexpected high gs for genotypes with the putative LT trait SC803, Tx7000, and 396 BTx642 (Table 1) suggest they had the non-LT trait. Discrepant results were also reported for 397 lentils where a genotype with the putative LT in the greenhouse displayed a non-LT trait in field 398 trials (Guiguitant et al., 2017). These observations imply that growth chamber experiments do 399 not fully represent field conditions. Phenotyping for TR-VPD in growth chambers is limited to 400 early growth stages and environmental conditions are more erratic in field settings. Indeed, prior 401 studies suggested that early growth stages in field conditions (Shekoofa et al., 2014) and 402 temperatures beyond 30 °C (Riar et al., 2015) suppress the expression of the LT trait.
- 403 The phenotype identified with each phenomic approach varied for most genotypes 404 (Figure 4). These differences can be attributed to the sample size utilized in each method (Figure 405 4), variability in canopy architecture, and stand count. The interaction of canopy architecture 406 traits with wind speed influences convective heat transfer and T_C (Gates, 1980; Leigh et al., 407 2017; Melcher et al., 1994). Canopy architecture traits can be more important determinants of T_C 408 than physiological traits (Still et al., 2021; Woods et al., 2018) since canopy architechture traits 409 and wind speed influence convective heat transfer and T_C. Genotypes with an acute leaf angle 410
- likely disrupted air movements through the canopy, leading to high T_{Cirt}, as observed in genotype
- 411 BTx642 (Figure 7A). Soil exposure in plots of genotypes with fewer leaves would have reflected 412 solar energy, contributing to increasing air temperature and $T_{\rm C}$ (Figure 7B). This would explain

413 the unexpectedly high T_{Cirt} in AB19 for genotypes DKS28-00 and SC1345 with the putative non-414 LT trait and low leaf area (Table S6).

415 Microenvironmental conditions created by low plant density and high tillering can

416 override the phenotypic expression of a target trait (Jones, 2007). The phenotype identified via g_s

417 for genotype Tx430 shifted from non-LT (Figure 4A, 4B) to LT (Figure 4C) when the plant

418 density was lower than 13 pl m⁻² (Figure S8A). Although the experiment was well-irrigated, at 419 low plant density, it is possible that soil evaporation surpassed plant transpiration. High tillering

419 fow plant density, it is possible that son evaporation surpassed plant transpiration. Fight therms 420 (Figure S8B) likely exacerbated water demand, promoting water competition among culms

421 (Borrell et al., 2014), leading to low g_s and high T_{Cimg} (Figures 4H and 4I).

422 T_{Cimg}, an HTPP approach, can accelerate breeding selection by bringing higher precision 423 (Deery et al., 2019), if such an approach can capture the nature of the target trait. Nevertheless,

424 H^2 for T_{Cimg} was close to zero and lower than H^2 for g_s in AB19. While the H^2 for g_s (0.16-0.55)

425 in our study aligned with an investigation for cotton (0.16 to 0.44) (Percy et al., 1996), the H^2 for

426 T_{Cimg} ranged from 0.05 to 05, which are lower than the values reported for wheat (0.3 to 0.8)

427 (Anderegg et al., 2021; Deery et al., 2019). Different H^2 for T_{Cimg} (Figure 5) among experiments

428 indicates that this approach is prone to microenvironmental conditions likely caused by the

429 variability of canopy architecture traits (Figure 7 and S7A). Additionally, stand count can

430 override the expression of the TR-VPD (Figure S7B and S7B). Overall, using T_{Cimg} to screen

431 large breeding populations would require considering the number of leaves and stand count as

432 covariate effects.

433 Non-linear function models better represent genetic variability for TR-VPD

Genetic variability identified via different phenomic approaches in this study provides candidate
donor lines to breed sorghum yields for water deficit conditions (Figure 4). Our study has shown
that genetic variability occurs under low and high (<2 kPa) VPD (Figure 6D). Genetic variability

437 for TR-VPD has been proposed as differences in slopes and restriction (breakpoint) in gas

438 exchange (Gholipoor et al., 2010; Shekoofa et al., 2014). In our study, observations indicated

that no genotypes with the non-LT trait increased g_s (Figure S3). Evidence of whether observed

440 data fit a linear or segmented regression was never shown or discussed in prior studies. For

441 instance, observed data for outdoor pots suggested asymptotic patterns for chickpeas with LT

and non-LT traits, but the study reported a breakpoint and LT and linear response (Zaman-Allahet al., 2011).

In our research, stomatal response to VPD fit different regressions (Figure 6), indicating
the breakpoint or asymptotic as the best model for a genotype with non-LT trait. Although

disparities between a breakpoint (Figure 6C) and an asymptotic model (Figure 6D) seem to be

447 negligible, independent data support that at higher VPD, g_s will reach its maximum value and

remain constant (Figure 6A). Our findings align with studies where transpiration under well-watered conditions and high VPD remained constant. A wheat study reported breakpoints

watered conditions and high VPD remained constant. A wheat study reported breakpoints
 response for 100% of commercial varieties tested in growth chambers (Schoppach et al., 2017).

451 For woody and herbaceous plants, transpiration under controlled environments reached a

452 maximum plateau at a VPD of 2.5 kPa (Turner et al., 1984). An asymptotic function better

453 portrays the stomatal response to VPD, as this non-linear model has a biological meaning and is

- 454 utilized to describe photosynthesis, light intensity, and light interception (Archontoulis and
- 455 Miguez, 2015). The correct mathematical representation of TR-VPD has a significant
- 456 implication when modeling the effect of the LT at crop system scales. Simulation studies using
- 457 segmented and nonlinear functions are most likely overestimating the impact of the LT trait on458 harvested yield.

459 The underlying mechanism of TR-VPD

- 460 The underlying physiological mechanism of TR-VPD remains enigmatic. The LT trait has been
- associated with aquaporin inhibition (Maurel et al., 2016) and low hydraulic conductivity
- 462 (Choudhary et al., 2013b) that causes partial stomatal closure. In arabidopsis and angiosperms,
- stomatal response to VPD is controlled by hydropassive and hydroactive processes (Merilo et al.,
- 464 2018; Wang et al., 2001). Nevertheless, studies in soybean and peanut claim that LT is
- 465 modulated via a hydropassive mechanism (Sinclair et al., 2017). While the hydropassive
- 466 response involves changes in the turgor cell, the hydroactive mechanism is modulated via
- 467 metabolic signaling (Merilo et al., 2018; Wang et al., 2001). Studies reported leaf-derived ABA
- 468 when plants are exposed to high VPD (Hu et al., 2016; Jalakas et al., 2021), although findings
- 469 indicate that *Open Stomata 1 (OST1)* kinase rather than ABA controls stomatal response to
- 470 humidity (Merilo et al., 2018). Otherwise, differences in leaf hydraulic conductivity reported for
- 471 genotypes SC1205 (dwarf, non-LT) and SC15 (tall, putative LT) (Choudhary et al., 2013b;
- 472 Ocheltree et al., 2013) can result from contrasting hydraulic anatomy (Du et al., 2020) and plant
- 473 architecture. Indeed, under well-water conditions, a study reported differences in hydraulic
- 474 anatomy (xylem diameter) for a short (ICSSH58, 1.5 m) and tall (ICSV25280, 1.7 m) sorghum
 475 (Guha et al., 2018).
- 476 Discrepancies in TR-VPD between non-LT and LT traits are hypothetically due to 477 stomatal openness (Sinclair et al., 2017) with expected open stomata for the genotype with the
- 478 non-LT trait and partial stomatal closure for a genotype with the LT trait (Figure 8A).
- 479 Determining whether partial stomatal closure is controlled by hydropassive or hydroactive
- 480 mechanisms requires testing the expression of metabolic signals (Figure 8B). Otherwise,
- 481 asymptotic patterns of TR-VPD (Figure 6D) suggest that the stomata remain open at high VPD
- 482 for genotypes with LT and non-LT traits (Figure 8C). Then, differences in gas exchange are
- 483 hypothetically due to variations in hydraulic anatomy traits such as variability in stomatal density
- 484 (Bheemanahalli et al., 2021), xylem diameter, or boundary layer thickness (Figure 8D). From a
- 485 breeding perspective, a hydroactive mechanism would allow identifying known genes such as
- 486 *OST1* or *slow anion channel 1 (SLAC1)*, which are ABA-dependent or -independent (Merilo et
- 487 al., 2018). Meanwhile, the hydropassive mechanism implies that the aquaporin response is
- 488 independent of any signaling pathway and unknown genes. Testing these hypotheses in
- 489 identified lines (DKS54-00, BTX2752, and SC979) can elucidate mechanisms underlying the
- 490 TR-VPD. Further quantifying the carbon assimilation trade-offs of the LT trait in near-isogenic
- 491 lines would indicate the impact of this trait on drought-prone regions. This knowledge can guide
- 492 breeding programs to consider different options when designing a breeding pipeline.

493 *Future steps*

- 494 The results of this study represent TR-VPD when soil water is not limited in the soil profile.
- 495 However, drought-prone regions are subjected to simultaneous high VPD and soil water deficit.
- 496 Results under a rainfed treatment and soil water deficit in a semiarid environment revealed a
- 497 decline in g_s as VPD increases (Figure 6A), suggesting that soil water deficit and high VPD lead
- to stomatal closure to avoid dehydration and physiological damage (Oren et al., 1999). So far,
- the confounding effect of VPD and soil moisture has been ignored in studies of LT. Then it
- 500 would be worthwhile to determine whether the effect of VPD on stomatal closure prevails over
- 501 the impact of soil water deficit on stomatal closure or vice versa. Genotypes with putative LT 502 traits identified in this study correspond to full-season backgrounds. A comparison of yield
- 502 components under well-watered and rainfed treatments in a semiarid environment (GR21)
- 504 demonstrated that early flowering time overrode the effect of the putative LT trait, either in
- 505 irrigated or rainfed treatments (Figure S10). In silico simulations showed that the LT trait can
- 506 increase yields by more than 5% in western regions of the sorghum belt where medium and
- soor increase yields by more than 5% in western regions of the sorghetin cent where incutatin and 507 short-season hybrids are planted. Then it would be needed to identify and introgress the LT trait
- 508 in short and medium-season backgrounds. Developing a mapping population and near-isogenic
- 509 lines with donors SC979, BTx2752 would allow identifying QTLs via T_{Cimg} ; nevertheless,
- 510 confounding effects detailed in this study need to be considered when using this approach.

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518 AUTHOR CONTRIBUTIONS

- 519 G.P.M., S.S.B., T.F., and R.R. contributed to the conception and design of the work. T.F. and
- 520 R.R. collected field experimental data. X.W was in charge of the UAS imaging. R.R. and X.W.
- 521 conducted data analysis, interpretation, and drafting of the article. G.P.M., S.S.B., T.F., R.R.,
- 522 X.W., J.P., A.E.L. contributed to the final manuscript.
- 523

524 **CONFLICT OF INTEREST**

- 525 The authors declare that they have no conflict of interest.
- 526

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- 719

720 FIGURES AND TABLES

721 Table 1. Summary of hypotheses on transpiration response to VPD classification for

sorghum genotypes reported by previous studies. Stomatal conductance (g_s) and canopy

723 temperature (T_{Cirt}) are proxies of transpiration response to VPD evaluated in field experiments

from 2019 to 2021 in Kansas and Colorado (see Table S1). Information on canopy temperature

was not collected in AB21.

726

Maturity	Inferred VPD response			$\mathbf{g}_{\mathbf{s}}$		T _{Cirt}			
	Growth chambers		Field	(years)			(years)		
	31 °C	37 °C		19	20	21	19	20	21
Med	LT ^a	non-LT ^d	LT ^e	+	+		+	+	
Med	LT ^a	non-LT ^d	-		+	+	+	+	+
Late	LT ^a	non-LT ^d	LT	+	+	+	+	+	+
Late	LT ^{ab}	non-LT ^d	-		+		+	+	
Late	LT ^a	non-LT ^d	LT ^e		+			+	
Late	LT ^a	LT ^d	-		+		+	+	
Med	LT ^b	-	-		+			+	
Early	non-LT ^a	-	-				+		
Late	non-LT ^a	-	-		+	+	+	+	+
Early	non-LT ^a	-	-				+		
Late	non-LT ^a	-	non-LT ^e	+	+	+	+	+	+
Late	NN	NN	NN	+	+	+	+	+	+
Med	NN	NN	NN			+			+
	Med Med Late Late Late Late Med Early Late <i>Early</i> Late <i>Late</i> Late	MaturityGrowth cl31 °CMedLatenon-LT aLatenon-LT aLateLateNN	MaturityGrowth chambersGrowth chambers31 °C37 °CMedLT anon-LT dMedLT anon-LT dLateLT abnon-LT dLateLT anon-LT dLateLT anon-LT dLateLT anon-LT dLateLT anon-LT dLateIT anon-LT dLateIT b-Earlynon-LT a-Latenon-LT a-Latenon-LT a-Latenon-LT a-LateNon-LT a-LateNNNN	MaturityGrowth chambersField31 °C37 °CFieldMedLT anon-LT dLTeMedLT anon-LT d-LateLT anon-LT dLTLateLT abnon-LT d-LateLT anon-LT d-LateLT bnon-LT d-LateLT abnon-LT d-LateLT abnon-LT d-LateLT abnon-LT d-LateNon-LT abEarlynon-LT abLatenon-LT abLatenon-LT abLatenon-LT abLateNNNNNN	MaturityGrowth chambersField31 °C37 °C19MedLT anon-LT dLTe+MedLT anon-LT d-+LateLT anon-LT dLT+LateLT anon-LT dLT+LateLT anon-LT d-+LateLT bnon-LT d-+LateLT anon-LT d-+LateLT anon-LT dLateIT bEarlynon-LT aLatenon-LT aLatenon-LT a-+-LateNNNNNN+	MaturityGrowth chambersField(years) $31 \circ C$ $37 \circ C$ 1920Med LT^a non- LT^d LT^e ++Med LT^a non- LT^d -+Med LT^a non- LT^d -+Late LT^{ab} non- LT^d LT^e +Late LT^a non- LT^d -+Late LT^a $non-LT^d$ -+Late LT^a $non-LT^d$ -+Med LT^b +Late $non-LT^a$ +Late $non-LT^a$ +Late $non-LT^a$ +Late $non-LT^a$ +Late NN NN NN ++	MaturityGrowth chambersField(years) $31 \circ C$ $37 \circ C$ 192021Med LT^a non- LT^d LT^e ++Med LT^a non- LT^d -++Late LT^{ab} non- LT^d LT^e ++Late LT^a non- LT^d -++Late LT^a non- LT^d -++Late LT^a $non-LT^d$ -++Med LT^b ++Late LT^b ++Earlynon- LT^a ++Latenon- LT^a ++Latenon- LT^a ++Late NN NN NN +++	MaturityGrowth chambersField(years)(31 °C37 °C19202119MedLT anon-LT dLTe+++MedLT anon-LT d-+++LateLT anon-LT dLT+++LateLT anon-LT dLT e+++LateLT anon-LT dLT e+++LateLT anon-LT d-+++LateLT b+++Earlynon-LT a+++Latenon-LT a+++Latenon-LT a+++LateNNNNNN++++	MaturityGrowth chambersField(years) $31 \circ C$ $37 \circ C$ 19 20 21 19 20 Med LT^a non- LT^d LT^e $+$ $+$ $+$ $+$ $+$ Med LT^a non- LT^d $ +$ $+$ $+$ $+$ $+$ Late LT^a non- LT^d LT $+$ $+$ $+$ $+$ $+$ Late LT^{ab} non- LT^d LT^e $+$ $+$ $+$ $+$ Late LT^a LT^d $ +$ $+$ $+$ $+$ Med LT^b $ +$ $+$ $+$ $+$ Late IT^a $100-LT^a$ $ +$ $+$ $+$ Latenon- LT^a $ +$ $+$ $+$ $+$ Late NN NN NN $+$ $+$ $+$ $+$ $+$ $+$

727 NN not known (Commercial checks)

728 - not available.

729 + genotypes evaluated in each experiment.

730 Germplasm characteristics: ¹ Has a Tx430 recombinant inbred line (RIL) family, ²Elite genotype (G2P

731 Bridge female tester), ³Nested association mapping parent (NAM) Parent, and ⁴Commercial hybrids.

References: ^a (Gholipoor et al., 2010), ^b (Choudhary et al., 2013a), ^d (Riar et al., 2015), and ^e (Shekoofa et al., 2014).

735 **Table 2. Regression parameters for linear, breakpoint and non-linear stomatal**

736 conductance (gs) response to VPD for five genotypes. Information represents data collected in

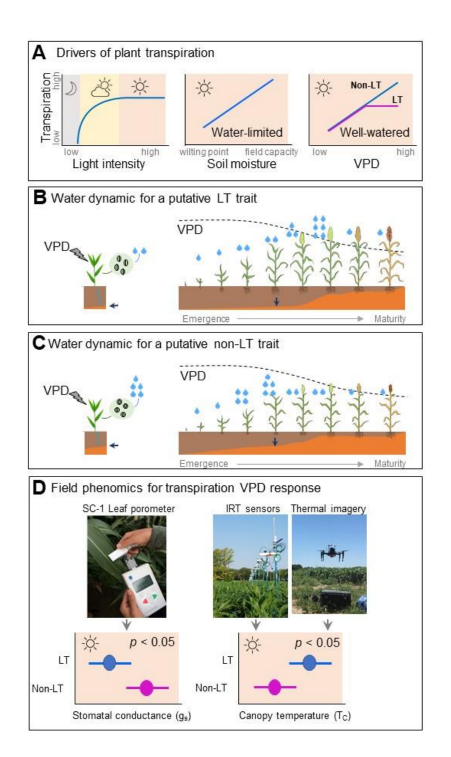
737 AB19, AB20, AB21, and GR21. 84G62:176 observations, DKS54-00: 173 observations, Tx430:

133 observations, Btx2752: 175 observations, and Sc979: 128 observations. RSE: residual

standard error.

740

Genotype	intercept	slope			R ²	RSE
Linear	mol m ⁻² s ⁻¹	mol m ⁻² s ⁻¹ kPa ⁻¹				
84G62 (NN)	0.70 ± 0.04	0.05 ± 0.01			0.03	0.23
DKS54-00 (non-LT)	0.80 ± 0.04	0.02 ± 0.02			0.00	0.27
Tx430 (non-LT)	0.98 ± 0.06	-0.08 ± 0.02			0.00	0.24
BTx2752 (LT)	0.64 ± 0.04	0.05 ± 0.02			0.04	0.24
SC979 (LT)	0.80 ± 0.04	0.02 ± 0.02			0.00	0.25
	intercept	slope1	slope2	BP	\mathbf{R}^2	RSE
Breakpoint	mol $m^{-2} s^{-1}$	mol m ⁻² s ⁻¹ kPa ⁻¹	mol m ⁻² s ⁻¹ kPa ⁻¹	kPa		
84G62 (NN)	0.37 ± 0.07	0.26 ± 0.04	$\textbf{-0.06} \pm 0.04$	2.08 ± 0.18	0.16	0.21
DKS54-00 (non-LT)	0.49 ± 0.08	0.20 ± 0.05	$\textbf{-0.07} \pm 0.03$	2.10 ± 0.20	0.09	0.22
TX430 (non-LT)	-0.057 ± 1.42	0.76 ± 1.21	-0.12 ± 0.29	1.29 ± 0.20	0.11	0.24
BTx2752 (LT)	0.47 ± 0.04	0.16 ± 0.09	0.00 ± 0.02	1.99 ± 0.45	0.06	0.21
SC979 (LT)	0.46 ± 0.16	0.13 ± 0.09	0.01 ± 0.04	2.07 ± 0.56	0.01	0.21
Asymptotic	plateau mol m ⁻² s ⁻¹	inflexion point mol m ⁻² s ⁻¹ kPa-1	start			RSE
84G62 (NN)	0.89 ± 0.02	1.47 ± 0.40	-0.00 ± 0.36			0.22
DKS54-00 (non-LT)	0.89 ± 0.02	1.96 ± 0.68	-0.09 ± 0.60			0.23
Tx430 (non-LT)	-	-	-			-
BTx2752 (LT)	0.81 ± 0.03	1.52 ± 0.81	0.03 ± 0.62			0.21
SC979 (LT)	0.48 ± 0.06	0.69 ± 1.86	0.48 ± 0.62			0.24



743 Figure 1. Drivers of plant transpiration, the hypothetical effect for transpiration response

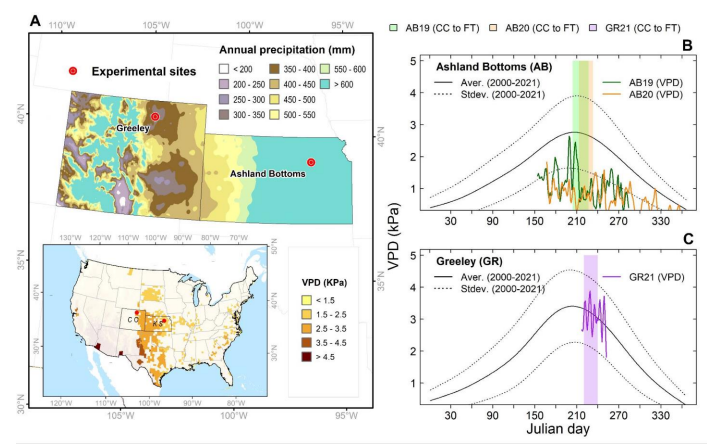
- 744 to VPD traits during a growing season, and predictions of field phenotyping for
- 745 **transpiration response to VPD traits**. A) Transpiration responses to light intensity represented
- 746 with an asymptotic function (Q), transpiration response to soil moisture represented with an
- 747 linear function (SW), hypothesis for the transpiration response to VPD for a putative non-LT
- (linear) and LT trait (breakpoint). Light intensity regulates stomatal response at night (2), from

- 749 sunrise to noon (答), or during any overcast period or day (答). LT, as a response to light
- saturation occurs around noon (*). In this period, transpiration is modulated by SW and VPD in
- 751 water-limited and well-watered conditions, respectively. B and C) Hypothesis of the effect of the
- putative LT and non-LT trait in water dynamics during the growing season. Leaves of a genotype
- with a putative non-LT trait are insensitive to high VPD (low humidity) and keep the stomata
- open. Leaves of a genotype with putative LT trait perceive a high VPD (low humidity), causing
- partial stomatal closure. Blue arrows indicate the effect of the LT on soil moisture during the
- 756 growing season. D) Phenomic approaches to discriminate TR-VPD under well-watered
- conditions phenotyped via stomatal conductance (g_s) and canopy temperature $(T_{Cirt} \text{ and } T_{Cimg})$.
- 758

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760

761 Figure 2. Study system to dissect transpiration response to VPD in sorghum germplasm. A)

762 Spatial variability of annual precipitation and maximum VPD in Kansas and Colorado.

763 Precipitation and VPDe deficit information for sorghum-producing areas were acquired from the

764 PRISM Climate Group (<u>https://prism.oregonstate.edu/</u>). B and C) The annual trajectory of daily

765 maximum VPD from 2000 to 2021 and daily maximum VPD for field experiments in Ashland

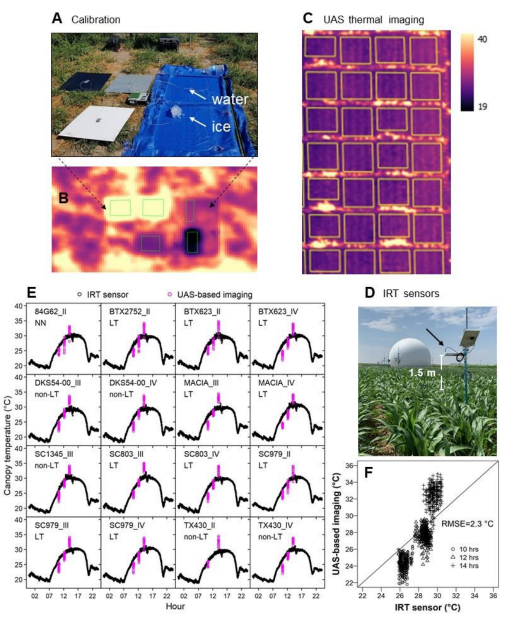
Bottoms (AB19, AB20) and Greeley (GR21). Maximum daily VPD at each site were acquired

767 from the PRISM Climate Group (<u>https://prism.oregonstate.edu/</u>). Maximum VPD for each

recorded with the ATMOS41 weather station. Information of AB21 not included

- in Figure 2B.
- 770
- 771

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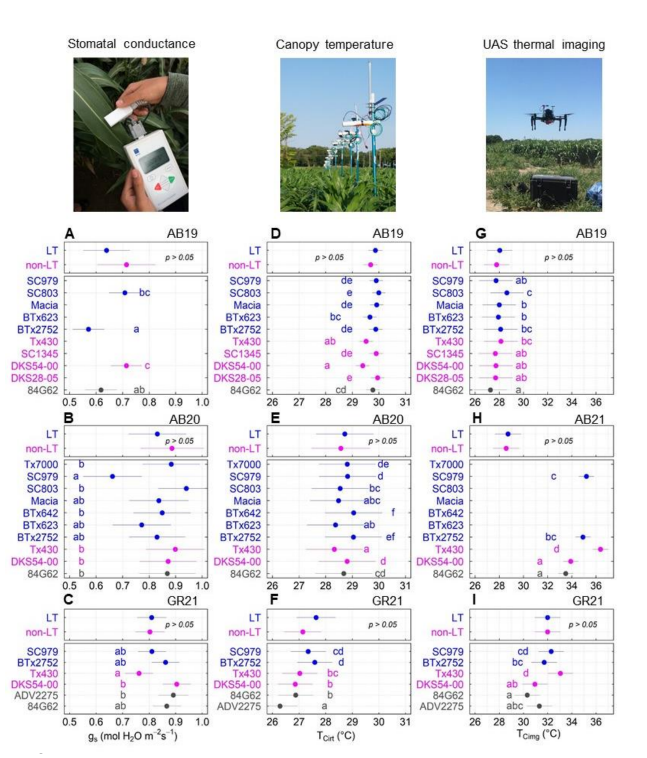
773 Figure 3. Approach for comparison of sources for canopy temperature (T_c) in AB19. A)

774 Multiple ground temperature reference panels, including stainless steel boards painted in white,

gray, and black, flat square containers filled with water and mixture of ice and water in AB19. B and C) Thermal image reflecting estimated T_C in AB19. D) IRT sensors in GR21. Each sensor

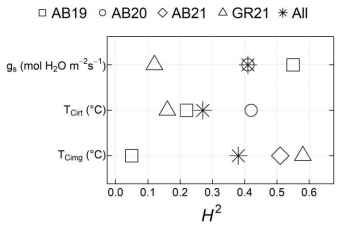
- and C) Thermal image reflecting estimated T_C in AB19. D) IRT sensors in GR21. Each sensor was located on top of the canopy at a 1.5-meter distance. E) Trajectory of T_C obtained with IRT
- 778 sensors (black symbol) and UAS-based imaging (magenta symbol) in 16 sorghum plots in
- Ashland Bottoms on July 26, 2019 (AB19). Black symbols represent T_{Cirt} every minute during
- 780 24 hours, and magenta symbols represent T_{Cimg} every two seconds during a flight time of 12
- 781 minutes. UAS-based imaging was collected in motion at 35 meters above ground level. F)
- 782 Comparison of T_C obtained with IRT sensors and UAS-based imaging in 16 sorghum plots at
- 783 10:00, 12:00, and 14:00 hours in Ashland Bottoms on July 26, 2019 (AB19).

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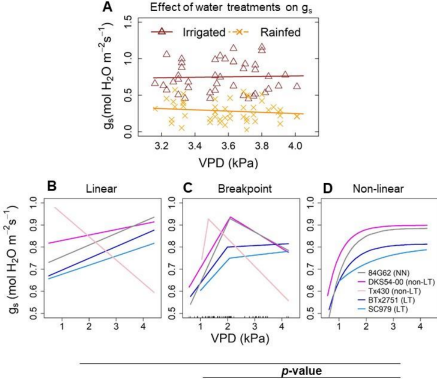


Figüre 4. Field Phenomic approaches to discriminate variability in transpiration response to VPD. A, B, a786C) Variability for stomatal conductance (g_s) D, E, and F) Canopy temperature from IRT sensors (T_{Ci7})7n hours with high VPD (12:00 to 16:00 hours). H, I and J) Canopy temperature from thermal imagery (T_{Ci7})87 Colors represent the reported putative TR-VPD (Table 1): not known (NN, gray), non-limited transfer ation (non-LT, blue) and limited transpiration (LT, magenta). Error lines indicate the standard error. Lett**290** indicate significant differences ($\alpha < 0.05$) of all pairwise comparisons using the Sidak test. g_s

repr**29d**nts individual leaves in the sunlit canopy, T_{Cirt} represents the 0.5 m² diameter of the sunlit canopy, whi**I** Θ **T**_{Cimg} represents the median T_C at plot level (3 m²)



- Figure 5. The broad sense of heritability (H^2) for different phenomic approaches to
- 795 **discriminate variability in transpiration response to VPD.** Stomatal conductance (g_s),
- canopy temperature from IRT sensors (T_{Cirt}), and canopy temperature from thermal imagery (T_{Cimg}).



	<i>p</i> -value					
Genotype (trait)	linear	breakpoint	Non-linea			
84G62 (NN)	ns	***	***			
DKS54-00(non-LT)	ns	***	**			
Tx430 (non-LT)	ns	ns	<i><i><i>u</i></i></i>			
BTx2752 (LT)	ns		ns			
SC979(LT)	ns	ns	ns			

799

800 Figure 6. Stomatal response to VPD for sorghum germplasm evaluated over three years of

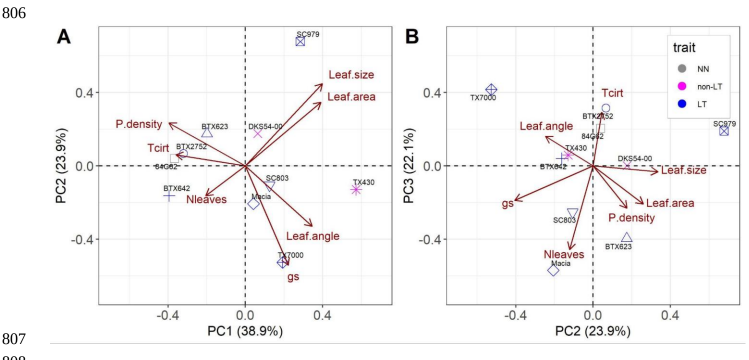
801 **field experiment**. A) Comparison of stomatal conductance (g_s) to VPD under irrigated and

rainfed treatments in GR21. The information comprises six genotypes evaluated for their

stomatal response to VPD in GR21. B, C and D) variability of stomatal response to VPD for

sorghum germplasm represented via linear, breakpoint and non-linear regression models. Each

805 line was fitted with observation indicated in Figure S2.



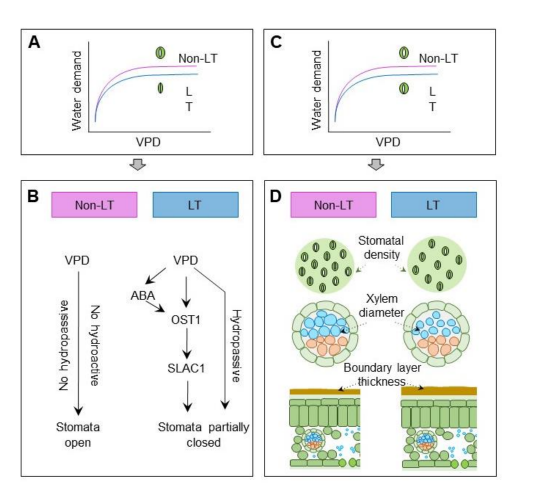
809 Figure 7. Principal component analysis (PCA) for genotypes characterized for their TR-

VPD and canopy architecture traits in AB20. P.density: plant density (plants m⁻²); Leaf.angle: 810

leaf angle (°); Leaf.area: leaf area per plant (leaf area plant⁻¹); Leaf.size : maximum leaf size 811

812 (cm²); N.leaf: maximum number of leaves. Details on each canopy architecture trait is provided

813 in Table S4 and Figures S3, S5, and S6.



815 Figure 8. Hypotheses on mechanisms underlying variation in transpiration VPD response.

A) Water demand varies due to stomatal response, where the genotype with putative non-LT
maintains open stomata, and the genotype with LT trait partially closes the stomata. B) Both

818 hydroactive and hydropassive mechanisms can cause stomatal closure, but hormonal signaling is

819 involved in the hydroactive mechanism. C) Genotypes with non-LT (pink line) and LT (blue)

820 maintain stomata open, and water demand varies due to differences in hydraulic anatomy. D)

821 Expected hydraulic anatomic traits for genotypes with putative non-LT and LT traits. The

822 hydropassive and hydroactive mechanism for the LT trait was adapted from Merilo et al. (2018).

823