# **High water use efficiency due to maintenance of photosynthetic capacity in sorghum under water stress**

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#### **Highlight**

JORIGATE THE STORIST INTERNATION THE CONSTRAINT THE CONTRAINT CONTRAINT THE CONTRAINST THAT CONTRAINST THAT CHANNEL THAT CHANNEL THAT CHANNEL THAT CHANNEL THAT THAT CHANNEL THAT THAT CHANNEL THAT THAT THAT THAT THAT THAT T High water use efficiency is linked to maintenance of photosynthesis independently of leaf hydraulic conductance in sorghum lines with different aquaporin-associated haplotypes.

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#### **ABSTRACT**

vulnerable agricultural systems with higher tolerance to water stress (WS) that<br>widely planted crops. To investigate physiological controls on *WUE* and its inherit<br>sorghum we screened 89 genotypes selected based on inheri Environmental change requires more crop production per water use to meet the rising global food demands. However, improving crop intrinsic water use efficiency (*iWUE*) usually comes at the expense of carbon assimilation. Sorghum is a key crop in many vulnerable agricultural systems with higher tolerance to water stress (WS) than most widely planted crops. To investigate physiological controls on *iWUE* and its inheritance in sorghum we screened 89 genotypes selected based on inherited haplotypes from an elite or five exotics lines, containing a mix of geographical origins and dry vs. milder climates, which included different aquaporin (AQP) alleles. We found significant variation among key highly heritable gas exchange and hydraulic traits, with some being significantly affected by variation in haplotypes among parental lines. Plants with a higher proportion of the non-stomatal component of *iWUE* still maintained *iWUE* under WS by maintaining photosynthetic capacity, independently of reduction in leaf hydraulic conductance. Haplotypes associated with two AQPs (SbPIP1.1 and SbTIP3.2) influenced *iWUE* and related traits. These findings expand the range of traits that bridge the trade-off between *iWUE* and productivity in C<sub>4</sub> crops, and provide possible genetic regions that can be targeted for breeding.

**Keywords**: C<sup>4</sup> Crops; water use efficiency; stomatal conductance; water stress; sorghum; hydraulic conductance; genotypic variation.

#### **INTRODUCTION**

Food security amid water scarcity is one of the key global challenges of the 21<sup>st</sup> century (UNCTAD, 2011). Sorghum (*Sorghum bicolor*) is globally important for fuel, fibre, food (Borrell *et al.*, 2014*b*), and animal feed (George-Jaeggli *et al.*, 2017). Sorghum, a C<sup>4</sup> species, was first domesticated in Africa, where it remains a key staple crop in the arid and semi-arid areas of sub-Saharan Africa, a region experiencing a rapid rise in population (Dillon *et al.*, 2007; Borrell *et al.*, 2014*a*). Such environments are heavily dependent on rainfall, which are expected to show more erratic patterns with climate change (Rippke *et al.*, 2016). With intensifying water scarcity, more attention is being paid to crop productivity per unit of transpired water (Passioura, 2006). This characteristic is termed transpiration efficiency or water use efficiency (*WUE*) (Passioura, 1977). At the leaf level, the physiological control of *WUE* is quantified as the ratio of leaf carbon assimilation (*An*) to stomatal conductance to water vapour (*gs*), and termed intrinsic water use efficiency (*iWUE*).

species, was first domesticated in Africa, where it remains a key staple crop in t<br>and semi-arid areas of sub-Saharan Africa, a region experiencing a rapid<br>population (Dillon *et al.*, 2007; Borrell *et al.*, 2014a). Such Selecting for higher *iWUE* in breeding programs of C<sub>4</sub> crops has been challenging for a number of reasons. First, *iWUE* is a complex trait with multiple physiological components contributing to the variations in *A<sup>n</sup>* and *g<sup>s</sup>* (Condon *et al.*, 2004). Secondly, the potential lack of heritable *WUE*-related traits that can be easily screened (Hammer *et al.*, 1997). Proxies for *iWUE* in C<sub>3</sub> crops such as carbon isotope discrimination are not easily applicable in C<sup>4</sup> counterparts (Condon and Richards, 1992; Henderson *et al.*, 1998; Rebetzke *et al.*, 2002; von Caemmerer *et al.*, 2014; Ellsworth and Cousins, 2016; Ellsworth *et al.*, 2020). Hence, finding genetic variation in *iWUE* among C<sup>4</sup> crops has mainly depended on gas-exchange parameters (Xin *et al.*, 2009). Consequently, improving *iWUE* in C<sup>4</sup> crops requires a better understanding of the mechanisms leading to genetic variation in gas exchange and *iWUE* (Jackson *et al.*, 2016).

Achieving higher *iWUE* can come at the expense of photosynthesis and biomass production (Martin *et al.*, 1999; Condon *et al.*, 2004; Passioura, 2006; Blum, 2009). This is because increases in *iWUE* may result from restricting water use via stomatal closure, which usually occurs during water stress (WS). But if the leaf can still maintain high photosynthesis rate at lower intercellular [CO2] (*Ci*) due to stomatal closure, then *iWUE* increases can benefit biomass production when water is scarce and allow water to remain in the soil for later phenological stages (Sinclair *et al.*, 2005; Vadez, 2019; Srivastava *et al.*, 2024). Still, higher *g<sup>s</sup>* and water use associated to high photosynthesis has led to higher yields in a number of crops under both WS (Blum *et al.*, 1982; Sanguineti *et al.*, 1999; Araus *et al.*, 2003; Vijayaraghavareddy *et al.*, 2020; Ouyang *et al.*, 2022; de Oliveira *et al.*, 2023) and well-watered (Reynolds *et al.*, 1994; Fischer *et al.*, 1998; Horie *et al.*, 2006) conditions. Therefore, a key challenge is to understand how to screen for greater *iWUE* without sacrificing greater productivity, especially under WS (Leakey *et al.*, 2019; de Oliveira *et al.*, 2022, 2023).

For the U.S. Therefore, increase that, 1994; Fischer et al., 1998; Horie et al., 2023) and well-watered (Reynolds et al., 1994; Fischer et al., 1998; Horie 2006) conditions. Therefore, a key challenge is to understand how *iWUE* depends on the *An*-*g<sup>s</sup>* relationship, which is almost linear at low to moderate *gs*, and reaches a plateau at high *g<sup>s</sup>* (Wong *et al.*, 1979; Gilbert *et al.*, 2011). Consequently, *A<sup>n</sup>* and *g<sup>s</sup>* contribute different proportions to *iWUE* depending on their operational position along the *An*-*g<sup>s</sup>* curve (Ghannoum, 2016). When comparing different plants, high *iWUE* may be due to higher *An*, and/or lower *g<sup>s</sup>* (Leakey *et al.*, 2019). The operation of the CO<sup>2</sup> concentrating mechanism (CCM) in  $C_4$  leaves leads to the saturation of  $A_n$  at lower  $C_i$ than in C<sup>3</sup> plants, and hence low *gs*, which means that operating with high *g<sup>s</sup>* may lose water without improving *An* (Srivastava *et al.*, 2024). On the other hand, some crop varieties can sustain high *iWUE* due to higher photosynthetic capacity per given *Ci*, and (Gilbert *et al.*, 2011) proposed a method to screen for variation in *iWUE* associated with stomatal or non-stomatal components of *A*<sup>n</sup> applied to soybean (C<sup>3</sup> dicot) and later applied by Li *et al.* (2017) in sugarcane (C<sup>4</sup> monocot). Finding such varieties is agronomically beneficial as it would alleviate the often-negative relationship between *iWUE* and photosynthesis or productivity.

The contribution of plant or leaf water status to WS responses can also be an important determinant of the trade-off between *iWUE* and photosynthesis. During WS, the hydraulic flux of water from the soil to the sites of transpiration within the leaves is often reduced, leading to a decrease in plant (*Kplant*) and leaf (*Kleaf*) hydraulic conductances. Consequently, leaves close stomata to maintain cell turgor and metabolism, and to reduce the risk of catastrophic hydraulic failure (Meinzer and Grantz, 1990; Mott and Franks, 2001; Meinzer, 2002; Brodribb *et al.*, 2003), which also reduces CO<sub>2</sub> supply for

photosynthesis. One byproduct of selecting for high *iWUE* under WS is obtaining varieties that favour water conservation in the soil, sometimes at the cost of photosynthesis (Choudhary *et al.*, 2013; Choudhary and Sinclair, 2014). This strategy often selects varieties with low *Kleaf* or that reduce *Kleaf* significantly during WS and specially at high VPD. However, lower *Kleaf* negatively impacts photosynthesis either directly, or indirectly via reducing *g<sup>s</sup>* and hence *Ci*. Hence, screening for variation in hydraulic responses to WS can identify varieties that maintain *A<sup>n</sup>* despite reduced *C<sup>i</sup>* under WS, attaining higher *iWUE*.

via reducing  $g_s$  and hence  $C_s$ . Hence, screening for variation in hydraulic responsive via reducing  $g_s$  and hence  $C_s$ . Hence, screening for variation in hydraulic responsive WS can identify varieties that maintain  $A_n$ A possible target that link photosynthesis, water relations and *iWUE* are aquaporins (Vadez *et al.*, 2014; Reddy *et al.*, 2015). Aquaporins (AQPs) are channel proteins embedded in the lipid bilayer of plant cellular membranes. AQPs strongly influence the flow of water and ions within the leaf, affecting physiological parameters such as *Kleaf* and *iWUE* (Maurel *et al.*, 2015), including in sorghum (Choudhary et al., 2013; Hasan et al., 2015, 2017; Liu et al., 2015; Zhang et al., 2019). More importantly, several AQPs in plants have been shown to be key CO<sub>2</sub> transporters (sometimes called cooporins) especially across the plasma membrane (Groszmann *et al.*, 2017). Hence, they could hypothetically increase CO<sup>2</sup> or H2O supply to the sites of carboxylation without increasing *gs*.

Screening for variation in physiological traits is laborious and time-consuming, and requires an extensive number of genotypes. We explored the rich genetic resources that are available for sorghum (Mace *et al.*, 2019), using variations in genomic regions associated with different AQP alleles (haplotypes) from a sorghum NAM population (see Methods). We curated more than 80 genotypes and grew them under two watering regimes to assess the degree of variation of *iWUE* and other plant traits in closely similar sorghum genotypes under WS and to use that variation to test the following hypotheses: 1) partitioning the stomatal and non-stomatal components of *iWUE* within this diversity will reveal genotypes that achieve high *iWUE* under WS by maintaining photosynthesis; 2) achieving high *iWUE* under WS due to maintenance of photosynthesis will be underpinned by maintenance of *Kleaf* and leaf water status; and finally 4) the maintenance of hydraulics and photosynthesis under WS will be linked to certain AQPs and their related haplotypes.

#### **MATERIALS & METHODS**

#### **Genotype selection**

allows for statistical robustness while retaining diversity of parental lines. NAM masone allelic diversity by breeding (and *backcrossing*) recombinant inbred lines from multiple parents with a single parent as a referenc The genotypes used in this study are a part of a Nested Association Mapping (NAM) population (Jordan *et al.*, 2011; Tao *et al.*, 2020). NAM is a type of selective breeding that allows for statistical robustness while retaining diversity of parental lines. NAM maintains some allelic diversity by breeding (and *backcrossing*) recombinant inbred lines (RILs) from multiple parents with a single parent as a reference line (**Fig. 1**). Hence, the progeny shares most of their genetic material, and phenotypic differences can be quickly linked to specific genetic regions. Genotypes used in our study came from a Sorghum NAM population that comprises an elite parental line R937945-2-2 (Recurrent Parent, RP) crossed with >100 exotic lines with geographical or racial diversity (Non-Recurrent Parent, NRP). The F1 progeny were backcrossed with the elite parent to produce  $BC_1F_1$ populations. BC<sub>1</sub>F<sub>1</sub> genotypes compromise  $\sim$  22-25% exotic (NRP) line genome with the rest being RP background (**Fig. 1**). Individual BC1F<sup>1</sup> populations are genotyped using high density single-nucleotide polymorphism (SNP) markers providing profiles of the exact exotic chromosomal segments, giving us information on what genes are coded for in the 22-25% NRP portion of the genome, and what genes are coded for in the remaining RP section of the genome. In addition to this resource, whole genome sequencing is available for many of the exotic parental lines and the elite line (Mace *et al.*, 2013).

We screened this sorghum NAM population for genes of 8 AQPs to select lines carrying non-synonymous SNP alleles of those genes. Specifically, the subpopulation was screened to identify individual lines with chromosomal segments harbouring the elite (RP) AQP allele (RP-Haplotype) or the exotic (NRP) AQP allele (NRP-Haplotype) of a specific AQP. The final 89 lines chosen were derived from 5 exotics (NRPs) containing a mix of geographical origins, with specific focus on a mix of dry vs. milder climates with the idea these would have greater extremes in the traits of interest due to necessary adaptations to their climate of origin (**Table 1**). This approach allowed us to create subpopulations within the 89 genotypes through focusing on one of the 8 AQPs, with each subpopulation containing two sets of genotypes, a set (>5) of genotypes containing the RP-haplotype for that AQP, and a set containing the NRP-haploype. Hence, any phenotypic difference

when comparing RP or NRP haplotypes associated with a certain AQP may be due to the specific AQP allele that characterize the RP or NRP haplotype or from the accompanying genes from that chromosomal segment (haplotype), creating a link between phenotype and genotype.

#### **Plant culture**

**Plant culture**<br>8 litre cylindrical pots were used to allow ample space for root development<br>implementation of the water stress treatment. The pots were adjusted to similar<br>(1.5 kg) by adding gravel (100-300 mm diameter), 8 litre cylindrical pots were used to allow ample space for root development before implementation of the water stress treatment. The pots were adjusted to similar weight (1.5 kg) by adding gravel (100-300 mm diameter), then the same amount of soil was added to all pots. Fly screen mesh (aluminium insect screen) was added to the bottom of the pots to minimize soil seeping through pot drainage holes. The potting mix was made of soil, sand and decomposed bark. It has large particle size for good drainage and root development. Granulated fertiliser (Osmocote Plus Organics All Purpose Fertiliser, Scotts Miracle-Gro Company, Marysville, Ohio) was pre-mixed with the soil, with more fertiliser added in the lower half of the pot where more roots will develop as the plant grows. To each pot, 3.5 kg of soil was added, making the total pot weight 5 kg. We left 2-3 cm at the op of the pot empty, making the volume of the soil filled 7.5 litres.

Seeds were directly sown into the upper soil layer in October 2019. Plants germinated and grew in a naturally lit, controlled-environment greenhouse (Plexiglas Alltop SDP 16; Evonik Performance Materials, Darmstadt, Germany) at the Hawkesbury Institute for the Environment, Western Sydney University, Richmond, New South Wales, Australia (- 33.612032, 150.749098). The ambient temperature was set at 30°C during the day period, with night temperatures set at 18°C. There was a 2 h period at 24°C between the temperature transitions. The day temperature started at 8 am, and night temperature at 8 pm, when sunrise was about 5 to 6 am and sunset at 7-8 pm, reaching midday maximums of 34-35°C and midday relative humidity of 40-50 %. (**Supplementary Fig. S1**). CO<sub>2</sub> concentration was kept at ambient levels. Due to the large number of plants, we needed three identical and adjacent greenhouse chambers (8 m long x 3 m wide x 5 m tall), which contained both well-watered and water stressed pots, and pots were swapped between the three chambers fortnightly during growth in a randomised fashion. Chamber conditions were monitored via a data logger (Tinytag plus 2, Omni Instruments) hung in the middle of the room at 2 metres height. Light levels were monitored occasionally using a light meter and were 1500 µmol m<sup>-2</sup> s<sup>-1</sup> at midday on sunny days at plant height level at measurement time (~2m from the ground).

#### **Watering Treatments**

All plants were well-watered the first six weeks of growth when half of the planotype were subjected to water stress (WS) and the other half continued und<br>watered conditions. When plants were five weeks old, pots were wei All plants were well-watered the first six weeks of growth when half of the plants by genotype were subjected to water stress (WS) and the other half continued under wellwatered conditions. When plants were five weeks old, pots were weighed in the late evening (*Wevening*), then watered at dusk and weighed again in the morning (*Wmorning*). This allowed pots to drain excess water with minimal loss via evaporation during the night and determine pot weight at field capacity (FC) by the repetition of this routine over three consecutive sunny days and taking the average of *Wmorning*. On each morning we also measured the volumetric soil water content (VSWC) with a sensor (Campbell Scientific, Logan, UT, USA) on each pot after measuring *Wmorning*. FC was 13-15 % for our soil. The difference between pot weight at FC and pot weight before watering in the evening (*Wmorning* – *Wevening*) represented the amount of water transpired by each plant during the day under well-watered conditions. After 6 weeks of growth, watering was withheld from half of the pots (WS, water stress treatment), while the other half continued to be watered at FC (WW, well-watered treatment). Stomatal conductance was monitored in WS plants until it reached around 0.1 mol  $m^{-2} s^{-1}$  or less at saturating light, with the plant also showing signs of wilting. When conductance reached the required level, and signs of wilting appeared, the VSWC was about 5% for most pots. At this point, we measured pot weight as described before to establish the amount of water lost to transpiration by the plants in the WS treatment (about 50 mL). Three folds this amount of water, equivalent to total plant transpiration during the day in the WS treatment for three days, was added every three days to the WS pots. Hence, plants under WS got just enough water for replacement of water loss via daytime evapotranspiration, and we ensured that water status of WS plants was not influenced by recent watering by the delaying of measurements to the third day after watering.

The two watering regimes were maintained until the end of the experiment, constituting the two treatments: WW – FC; WS – 50 mL every day or 150 mL every 3 days. Impact of WS was visible 2 weeks after water withholding for most genotypes (plants were 8 weeks old). There were three replicates (pots) per genotype and water treatment. Hence, each genotype had 6 pots total, with 3 for each treatment (*n*=3), except for the elite parent R937945-2-2 (the RP) which had 6 pots per treatment (*n*=6).

#### **Time of measurements and sampling**

Plants were sampled between weeks 9 and 12 after germination, when they had 10–12 fully expanded leaves. WS plants were measured at least 3 weeks after the onset of the drought treatment. In total, sampling lasted for about a month (mid-December 2019 to mid-January 2020), which represents the peak of the Australian summer. Priority for physiological sampling was given to plants at the booting stage so that all plants were measured before or at the start of flowering.

#### **Midday leaf gas exchange**

**Time of measurements and sampling**<br>
Plants were sampled between weeks 9 and 12 after germination, when they had<br>
fully expanded leaves. WS plants were measured at least 3 weeks after the onse<br>
drought treatment. In total Midday leaf gas exchange rates were measured between 10 am and 2 pm on sunny days. The photoperiod was 14-15 hours and solar midday was around 1–1:30 pm. A Li-6400XT infra-red gas analyser with a LED light source and an area of 6 cm<sup>2</sup> (LiCor Biosciences, Lincoln, Nebraska, USA) was used to obtain light-saturating rates of  $CO<sub>2</sub>$  assimilation (*An*), stomatal conductance to water vapour (*gs*) and transpiration flux (*E*); cuvette conditions were set at:  $30^{\circ}$ C block temperature, flow rate of 500 µmol m<sup>-2</sup> s<sup>-1</sup>, photosynthetic photon flux density (PPFD) of 2000 µmol m<sup>-2</sup> s<sup>-1</sup> (10% blue light), ambient CO<sup>2</sup> concentration set to 400 ppm using CO<sup>2</sup> cylinder mixer, and relative humidity of 40- 60%. The leaf was inserted into the gas exchange cuvette under those conditions avoiding the midrib and with the entire  $6 \text{ cm}^2$  area of the cuvette filled. The leaf was left to acclimate to those conditions until gas exchange and  $CO<sub>2</sub>$  concentration in the substomatal cavity (intercellular CO2, *Ci*) stabilised. Intrinsic water use efficiency (*iWUE*) was calculated as the ratio of *A<sup>n</sup>* to *gs*. All measurements were taken from the middle of the youngest fully expanded leaf (YFEL) of the plant, corresponding to the 9<sup>th</sup>-12<sup>th</sup> leaf depending on genotype. Ambient light level at the YFEL was around 1500  $\mu$ mol m<sup>-2</sup> s<sup>-1</sup>.

#### **Leaf water potential and hydraulic conductance**

covered with cling wrap and tin foil to prevent transpiration and allow the leaf to equal of the day for at least 6 hours (usually they were covered before gas exchange measure started or the day before and collected at t A leaf adjacent to the gas exchange leaf was used to measure midday leaf water potential (*Ψmidday*) using a Scholander-type pressure chamber (Model 1505D Pressure Chambers, PMS Instrument Company, Albany, Oregon, USA). The leaf below the *Ψmidday* leaf was covered with cling wrap and tin foil to prevent transpiration and allow the leaf to equilibrate for at least 6 hours (usually they were covered before gas exchange measurements started or the day before and collected at the end of the day and taken to the lab). This leaf was then used to estimate midday stem water potential (*Ψstem*). Pre-dawn leaf water potential (*Ψpre-dawn*) was sampled on different leaves before daybreak, usually taking leaves in the lower canopy. In each case, the leaf was cut at the ligule and placed in a plastic bag that was exhaled into before sealing. The bags were stored in ice boxes, then transported from the greenhouse to the lab where leaf water potentials were measured within 1-2 h of excision.

Leaf hydraulic conductance was calculated as shown in (Simonin *et al.*, 2015):

$$
K_{leaf} = \frac{E}{(\Psi_{stem} - \Psi_{midday})}
$$
 (1)

where *E* refers to the leaf transpiration rate at the time of excision, estimated by measuring incident PPFD at the time of leaf excision and then *E* at that PPFD level estimated from light-response curves conducted on the same plant. Soil-to-leaf hydraulic conductance (referred to as plant hydraulic conductance, *Kplant*) was calculated as shown in (Robson *et al.*, 2012):

$$
K_{plant} = \frac{E}{(\Psi_{predawn} - \Psi_{midday})}
$$
 (2)

Leaf hydraulic resistance (*Rleaf*) was calculated as 1 / *Kleaf*. Hydraulic resistance of the rest of the plant (*Rrest*) was calculated as (1/*Kplant* ) – *Rleaf*.

#### **Plant and Leaf Morphology**

Leaf width (*LW*) was measured at the same leaf area where gas exchange measurements were made. Leaf length (*LL*) was also measured. Leaf thickness (*LT*) was measured using a Photosynq Multispec (Photosynq, East Lansing, Michigan, USA). At the end of

distilled water and kept in darkness and at 4°C overnight before measuring ag<br>distilled water and kept in darkness and at 4°C overnight before measuring ag<br>turgid weight. Emally lead discs were placed inside an oven at 68 the experiment and before biomass harvest, plant height (*PH*) and number of leaves (*LN*) of each plant were recorded. In this same leaf and area of leaf we measured gas exchange, we collected three leaf discs of  $0.5 \text{ cm}^2$  each to measured leaf mass per area (*LMA*) and relative water content (*RWC*). First, we placed leaf discs inside Eppendorf tubes in ice to quickly measure fresh weight in a four positions balance, then we added distilled water and kept in darkness and at 4ºC overnight before measuring again the turgid weight. Finally leaf discs were placed inside an oven at 65ºC for 48 hours to measure dry weight. *LMA* was calculated as dry weight / leaf discs area (g m-2 ) and *RWC* as: (fresh weight – dry weight)/(turgid weight – dry weight). Plants were harvested after 95-100 days and total aboveground biomass was separated into panicle and vegetative (i.e. leaves and stem) to dry in an oven at 40°C for 10 days before measuring dry biomass, but we present aboveground biomass in the data below as encompassing panicles and vegetative.

## **Relative chlorophyll content (***SPAD***) & Quantum efficiency of PSII (***ΦPSII***)**

Relative chlorophyll content was estimated by a SPAD meter that is embedded in the Photosynq Multispeq (Kuhlgert *et al.,* 2016). SPAD meters measures absorbance at 650 and 940 nm, and then relative values for chlorophyll content are produced. The Multispec was also used to record *ΦPSII* using a pulse-amplitude fluorometer at ambient light. Measurements were conducted on the same leaf used for gas exchange.

## **Components of** *iWUE*

To partition the relative contribution of *A<sup>n</sup>* and *g<sup>s</sup>* to variation in *iWUE* in our population, the approach of Gilbert *et al.* (2011) was used as modified by Li *et al.* (2017). Briefly, because of the curvilinear relationship between *A<sup>n</sup>* and *gs*, it is expected that *A<sup>n</sup>* and *g<sup>s</sup>* will contribute in different proportions to *iWUE* depending on the position of the genotype along the curve and in respect to the mean population value.

From each measurement of gas exchange, we constructed a curve of *iWUE* vs *g<sup>s</sup>* encompassing all treatments. We then calculated the average *iWUE* of all measurements for each treatment. To get variation in *iWUE* due to *g<sup>s</sup>* (*ΔiWUEgs*), the *iWUE* expected if *iWUE* was calculated from our reference curve (*iWUE* vs. *g<sup>s</sup>* – i.e. constant *An*) and then

and component- was then calculated as the difference between the actual method is stomatal component- was then calculated as the difference between the actual method in the calculated as the difference between the actual *ΔiWUEgs* was expressed as the deviation of the calculated *iWUE* from the population mean of *iWUE* for that treatment. This results in a value that highlights how impactful *g<sup>s</sup>* was in deviating that genotypic *iWUE* from the population mean assuming fixed *A<sup>n</sup>* (negative value for *ΔiWUEgs* would mean *g<sup>s</sup>* increase for that genotype reduced *iWUE* by that level compared to the mean). Variation in *iWUE* due to *A<sup>n</sup>* (*ΔiWUEpc*)-the nonstomatal component- was then calculated as the difference between the actual measured *iWUE* and calculated *iWUE* based on *gs* variation. Basically *ΔiWUEpc* represents the remaining "difference" between the population mean *iWUE* and genotypic *iWUE* that was not covered by *ΔiWUEgs* This means that variation in these two components can highlight how each of *g<sup>s</sup>* and *A<sup>n</sup>* contribute to *iWUE*. For example, for a given genotype, if *ΔiWUEgs* is small but *ΔiWUEpc* is big (both positive), it means that *iWUE* is higher than the population mean because of higher photosynthesis mainly and lower conductance secondarily (see **Supplementary Fig. S2** an illustration). We also compared *ΔiWUEgs* and *ΔiWUEpc* values if taken from a reference curve that is based on a reference genotypes and it showed complete agreement (*R*=0.98, **Supplementary Fig. S3**).

### **Calculating the magnitude of change in** *A<sup>n</sup>* **and** *C<sup>i</sup>* **in response to water stress**

To investigate how genotype response to WS enables the achievement of high *iWUE* by amplifying one of its two components highlighted earlier, we calculate the "degree of change" in a hypothetical *An*-*C<sup>i</sup>* curve based on genotype mean value change between WW and WS (Rowland *et al.*, 2023). This method estimates both the magnitude of the change and the direction of the phenotypic change vector (the angle) between two contrasting environments. The change in the angle, *θ*, represents change in trait covariation, in our case the dependence of *A<sup>n</sup>* on *Ci*. Small changes in *θ* would indicate a large decrease in *C<sup>i</sup>* but a small decrease in *An*, indicating drought resilience, meaning the achievement of higher *iWUE* due to stomatal closure but also maintenance of photosynthesis rates. A large *θ* would indicate a combined plummeting in *A<sup>n</sup>* with *Ci*, meaning that *iWUE* would increase less due to photosynthesis and more due to stomatal closure under WS (see Result for more clarification).

#### **Genetic variation**

Broad-sense heritability was calculated as in Li *et al* ( 2017):

$$
H_b^2 = \frac{\sigma_g^2}{\sigma_p^2} \tag{3}
$$

where  $\sigma_{g}^2$  and  $\sigma_{p}^2$  are the genotypic and phenotypic variances respectively.  $\sigma_{g}^2$  was obtained as the sq. of mean from the ANOVA output.  $\sigma_{p}^{2}$  was calculated as:

$$
\sigma_p^2 = \sigma_g^2 + \frac{\sigma_g^2 \times treatment}{number \space of \space treatments} + \frac{\sigma_e^2}{number \space of \space replicates}
$$
 (4)

where  $\sigma_{g}^2$  and  $\sigma_{p}^2$  are the genotypic and phenotypic variances respectively. o<br>
obtained as the sq. of mean from the ANOVA output.  $\sigma_{p}^2$  was calculated as:<br>  $\sigma_{p}^2 = \sigma_{g}^2 + \frac{\sigma_{g\_streament}^2}{number \space of \space reaches} + \frac{\sigma_{g}^2}{number$ where *σg x treatment<sup>2</sup>* and *σ<sup>e</sup> <sup>2</sup>* are the genotype \* treatment interaction and error variances respectively. *σg x treatment<sup>2</sup>* was obtained as the mean squared of the genotype \* treatment interaction and  $\sigma_e^2$  was obtained as the sq. of mean residual error. Because the heritability analysis encompasses both treatments, the number of replicates was standardized as 5 (as opposed to 6; 3 WW and 3 WS) to account for genotypes not in both treatments. The genotypic coefficient of variation (*GCV*) and the phenotypic coefficient of variation (*PCV*) were calculated as:

$$
GCV = \frac{\sigma_g}{mean} \times 100
$$
 (5)  
PCV =  $\frac{\sigma_p}{mean} \times 100$  (6)

where *σ<sup>g</sup>* and *σ<sup>p</sup>* are the genotypic and phenotypic standard deviation. The *mean* refers to the mean of all the measurements across treatments for the variable in question. For the mean value of *iWUEgs* and *iWUEpc* where averages are near zero or negative (because these values are expressed as deviations from the average of all observations), the value used for *mean* was that for *iWUE*.

#### **Statistical analyses**

Statistical analysis and data visualisation were performed using R software (R Core Team (2020). R: A language and environment for statistical computing. R Foundation for Statistical Computing, Vienna, Austria. URL https://www.R-project.org/). Normality was checked by plotting a generalized linear model and inspecting residual plots. Analysis of

variance (ANOVA) and multiple ANOVA (MANOVA) was carried out using linear mixedeffects models (package nlme), with replicate and genotype as the random variable respectively, and the fixed variables being AQP Haplotype\* Water Treatment \* to get the *P*-value associated with the model (**Fig. 2** and **Table 3** respectively). Variance within groups was performed afterwards using a *post hoc* Tukey test. Regression analysis was carried in R using linear modelling (lm). A Pearson product moment correlation analysis was performed to test statistical significance of relationships at *P*<0.05 and obtain correlation coefficients *R* (which was then converted into *R<sup>2</sup>* ).

#### **RESULTS**

#### *Genotypic variation among key traits*

carried in R using linear modelling (lm). A Pearson product moment correlation<br>carried in R using linear modelling (lm). A Pearson product moment correlation<br>was performed to test statistical significance of relationships Gas exchange variables varied among the genotypes under both watering regimes. We excluded the means for genotype R-05012-1 under WS as it responded very poorly to WS and exhibited mean carbon assimilation rate of 1.79  $\mu$ mol m<sup>-2</sup> s<sup>-1</sup> and stomatal conductance of 0.01 mol H<sub>2</sub>O m<sup>-2</sup> s<sup>-1</sup>, which was extremely low. Mean genotype CO<sub>2</sub> assimilation rate  $(A_n)$  experienced a 2.2-fold variation (17.6–39.3 µmol m<sup>-2</sup> s<sup>-1</sup>) under WW conditions and 6.1-fold variation (6.8—32.0 µmol m<sup>-2</sup> s<sup>-1</sup>) under WS conditions (**Table** 2). Similarly, mean stomatal conductance  $(g_s)$  experienced 2.9- (0.11 to 0.33 mol m<sup>-2</sup> s<sup>-1</sup>) and 6.4-fold variation (0.01 to 0.16 mol  $m<sup>-2</sup>$  s<sup>-1</sup>) under WW and WS conditions, respectively (Table 2). Operational intercellular  $CO<sub>2</sub>$  concentration  $(C<sub>i</sub>)$  was similarly variable (**Table 2**). Intrinsic water use efficiency (*iWUE*) experienced less variation, with fold change of 1.9 and 1.8 under WW (92 to 170  $\mu$ mol CO<sub>2</sub> mol<sup>-1</sup> H<sub>2</sub>O) and WS (121 to 216 µmol CO<sup>2</sup> mol-1 H2O) conditions respectively (**Table 2**).

*A<sup>n</sup>* and *g<sup>s</sup>* had higher *GCV* than *iWUE* and *C<sup>i</sup>* (**Table 2**). All those variables exhibited high *H<sup>b</sup> <sup>2</sup>* of ≥0.7 alongside hydraulic variables such as *Ψmidday*, apart from *C<sup>i</sup>* (*H<sup>b</sup> <sup>2</sup>*=0.66) (**Table 2**). *PCV* was also similarly high (30-50%) for all those variables (**Table 2**), indicating that environmental factors played a role in determining variation. The genotype \* treatment variance was lower than the genotype variance, indicating that most genotypes responded similarly. Final harvest parameters such as aboveground biomass also varied significantly (fold change > 15), and displayed high *GCV* and *PCV* (**Table 2**). Mean values

(with standard error) of all measured variables for every genotype under both conditions are shown in **Supplementary Table S1**.

#### *Influence of AQP-associated haplotypes on leaf intrinsic water use efficiency*

different AQP-associated haplotypes (see Methods). The results of this statistical are presented in **Table 3.** Haplotypes associated with two AQPs, SbPIP1.1 and Sb<br>had a significant impact on a number of key traits. For S We focused on the variation caused by differences between genotype groups with different AQP-associated haplotypes (see Methods). The results of this statistical analysis are presented in **Table 3**. Haplotypes associated with two AQPs, SbPIP1.1 and SbTIP3.2, had a significant impact on a number of key traits. For SbPIP1.1, the RP haplotype was associated with significantly higher *A<sup>n</sup>* and *g<sup>s</sup>* (**Fig. 2A-B**), while the NRP haplotype had higher *iWUE* (including its *g<sup>s</sup>* component *ΔiWUEgs*) (**Fig. 2C-D**), leaf mass per area (*LMA*) (**Fig. 2E**) and total aboveground biomass (**Fig. 2F**). The SbPIP1.1 NRP haplotype also had the highest *ΔiWUEpc* of all haplotypes under WS (**Table S2** and **Fig. S2H**). For SbTIP3.2, The RP haplotype had higher overall *An*, *gs*, *SPAD*, *ΦPSII* (**Fig. 2G, H, J, K** respectively), and higher *Ψmidday* (**Fig. 2L**), especially under WS, while the NRP haplotype of SbTIP3.2 had higher *iWUE* and plant hydraulic resistance excluding the leaf (*Rrest*) (**Fig. 2I** and **Table 3**), without an effect on biomass. RP haplotype of SbTIP3.2 also maintained *Kleaf* under WS (**Table 3**). In summary, a common trade-off was observed between photosynthesis (*An*, *ΦPSII*) and water use (*gs*, *Kleaf*) for both haplotypes. Hence, genes in that chromosomal region (haplotype) likely influence those traits, including the AQP gene.

#### *WS increased iWUE, which was positively associated with aboveground biomass*

Taken together all the genotypes, we observed that *A<sup>n</sup>* and *g<sup>s</sup>* correlated positively as expected (*R<sup>2</sup>*=0.91; *P*<0.0001; **Fig. 3A**), with both correlating negatively with *iWUE* (*R<sup>2</sup>*=0.92; *P*<0.0001; **Fig. 3B**), especially under WS for *A<sup>n</sup>* (*R*=0.61; *P*<0.0001; **Fig. 3C**). *A<sup>n</sup>* correlated positively with *Kleaf* under WS (*R*=0.53; *P*<0.0001; **Fig. 3D**), and *g<sup>s</sup>* increased with higher *Ψmidday* (*R<sup>2</sup>*=0.49; *P*<0.0001; **Fig. 3E**). Subsequently, *iWUE* correlated negatively with *Kleaf* (*R*=0.54; *P*<0.0001; **Fig. 3F**) as well as with more negative *Ψmidday* and increasing *Rrest* and *Rleaf* (**Supplementary Table S2**). Despite this, aboveground biomass was only marginally associated with *A<sup>n</sup>* when considering both WW and WS plants, but positively correlated with *iWUE* within each watering treatment (**Supplementary Table S2**). Overall the aboveground biomass production across all the

genotypes under WS was regulated by a reduction in leaf area under WS (**Supplementary Fig. S4D**), and increasing leaf mass per are (*LMA*) (**Supplementary Table S2C**).

#### *Components of iWUE under both well-watered and water stress conditions*

We separated *iWUE* into a component attributed to the variation in  $A_n (\Delta iWUE$ <br>another attributed to variation in  $g_s (\Delta iWUE_{gs})$  (Supplementary Fig. S2). T<br>components did not correlate with each other (Fig. 4A), but both pos We separated *iWUE* into a component attributed to the variation in *A<sup>n</sup>* (*ΔiWUEpc*) and another attributed to variation in *g<sup>s</sup>* (*ΔiWUEgs*) (**Supplementary Fig. S2**). The two components did not correlate with each other (**Fig. 4A**), but both positively correlated with *iWUE* (*R*=0.45-0.7, *P*<0.0001; **Fig. 4C-D**). *ΔiWUEgs* was significantly higher under WW than *ΔiWUEpc*, while the opposite was true under WS (**Fig. 4B**). Increased *iWUE* associated with *ΔiWUEpc* under WS occurs because photosynthesis decreases less than *C<sup>i</sup>* (lower *θ*, **Fig. 5A**) due to the maintenance of CCM under WS. Indeed, genotypes that increased *ΔiWUEpc* under WS compared to WW had lower *θ* (*R<sup>2</sup>*=0.58; *P*<0.0001; **Fig. 5C**), while genotypes that increased their *ΔiWUEgs* under WS showed a weak association with increasing *θ* (*R*=0.4; *P*<0.05; **Fig. 5B**). Hence, genotypes that maintained photosynthetic capacity under low *C<sup>i</sup>* can combine *iWUE* with photosynthetic performance under WS.

We examined the link between increased *ΔiWUEpc* under WS with a better hydraulic response. No correlation was found between the increase in *ΔiWUEpc* under WS and higher *Kleaf* (**Fig. 6B**). Instead, increasing *ΔiWUEgs* was associated with lower *Kleaf* (*R*=0.43; *P*<0.05; **Fig. 6A**) and more negative *Ψmidday* (**Supplementary Fig. S5C**), but this did not apply to *ΔiWUEpc* (**Supplementary Fig. S5D**).

## **DISCUSSION**

This study screened a large number of sorghum genotypes that shared most of their genetic composition but differed in key gene blocks (haplotypes) that are associated with certain aquaporins (AQPs) genes inherited from the elite or exotic parental lines. This population was used to test for genetic variation in the response of *iWUE* and its components to WS, and their relationship with productivity and plant hydraulics. Our key findings were: 1) there was significant diversity in many variables related with productivity which also presented high broad-sense heritability; 2) some of this diversity is

underpinned by differences in haplotypes associated with some AQPs especially for gas exchange and hydraulic parameters; 3) the non-stomatal component of *iWUE* (*ΔiWUEpc*) was associated with higher *iWUE* under both WW and WS conditions; 4) genotypes with higher *ΔiWUEpc* were not sensitive to low *Kleaf* under WS. We discuss those findings below.

#### **Breeding for high** *iWUE* **and possible impact of SbAQPs**

Breeding for high *iWUE* and possible impact of SbAQPs<br>Breeding for high *iWUE* in C<sub>4</sub> crops, and particularly in sorghum, has been disco<br>due to lack of sufficient variation among genotypes reported in earlier studies,<br>t Breeding for high *iWUE* in C<sup>4</sup> crops, and particularly in sorghum, has been discouraged due to lack of sufficient variation among genotypes reported in earlier studies, lack of traits that could be easily measured in large-scale screens, and the complex physiology of *iWUE,* where its components such as *g<sup>s</sup>* can be easily influenced by environmental factors such as VPD or WS (Condon *et al.*, 2004; Sinclair *et al.*, 2005). High *H<sup>b</sup> <sup>2</sup>*of key parameters such as *An*, *gs*, *iWUE*, *LW* and *SPAD* under environmental variation within inbred sorghum genotypes is a significant finding (**Table 2** – see similar high *H<sup>b</sup> <sup>2</sup>*in other C4 crops (Basnayake *et al.*, 2015; Jackson *et al.*, 2016; Li *et al.*, 2017; Ferguson *et al.*, 2023)), considering: 1) the genotypes shared 75% of their genetic material (**Fig. 1**); 2) later attempts at finding variation in *iWUE* were not always promising (Hammer *et al.*, 1997; Blum, 2009; Leakey *et al.*, 2019; Pan *et al.*, 2022; Zhi *et al.*, 2022; Al-Salman *et al.*, 2023) and 3) previous key improvements in sorghum, such as the stay-green trait, were achieved via significant breeding contribution from wild sorghum relatives (Ochieng *et al.*, 2021). Despite the high  $H_b^2$ , the high *PCV* of *iWUE* (Table 2) meant that environmental factors that affect *g<sup>s</sup>* played an important role in driving variation of *iWUE*.

Hence, success in breeding for high *iWUE* is dependent on understanding the effect of different adaptive traits on *iWUE* and vice versa under different environments (Reynolds et al., 1994a; Araus et al., 2002). To screen for and expand the suite of such adaptive traits, we partitioned *iWUE* into a non-stomatal component (*ΔiWUEpc*) and a stomatal component (*ΔiWUEgs*) (as in Gilbert *et al.* (2011) and Li *et al.* (2017)), allowing us to reconcile high *iWUE* with photosynthetic performance and to link *iWUE* components to traits such as *Kleaf* or *θ*. Also, the variation in *iWUE* we found was associated with haplotypes where specific AQP genes were positively ascribed to parental lines from contrasting geographical regions and climates. This genetic information may be used for

further specific studies addressing the role of such AQPs, or the accompanied genes, in sorghum performances under both WW and WS conditions.

facilitating water transport from the apoplastic region to the inner cells, and vice<br>from the xylem to the stomata in the leaves, and hence keeping leaf cells hydrated<br>transpiration (Shope et al., 2008; Mott and Peak, 201 Our results hint at a possible role for two AQPs (SbPIP1.1 and SbTIP3.2) that might influence *iWUE* and related traits (**Fig. 2**). AQPs change permeability of cell membranes, facilitating water transport from the apoplastic region to the inner cells, and vice versa from the xylem to the stomata in the leaves, and hence keeping leaf cells hydrated during transpiration (Shope *et al.*, 2008; Mott and Peak, 2010; Chaumont and Tyerman, 2014; Li *et al.*, 2014). Water needs to enter guard cells for stomatal opening and increasing *g<sup>s</sup>* (Franks and Farquhar, 2007; Rockwell *et al.*, 2014; Buckley *et al.*, 2017), which subsequently increases *A<sup>n</sup>* and reduces *iWUE* (**Fig. 2**). The ability to maintain higher *g<sup>s</sup>* can be related to improved leaf hydraulic traits (Brodribb *et al.*, 2005). For example, the RP SbTIP3.2 haplotype had higher *Ψmidday*, higher *g*<sup>s</sup> and maintained *Kleaf* under WS compared to the NRP (**Fig. 2L**, **Table 3** and **Supplementary Table S3**). TIP AQPs are localised in the vacuolar membrane (tonoplast) and play a key role in maintaining cell turgor, possibly explaining the effect on leaf water status of SbTIP3.2 (Chaumont and Tyerman, 2014). Ectopic expression of a TIP gene has demonstrated that increased aquaporin activity generally leads to anisohydric behavior by promoting water transport within the plant and preventing stomatal closure (Maurel *et al.*, 2015). Furthermore, TIPs and PIP2s are known to transport the most abundant reactive oxygen species  $(H_2O_2)$ , which may have a role in plant cell signalling and even in ROS detoxication (Maurel *et al.*, 2015). However, WS also alters leaf pH and triggers ABA production and transport, which impact the activity of proton pumps associated with AQP activation and likely reducing AQP expression levels (Alexandersson *et al.*, 2005; Miyazawa *et al.*, 2008; Shatil-Cohen *et al.*, 2011; Pantin *et al.*, 2013; Shivaraj *et al.*, 2021). Therefore, it is also likely that other genes within that haplotype contribute to this response. Increased *A<sup>n</sup>* in RP SbTIP3.2 may be attributed to the higher *g*s, but also to more efficient ROS scavenging systems, which is in agreement with their higher chlorophyll content (as surrogated by SPAD) and electron transport rate, as inferred by higher *ΦPSII* (**Fig. 2J**  and **2K**). We did find significant differences in those two parameters between the RP and NRP haplotypes associated with SbTIP4.3/4.4 but with no impact on  $A_n$  (Table 3). Given that SbTIP4.3/4.4 genes are located in chromosome 3, but SbTIP 3.2 and SbPIP 1.1 in

chromosome 6 one close to the other (Reddy *et al.*, 2015), and that both haplotypes are from the elite parental line used in the Australian breeding program (RP SbTIP 3.2 and RP SbPIP 1.1) had higher both *A*<sup>n</sup> and *g*s, although lower *iWUE* than NRP haplotypes, they can be exploited to increase *A<sup>n</sup>* under predominantly well-watered conditions.

However, the NRP haplotype (associated with the parental line IS9710 originate<br>the dry region of Sudan) of the AQP SbPIP 1.1 had significantly higher  $\Delta \dot{W}U_{\rm Ggs}$ ,<br>aboveground biomass and LMA than the RP SbPIP 1.1 hap However, the NRP haplotype (associated with the parental line IS9710 originated from the dry region of Sudan) of the AQP SbPIP 1.1 had significantly higher *ΔiWUEgs*, *iWUE*, aboveground biomass and *LMA* than the RP SbPIP 1.1 haplotype under WW, suggesting a trade-off between higher carbon assimilation by unit of leaf area of the RP Australian line, but total plant assimilation of the NRP Sudanese line. This same haplotype (NRP SbPIP1.1) had the highest *ΔiWUEpc* of all haplotypes under WS, but also the highest aboveground biomass and highest *iWUE* of all haplotypes under WS, suggesting a probable function of SbPIP1.1 from the Sudanese haplotype also in WS response. Further studies are required to ascertain the functions of SbAQPs genes, and related genes associated to the haplotypes, identified in this study and the precise role of the highlighted AQPs in abiotic stress responses.

## **Screening for both high** *A<sup>n</sup>* **and** *iWUE* **under WS may be achieved through** *C<sup>i</sup>* **and might be associated with aboveground plant biomass**

In C<sup>4</sup> plants, increased *g<sup>s</sup>* under WW conditions may not be advantageous because C<sup>4</sup> photosynthesis saturates close to their operational *Ci*, resulting in the strong dependence of *iWUE* on *g<sup>s</sup>* (**Fig. 3B** and **4B**) as observed in previous studies (Jackson *et al.*, 2016; Cano *et al.*, 2019; Pignon *et al.*, 2021*b*; Pan *et al.*, 2022; Al-Salman *et al.*, 2023). Under WS, lower *g<sup>s</sup>* increases *iWUE* overall but also imposes a diffusional limitation on *A<sup>n</sup>* by lowering *Ci*. Hence, variation in photosynthetic capacity can overcome this diffusional limitation and increase *iWUE* by maximizing *A<sup>n</sup>* for a given *g<sup>s</sup>* (**Fig. 4B**), or rather *C<sup>i</sup>* as shown for genotypes with higher *ΔiWUEpc* having smaller *A<sup>n</sup>* reductions compared to *C<sup>i</sup>* (**Fig. 5C**) (Collyer and Adams, 2007; Gilbert *et al.*, 2011; Li *et al.*, 2017). *C<sup>i</sup>* can then be an indicator of not just *iWUE*, but of *ΔiWUEpc* (see strong association of *C<sup>i</sup>* with *ΔiWUEpc* compared to *ΔiWUEgs* in **Table S2**), confirming previous assumptions about *C<sup>i</sup>* as an integrator of *iWUE* and productivity in C<sup>4</sup> plants (Ghannoum, 2016; Jackson *et al.*, 2016; Condon, 2020). However, we found no strong relationship between *A<sup>n</sup>* or *ΔiWUEpc* with

biomass, apart from a weak relationship between *A<sup>n</sup>* and panicle size when both treatments are grouped (**Table S2A**). We also detected a weak (*R*=0.27) but statistically significant relationship between *iWUE* and total biomass (**Table S1B** & **S1C** - We note here that the significant, but low *R<sup>2</sup>* (and *R*) values displayed in our data are typical of studies focus on intra-specific diversity especially within crops and especially when exploring complex physiological traits that are underpinned by several processes (Pignon *et al.*, 2021*a*; Li *et al.*, 2022; Zhi *et al.*, 2022).

exploring complex physiological traits that are underpinned by several processes (et al., 2021 a; Li et al., 2022; Zhi et al., 2022).<br>
Efficient use of water at the leaf scale (higher leaf  $\Psi_{mvday}$  and lower plant hy<br>
re Efficient use of water at the leaf scale (higher leaf *Ψmidday* and lower plant hydraulic resistance (*Rrest*) (**Supplementary Table S2B** & **S2C**) combined with morphological adaptations such as narrower leaves (**Supplementary Fig. S4A**) and higher leaf density (as *LMA* increased but leaf thickness only marginally reduced under WS (**Supplementary Fig. S4C** & **S4D**)) can lead to reduced *g<sup>s</sup>* (see positive association between *g<sup>s</sup>* and *LW* in **Supplementary Table S2** also found in Pan *et al.* (2022) and Al-Salman *et al.* (2023)). This results in reduced water use and high *iWUE* leading to water conservation in the soil for biomass accumulation later in the season (Seneweera *et al.*, 2001; Vadez, 2019). Previous work on stay-green sorghum (most of our population is stay-green) showed that plant water use is lower during vegetative and early-reproductive stage, which is when we measured gas exchange, before ramping up during grain filling (Borrell *et al.*, 1999, 2014*b*,*a*, 2023; George-Jaeggli *et al.*, 2017). There is still scepticism about how much *iWUE* or photosynthesis *per se* can help drive productivity in future environments (Sinclair, 2012; Sinclair *et al.*, 2019), especially in C<sup>4</sup> crops (Sales *et al.*, 2021), since the yield of grain crops are heavily influenced by changing source-sink relationships and seasonal timings (Dingkuhn *et al.*, 2020; Fabre *et al.*, 2020). The impact of leaf-level physiological traits on whole-plant productivity under different conditions requires a comprehensive approach (Sreeman *et al.*, 2018; Tardieu *et al.*, 2018).

#### **Road map to select promising sorghum genotypes under soil water deficit**

A comprehensive physiological approach of crop drought response requires understanding of the relevant traits in response to the specific environment (Tardieu *et al.*, 2018). Too high *iWUE* under soil water deficit due to lowering *g<sup>s</sup>* is not desirable because this indicates the plant is experiencing moderate to severe water stress and has

and Europy and the method of properations. The control of the benefits of the conducted a physiologically extensive screen of more than 80 sorghum geodesicial conducted a physiologically extends the method of the conducti an overall lower plant water status and reduced *Kleaf* (Blum, 2009; Sinclair, 2012, 2018). Indeed, reductions of *Kleaf* and *Ψmidday* were associated with increasing *ΔiWUEgs* (and more closed stomata) (**Fig. 6A**, **Supplementary Fig. S5C** and **Supplementary Table S2**, coming at the expense of photosynthesis (see negative correlation between *ΦPSII* and *ΔiWUEgs* under WS (**Supplementary Fig. S5A**)). Higher *Kleaf* can help maintain *A<sup>n</sup>* under low *Ci*. Selecting for genotypes that respond to soil drought by taking some hydraulic "risks" (maintaining *Kleaf*) and keeping stomata relatively open under increasing WS may increase *iWUE* by increasing carbon accumulation as seen already in some grasses (Holloway-Phillips and Brodribb, 2011). Such a genotype would operate where the minimum *g<sup>s</sup>* is attained for the maximum *A<sup>n</sup>* (hence, high *iWUE* associated with high *ΔiWUEpc*) (**Fig. 5**). Traits that enable "risky" hydraulic behaviour without risk of cavitation can include deeper and more conductive roots, wider xylem vessels (Scoffoni *et al.*, 2011) and higher leaf vein density (already associated with higher *iWUE* in sorghum (Pan *et al.*, 2022; Al-Salman *et al.*, 2023). Other important traits can be related to extra-xylem conductivities such as enhanced mesophyll conductance (of  $CO<sub>2</sub>$  or H<sub>2</sub>O), reduced bundle sheath conductance, reduced airspace and more compact mesophyll structure around veins (Buckley, 2015; Buckley *et al.*, 2015; Sack *et al.*, 2015; Fiorin *et al.*, 2016; Xiong *et al.*, 2017, 2018; Pathare *et al.*, 2020; Al-Salman *et al.*, 2023), which are all processes influenced by AQPs (Maurel *et al.*, 2015; Negin and Moshelion, 2016; Groszmann *et al.*, 2017; Ermakova *et al.*, 2021). Combining water use strategy with gas exchange mechanisms is crucial to clarifying the benefits of increasing *iWUE* under different conditions (Liang *et al.*, 2023).

## **Conclusion**

We conducted a physiologically extensive screen of more than 80 sorghum genotypes selected based on differences in haplotypes originating from different parents from different origins and climates. We found significant variation among key traits, with some underpinned by differences between AQP-associated haplotypes inherited from an elite and exotic parent, providing a possible target genomic regions for beneficial traits. Partitioning the components of *iWUE* into stomatal anon non-stomatal components of *A*<sup>n</sup> allowed us to find a physiological mechanism that can lead to attainment of high *iWUE* without hindering photosynthesis or drought tolerance. We explained this mechanism through the connection between leaf and plant hydraulic conductivities and the maintenance of assimilation rates under low *Ci*. These findings provide a possible roadmap to expand the range of traits linked to *iWUE* in C<sup>4</sup> crops, offer possible avenues to bridge the trade-off between *iWUE* and productivity, and strengthens the case for AQPs as possible key players in this endeavour.

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## **Author contribution**

YA and FJC designed the experiment based on original ideas by FJC, MG and OG. EM oversaw genotype selection along with DJ and MG. YA led data collection alongside FJC. YA analysed all the data and wrote the manuscript with help from all the authors. OG oversaw project execution.

## **Conflict of interest**

The authors declare no conflict of interest.

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## **Data Availability**

The data generated and analysed for this study are available from the corresponding authors by request.

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**Table 1.** The Elite (RP) and Exotic (NRP) Parents used in the nested association mapping breeding program. Information about the parents and the production of the NAM population can be found in Jordan *et al.* (2011).





**Table 2.** Statistical summary of measured traits along with the calculated heritability and genetic variation information



*An*: Carbon assimilation rate; *gs*: Stomatal conductance; *iWUE*: intrinsic water use efficiency; *Ci*: Sub-stomatal carbon dioxide concentration; *ΦPSII*: Operating quantum yield of photosystem II; *SPAD*: Chlorophyll content measured by SPAD; *Ψmiddayf*: Midday leaf water potenial; *Ψpre-dawn*: Pre-dawn leaf water potenial; *LMA*: Leaf mass per area; *RWC*: Leaf relative water content; *Aboveground Biomass*: Total aboveground biomass; *Hb*: Broad-sense heritability; *GCV*: Genetic coefficient of variation; *PCV*: Phenotypic coefficient of variation.

**Table 3**. Summary of *P-*values from the mixed effect MANOVA of the parameters.(Bold: *P*<0.05). Population refers to the comparison between genotypes that have the recurrent parent (RP) haplotype for the AQP and genotypes that have the non-recurrent parent (NRP) haplotype for that AQP. Treatment means the watering level: well-watered and water-limited. Both comparison have two levels (df=1). *n* (6-63) for the number of independent genotypes per haplotype group for that specific aquaporin is in Table S3.





Abbrevations:- *An*: Carbon assimilation rate; *gs*: Stomatal conductance; *iWUE*: Instantaneous water use efficiency; *ΦPSII*: Operating quantum yield of photosystem II;  $\psi_{miday}$ . Midday leaf water potenial; Kleaf. Leaf hydraulic conductivity; LMA: Leaf mass per area; Rleaf. hydraulic resistance of plant leaf; Rrest. hydraulic resistance of rest of the plant; SPAD: Relative chlorophyll content using SPAD; *Tot Biom*: Total aboveground biomass; *ΔiWUEgs*: *iWUE* attributed to variation in *gs*; *ΔiWUEpc*: *iWUE* attributed to variation in *An.*

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**Fig. 1.** A simplified illustration of how recombinant inbred lines (RILs) are produced using nested association mapping (NAM).

**Fig. 2.** Bar charts showing the effects of two aquaporin haplotypes on key traits. The two AQPs shown are the ones that showed significant differences in several key traits between haplotype genotype populations, with the full analysis for all AQPs shown in Table 3. Each bar represents the mean of all individual replicates belonging to the genotypes of that population (*n*=18-63; see Table S2 for the number of genotypes for each Haplotype \* Treatment combination). Statistics shown are the result of ANOVA test and post-hoc Tukey test. Bars that share the same letter have no significant differences between them at *P*<0.05. For information about the approach to genotype selection, see materials and methods. Each population (RP & NRP) refers to a set of genotypes that have either inherited the AQP haplotype block from the elite parent (RP) or from the exotic parent (NRP). **(A)** & **(G)**The traits shown are: carbon assimilation rate (*An*); **(B)** & **(H)**  stomatal conductance (*gs*); **(C)** & **(I)** intrinsic water use efficiency (*iWUE*); **(D)** variation in *iWUE* due to *g<sup>s</sup>* (*ΔiWUEgs*); **(E)** leaf mass per area (*LMA*); **(F)** Aboveground biomass; **(J)** Leaf chlorophyll content (*SPAD*); **(K)** operating efficiency of photosystem II (*ΦPSII*); **(L)** midday leaf water potential ( $\Psi_{midadv}$ ).

**Fig. 3.** Relationship between leaf gas exchange parameters and hydraulic parameters. Data was collected on the YFEL and measured at saturating light levels (see Materials & Methods). Each point in scatter plots represent genotype mean (*n*=3). Standard error is presented in Supplementary Table S1. R<sup>2</sup> value and related significant level (\*\*\*=*P*<0.0001 & \*\*=*P*<0.05) are from a Pearson product-moment correlation analysis or from the exponential fit models. Leaf water potential measurements were collected from the leaf adjacent to the YFEL used for gas exchange. **(A)** Net carbon assimilation rate (*An*) vs stomatal conductance (*gs*); **(B)** intrinsic water use efficiency *(IWUE)* vs *gs*; **(C)** *iWUE* vs *An*; **(D)** *A<sup>n</sup>* vs. leaf hydraulic conductance (*Kleaf*); **(E)** *g<sup>s</sup>* vs midday leaf water potential *(Ψmidday)*; **(F)** *iWUE* vs *Kleaf*.

Table S2 for the number of genotypes for each Haplotype \* Treatment combination). Stats<br>howen are the result of ANOVA test and post-hoc Tukey test. Bars that share the same is now are the result of ANOVA test and the AQP. **Fig. 4.** The distribution of the components of intrinsic water use efficiency (*iWUE*) and their relationship with each other. The values in each scatter plot compromise the mean of every genotype (*n*=3) per treatment. For the bar chart the mean is of the genotype population (*n*=89 for WW and *n*=61 for WS). Data was collected on the YFEL and measured at saturating light levels (see Materials & Methods). Each point in scatter plots represent genotype mean (*n*=3). Standard error is presented in Supplementary Table S1.  $R^2$  value and related significant level (\*\*\*=*P*<0.0001 & \*\*=*P*<0.05) are from a Pearson product-moment correlation analysis or from the exponential fit models.  $R^2$  value from a Pearson product-moment correlation analysis.  $(A)$ variation in *iWUE* due to stomatal conductance (*ΔiWUEgs*) vs variation in *iWUE* due to photosynthetic capacity (*ΔiWUEpc*); **(B)** Bar chart showing the treatment effect on *ΔiWUEgs* & *ΔiWUEpc*; **(C)** *iWUE* vs *ΔiWUEgs*; **(D)** *iWUE* vs *ΔiWUEpc*.

**Fig. 5 (A)** Conceptual representation of the change in net assimilation rate  $(A_n)$  vs operational intercellular CO<sub>2</sub> concentration (C<sub>i</sub>) under progressive water stress driven mainly by stomatal limitation (black curve) or by concomitant decrease in both stomatal and non-stomatal limitations (grey line). The figure shows potential change in *C<sup>i</sup>* (*ΔCi*), the accompanying change in  $A_n$  (Δ $A_n$ ) and the degree of change in the  $A_n$  -  $C_i$  relationship (the angle  $\theta$ , with  $\theta$  = tan<sup>-1</sup>(Δ $A_n$ / *ΔCi* )) when the plant experiences water-limitation. **(B-C)** Relationship between degree of change in the *A<sup>n</sup>* - *C<sup>i</sup>* relationship (*θ*) from WW to WS with the change in the contribution of each component of intrinsic water use efficiency (i.e. *iWUE* variation due to stomatal conductance (*ΔiWUEgs*) and non-stomatal conductance or photosynthetic capacity (*ΔiWUEpc*)) also between

WW and WS (i.e. WS – WW). Data was collected on the YFEL and measured at saturating light levels (see Materials & Methods). Each point in scatter plots represent genotype mean (*n*=3). Standard error is presented in Supplementary Table S1.  $R<sup>2</sup>$  value and related significant level (\*\*\* *P*<0.0001 and \*\* *P*<0.05) are from a Pearson product-moment correlation analysis or from the exponential fit models

use efficiency from WW to WS (i.e. WS – WW) with lead hydraulic conductance ( $K_{\text{sub}}$ ) unrated that its and measured at staurating light levels (see Materials & Methods). Each point in scatter plots represents genotype me **Fig. 6** Relationship between the change in the contribution of each component of intrinsic water use efficiency from WW to WS (i.e. WS – WW) with leaf hydraulic conductance (*Kleaf*) under WS. Data was collected on the YFEL and measured at saturating light levels (see Materials & Methods). Each point in scatter plots represents genotype mean (*n*=3). Standard error is presented in Supplementary Table S1. R<sup>2</sup> value and related significant level (\*\*\*=*P*<0.0001 & \*\*=*P*<0.05) are from a Pearson product-moment correlation analysis or from the exponential fit models. Leaf water potential measurements were collected from the leaf adjacent to the YFEL used for gas exchange **(A)** Change in *iWUE* variation due to stomatal conductance (*ΔiWUEgs*) vs *Kleaf WS*; **(B)** Change in *iWUE* variation due to photosynthetic capacity (*ΔiWUEpc*) vs *Kleaf WS.*



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Figure 2



Figure 3







Figure 5



