# **Research** Article

# **Stay-Green QTLs Response in Adaptation to Post-Flowering Drought Depends on the Drought Severity**

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Stay-green trait enhances sorghum adaptation to post-flowering drought. Six stay-green backcross introgression lines (BILs) carrying one or more stay-green QTLs (Stg1-4) and their parents were characterized under non-stress ( $W_{100}$ : 100% of soil field capacity (FC)) and two levels of post-flowering drought ( $W_{75}$ : 75% FC;  $W_{50}$ : 50% FC) in a controlled condition. We aimed to study the response and identify the drought threshold of these QTLs under different levels of post-flowering drought and find traits closely contributing to grain yield (GY) under different drought severity.  $W_{50}$  caused the highest reduction in BILs performance. From  $W_{100}$  to  $W_{50}$ , the GY of the recurrent parent reduced by 70%, whereas that of the BILs reduced by only 36%.  $W_{75}$  and  $W_{50}$  induce different behavior/response compared to  $W_{100}$ . Harvest index contributed to the GY under the three water regimes. For high GY under drought transpiration rate at the beginning of drought and mid-grain filling was important at  $W_{75}$ , whereas it was important at mid-grain filling and late-grain filling at  $W_{50}$ . Stay-green trait can be scored simply with the relative number of green leaves/plants under both irrigated and stress environments. QTL pyramiding might not always be necessary to stabilize or increase the GY under post-flowering drought. The stay-green QTLs increase GY under drought by manipulating water utilization depending on drought severity.

# 1. Introduction

The global population will increase to 9 billion by 2050 and most of the increase will occur in sub-Saharan Africa [1], increasing the risk of food insecurity in this region [2]. Therefore, making plants resilient to the challenges of a water-scarce planet where climate change and global warming threaten food supplies is the major challenge facing the humanity [3]. Drought is perhaps the most important abiotic stress limiting crop productivity in the rain-fed agriculture around the world [4]. Sorghum (*Sorghum bicolor* (L.) Moench) is a grain crop that is well adapted to hot and dry climates. The productivity of sorghum, the major cereal crop grown for food, feed, and fuel, is usually under threat of terminal drought, which is likely to occur in rainfed environments during grain filling [5]. It is reported that stay-green genotypes that exhibited the ability to retain green leaf area during grain filling under terminal drought produce higher grain yield than the non-stay-green genotypes [6–10]. Over the last 30 years, the stay-green trait has been used in breeding programs for improvement of sorghum terminal drought tolerance [11, 12]. Several sources for stay-green have been identified, including B35, SC56, and E-36 [13–15]. The quantitative trait loci (QTLs) that contribute to the stay-green trait have been mapped in a range of populations, mostly derived from crosses with B35, a derivative of an Ethiopian dura landrace [16–23]. Xu and Sanchez [23, 24] identified four major QTLs in B35, including Stg1 (Stay-green QTL 1) and Stg2 both located on the linkage group SBI-03, Stg3 on SBI-02 and Stg4 on SBI-05. These QTLs explain 20, 30, 16, and 10%, respectively, of the phenotypic variation of the stay-green under post-flowering drought stress. Reddy [25] validated those QTLs reported in earlier studies and indicated that Stg2 and Stg3 were prominent in their expression.

The exact physiological mechanism of stay-green and the role of each individual QTL on the final phenotype of the stay-green genotypes under post-flowering drought stress are still unclear. However, a number of reports increased our understanding of this complicated drought adaptation mechanism. Borrell and Hammer [26, 27] explained the delayed onset and reduced rate of leaf senescence in staygreen genotypes by the high specific leaf nitrogen and nitrogen uptake during grain filling. Harris [28] reported that, under post-flowering water deficit, Stg2, Stg3, and Stg4 near isogenic lines (NILs) exhibited delayed onset of leaf senescence compared with the non-stay-green genotype, RTx7000, while significantly lower rates of leaf senescence in relation to RTx7000 were displayed by all of the stay-green NILs to varying degrees, but particularly by the Stg2 NIL. The Stg1 and Stg4 NILs exhibited greener leaves at flowering relative to RTx7000, indicated by higher SPAD values. Borrell [29] reported that hybrids containing B35, the source of staygreen, have higher transpiration efficiency (TE) than other eight hybrids examined. They suggested that the higher TE was due to increased photosynthetic capacity associated with higher specific leaf nitrogen, rather than reduced stomatal conductance. Vadez [9] studied the effect of different staygreen QTLs on modification of tillering and leaf area at flowering, transpiration efficiency, water extraction, harvest index, and grain yield under both terminal drought and fully irrigated conditions in 29 introgression lines with different stay-green QTLs in two backgrounds. They concluded that StgB and Stg1 modify the TE and water extraction depending on the background. Stay-green QTLs decrease the canopy size before flowering to conserve soil water for use during grain filling; the increased water uptake during grain filling in staygreen NILs relative to the non-stay-green parent RTx7000 resulted in higher biomass production, grain number, and yield [5, 30]. Moreover, Jaegglia [31] explained that tiller leaf area rather than transpiration efficiency, or transpiration per leaf area, was the main driver of weekly transpiration and the reduced pre-flowering water use in stay-green lines. According to Vadez and Borrell [5, 9, 30], the differences in TE are still unexplained and work is ongoing to investigate traits that might be related to leaf conductance aspects. In soils with good water-holding capacity, any water savings during the pre-flowering period increases water availability during the post-flowering period, therefore allowing plants to retain the photosynthetic capacity for longer by 'staying green' during grain filling [31].

The effect of the stay-green QTLs on modification of plant parameters under different levels of post-flowering drought is not extensively studied. Effect comparison between single QTL and QTL pyramiding under a specific level of drought severity has not been investigated adequately. Previously, we introgressed the stay-green trait into the Sudanese sorghum cultivar 'Tabat' [32, 33] and evaluated the response of the  $BC_2F_4$  stay-green lines under irrigated and drought conditions; we concluded that identification of the drought threshold is needed for better understanding the physiological reactions under specific drought severity. In this study, we analyzed the effect of single or more stay-green QTLs on modification of plant performance under controlled, nonstress condition and two levels of post-flowering drought stress by examining the plant physiological status through direct measurement of the leaves activity under drought. Also, we aimed to identify the level of drought to which those stay-green QTLs can confer drought tolerance. Moreover, we studied traits that mostly contribute to grain yield under different drought severity.

#### 2. Materials and Methods

2.1. Plant Materials. Six  $BC_3F_3$  stay-green backcross introgression lines (BILs) derived from a cross of drought sensitive cultivar 'Tabat' (abbrev. TAB, the recurrent parent) and staygreen donor B35 were used in this study. The backcross lines were produced at Agricultural Research Corporation, Sudan [32, 33]. Out of the six BILs, four carry a single stay-green QTL (Stg1, Stg2, Stg3, and Stg4, respectively), one BIL carries two QTLs (Stg1+4), and the other BIL carries three QTLs (Stg1+2+4). TAB is an improved Sudanese variety released for irrigated areas [34, 35] and B35 is a partially converted selection of the durra sorghum IS12555 from Ethiopia [13]. There was no big variation in the flowering of the genotypes and all flowered in the range from 70 to 76 days after sowing.

2.2. Pot Experiment. A pot experiment was conducted in a glasshouse in the Arid Land Research Center (ALRC), Tottori University (Tottori, Japan;  $35^{\circ}32'$ N,  $134^{\circ}13'$ E) from June to November. The pots were filled with 15 kg of Tottori sandy soil. The chemical properties were reported by Fujiyama and Nagai [36]. As reported by Sohail et al. [37], we used three inorganic fertilizers: compound macronutrients N:P:K (16:16:16) in a rate of 0.4g/Kg (Central Glass Co., Ltd, Tokyo, Japan) and Ca:Mg (21:0.6) (Hitachi Chemical Co., Ltd, Tokyo, Japan) at a rate of 0.7g/Kg and micronutrients Mg:Mn:B (8.4:0.3:0.3) (MC Ferticom Co., Ltd, Tokyo, Japan) at a rate of 0.3g/Kg.

Five seeds were sown per pot, and two weeks after sowing the seedlings were thinned and only one plant was allowed to grow beyond until maturity stage. After flowering, we applied three different drought levels (soil water regimes):  $W_{100}$  (100% of soil field capacity (FC); water content was 120 ml/kg soil);  $W_{75}$  (75% of soil FC; water content was 90 ml/kg soil) representing moderate drought; and  $W_{50}$ (50% of soil FC representing severe drought; water content was 60 ml/kg soil). The experiment was arranged in a completely randomized design with three replications. The position of each pot was randomized and changed weekly in the glasshouse to ensure uniform environmental conditions. Usually, the pots were weighted every day and irrigated with tap water to keep the specific FC; usually, pots in  $W_{100}$  were irrigated before reaching the FC of the pots in  $W_{75}$ , and pots in  $W_{75}$  were irrigated before reaching the FC of the pots in  $W_{50}$ , whereas pots in  $W_{50}$  were irrigated before reaching FC of 35%. During the experiment, the average maximum temperature in the glasshouse ranged from 34 to 23°C and the minimum ranged from 19 to 10°C.

2.3. Morphological Traits. Days to heading (DTH) were calculated as the number of days between the sowing date and the date when 50% of all the shoots in a pot had fully emerged spikes. At physiological maturity, plant height (PH) was measured in centimeters (cm) from the ground to the tip of the spike in each pot before harvesting. Days to maturity (DTM) were calculated from sowing date to 50% senescence of the spikes. Finally, grain yield per pot (GY) was determined as the weight (grams) of the grain from each pot; Biomass (BM) was determined as the weight (grams) of the aboveground fresh biomass, and harvest index (HI) was calculated using the formula

$$HI = \frac{Grain \ yield}{above \ ground \ biomass} * 100.$$
(1)

Yield susceptibility index (YSI) was calculated according to Fischer and Maurer [37]:

$$YSI = \frac{(1 - Y/Yp)}{D}$$
(2)

where *Y* is the GY of the genotype at drought, *Yp* is the mean GY of the genotypes at control, and *D* (stress intensity) = 1 - X/Xp, where *X* is the mean *Y* of all genotypes and *Xp* is the mean *Yp* of all genotypes. Genotypes were classified as highly tolerant (YSI  $\leq 0.50$ ), moderately tolerant ( $0.50 < YSI \leq 1.00$ ), or sensitive (YSI > 1.00) to drought [38, 39].

2.4. Leaf Measurement for Chlorophyll Content and Relative Number of Green Leaves/Plants. Detailed leaf observations were made on three replicates in each treatment. Fully expanded and senesced leaf number was recorded as described by Hammer [40] at one week before drought (WD), mid-grain filling (GF), and maturity (M).

A leaf was considered fully expanded when its ligule became visible above the enclosing sheath of the previous leaf. A leaf was considered senesced when more than 50% of its area turned yellow. Relative number of green leaves/plants (GN) was calculated as

$$\frac{\text{a number of green leaves per plant}}{\text{total number of leaves per plant}} * 100.$$
(3)

Leaf area of each individual fully expanded leaf was estimated nondestructively from the product of its length, greatest width, and a shape factor of 0.57, which was established by regressing the product of width and length of a leaf against its actual leaf area measured destructively at the end of the experiment. These estimates of individual leaf sizes, combined with observations of fully expanded and senesced leaves, allowed the estimation of green leaf area [41]. For the chlorophyll content (SPAD) data was represented as relative chlorophyll content (RCC) to ease the explanation and understanding of the degradation of leaf chlorophyll (senescence). Chlorophyll content was measured by the chlorophyll meter SPAD-502 (Konica Minolta). The arbitrary SPAD values can be translated into the actual value of total chlorophyll/unit area (mg cm<sup>-2</sup>) using the equation: Chlorophyll content= SPAD values x 0.003 — 0.048, as described by Xu [23].

2.5. Leaf Gas Exchange Measurement. The fully expanded second leaf from the apex position and flag leaf were used for measurements. The photosynthesis rate (PR) and transpiration rate (TR) were measured using LI-6400 portable photosynthesis system (LI-COR Bioscience, Lincoln, NE, USA), at three growth stages: WD, GF, and late-grain filling (LGF) during sunny days. During measurement the chamber temperature was 25°C, the reference CO<sub>2</sub> concentration was 400  $\mu$ mol mol<sup>-1</sup>, the relative humidity was approximately 25%, and the irradiance was 1200  $\mu$ mol m<sup>-2</sup> s<sup>-1</sup>.

2.6. Statistical Analysis. Two-way ANOVA was performed to assess the effect of genotype (G), drought treatment (W), and genotype-by-drought treatment (G×W) interaction for the different traits measured using GenStat version 17. The ANOVA was followed by Fisher's protected least significant difference (PLSD) test at P < 0.05. PCA analysis was performed using STAR software (STAR, version 1.4., International Rice Research Institute, Los Baños, Philippines; http://bbi.irri.org/products). Simple linear regression was performed using a linear regression model.

#### 3. Results

3.1. The Drought Threshold of the Stay-Green BILs. The drought treatment effect was significant for all traits studied under  $W_{100}$ ,  $W_{75}$ , and  $W_{50}$  (P<0.0001). In all traits except BM, PH, PR, and TR at WD,  $W_{100}$  had the highest mean values compared to  $W_{75}$  and  $W_{50}$  with the latter being the lowest (Tables 1 and 2). In the case of BM,  $W_{100}$  and  $W_{75}$  did not differ significantly, whereas, in the case of PH,  $W_{75}$  and  $W_{50}$  did not differ significantly. These findings indicated clearly that  $W_{50}$  as drought treatment was more severe than  $W_{75}$ . From  $W_{100}$  to  $W_{50}$ , the GY of the recurrent parent reduced by 70% whereas that of the BILs reduced by only 36%. The reduction from  $W_{100}$  to  $W_{50}$  did not exceed 50% in all traits except GLA and PR at LGF. Nevertheless, the BILs showed clear stay-green expression under  $W_{75}$  (moderate drought) (Tables 1 and 2).

3.2. The Effect of the Introgressed QTLs under No Drought at  $W_{100}$ . Generally, all the BILs with the different QTLs were comparable to or even better than TAB in their performance at  $W_{100}$  (Table 1 and Figures 1–5). However, BIL Stg1+4 had lower GY and HI, and BIL Stg3 had lower BM than TAB (Table 1). BILs Stg1, Stg2, and Stg1+4 had higher GLA than TAB at all stages (Figure 1(a)). BILs Stg3 and Stg1+2+4 had lower TR than TAB at WD and LGF, whereas at GF all BILs

		<u> Frain yield (g</u>		% Ον <b>ε</b>	er TAB		Biomass (g,		Ha	rvest index ('	(%	P	ant height (ci	m)
Genotypes	$W_{100}$	$W_{75}$	$W_{50}$	$W_{75}$	$W_{50}$	$W_{100}$	$W_{75}$	$W_{50}$	$W_{100}$	$W_{75}$	$W_{50}$	$W_{100}$	$W_{75}$	W <sub>50</sub>
TAB	48.4	28.0	14.4		I	300.0	255.0	210.0	16.1	12.1	6.4	167.3	145.0	142.3
B35	15.3	16.6	14.2		I	280.0	256.0	236.0	6.1	8.9	6.1	115.7	116.7	117.0
Stg1	46.5	35.5	27.0	26.9	87.5	338.3	320.0	260.0	12.1	10.4	7.1	166.3	136.0	134.0
Stg2	44.4	13.5	3.2	-51.8	-77.9	320.0	350.0	333.3	14.0	3.8	1.2	161.7	150.3	142.3
Stg3	42.2	43.3	41.2	54.6	186.1	250.0	246.7	206.7	16.5	17.1	1.61	172.3	162.3	166.3
Stg4	41.2	33.4	29.1	19.2	102.3	275.0	290.0	270.0	15.5	10.5	7.8	160.3	158.7	146.3
Stg1+4	39.0	35.0	31.0	25.1	115.3	333.3	310.0	315.0	10.7	8.3	4.0	160.0	156.3	149.0
Stg1+2+4	43.6	42.1	32.1	50.2	122.9	280.0	276.7	213.3	14.3	14.3	14.9	162.0	163.3	160.3
Mean	40.1	30.9	24.0			297.1	288.0	255.5	13.2	10.7	8.3	158.2	148.6	144.7
SEM±	2.195	2.210	2.560			7.510	7.860	9.740	0.740	0.860	1.140	4.290	3.330	3.180
<sup>a</sup> LSD ( <sup>b</sup> G)	8.880	6.180	8.180		I	38.060	34.840	23.690	3.2	3.1	1.2	24.9	10.2	12.0
LSD (G x <sup>c</sup> T)	3.980	3.980	3.980	I	Ι	18.650	18.650	18.650	1.630	1.630	1.630	9.41	9.41	9.41
P value (G)	< 0.0001	<0.0001	< 0.0001	I	Ι	0.002	0.000	< 0.0001	< 0.0001	<0.0001	< 0.0001	0.007	< 0.0001	<0.000
<i>P</i> value (G x T)	< 0.0001	< 0.0001	< 0.0001			0.004	0.004	0.004	< 0.0001	<0.0001	< 0.0001	$0.15 \mathrm{ns}^{\mathrm{d}}$	0.15ns	0.15ns
CV (%)	26	35	41			12	13	18	27	39	45	13	11	10

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TABLE parents	2: Green le evaluated	eaf area, rela under conti	ative chlorc rol conditio	pphyll conter on (W <sub>100</sub> , 10	nt, relative nu 0% soil field c	umber of gre apacity) and	en leaves/pla 1 two post-flc	unts, photos owering dro	ynthesis rate ught conditi	, and transpions $(W_{75}, 75)$	iration rate c % soil field c	of the six stay apacity; W <sub>50</sub>	y-green intr , 50% soil fi	ogression lir ield capacity	ies and their).
		<b>Freen</b> leaf a	rea	Re	lative chlorop	hyll	Rel	ative numbe	er of	Ph	otosynthesis	rate		Transpiration	ı rate
		$(cm^2)$			content		gre	en leaves/pl	ants	u <i>н</i> )	nol $CO_2 \text{ m}^{-2}$	$s^{-1}$ )	(n	$1 \text{ mol H}_2 \text{O} \text{ m}$	$^{-2} s^{-1}$ )
	WD	GF	Μ	WD	GF	Μ	WD	GF	Μ	WD	GF	LGF	WD	GF	LGF
$W_{100}$	$679^{a}$	$585^{a}$	$262^{a}$	I	I		$50.8^{a}$	$43.8^{a}$	$21.0^{a}$	$27.5^{a}$	$27.2^{a}$	$16.9^{a}$	$4.0^{a}$	$3.4^{a}$	$2.7^{a}$
$W_{75}$	$619^{a}$	$539^{\mathrm{b}}$	$251^{a}$	$95.2^{a}$	$95.2^{a}$	$53.8^{a}$	$45.8^{\mathrm{b}}$	$40.4^{\mathrm{b}}$	$18.5^{\mathrm{b}}$	$23.8^{\mathrm{b}}$	21.2 <sup>b</sup>	$10.2^{b}$	$4.1^{\mathrm{ab}}$	$1.4^{\mathrm{b}}$	$0.6^{\mathrm{b}}$
$W_{s0}$	$604^{a}$	$449^{c}$	$193^{\mathrm{b}}$	$92.8^{\mathrm{b}}$	77.9 <sup>b</sup>	$46.4^{\mathrm{b}}$	$43.5^{\mathrm{b}}$	$33.1^{\circ}$	$13.9^{\circ}$	$24.9^{\mathrm{ab}}$	$17.6^{\circ}$	$8.6^{\circ}$	$4.2^{\mathrm{b}}$	$2.2^{\circ}$	$1.8^{\circ}$

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	12 00	C.CI
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	42 CV	<b>C.C</b> <sup>‡</sup>
	dr zh	40.4
	77.0b	11.7
	do co	22.0
	qcui	C71
	1 10C	447
	60.4ª	004
2		50

WD: one week before drought; GF: mid-grain filling; LGF: late-grain filling; M: maturity. Different letters are significantly different (Fischer PLSD, P<0.05).



FIGURE 1: Green leaf area (GLA) at one week before drought (WD), mid-grain filling (GF), and maturity (M) under control  $W_{100}$  (a), 75% field capacity  $W_{75}$  (b), and 50% filed capacity  $W_{50}$  (c) of the six stay-green sorghum introgression lines evaluated with their parents under  $W_{100}$ ,  $W_{75}$ , and  $W_{50}$  of soil field capacity. Asterisks indicate significant difference from Tabat (TAB) (P < 0.05, Fisher's PLSD test).



FIGURE 2: Relative chlorophyll content (RCC) at one week before drought (WD), mid-grain filling (GF), and maturity (M) under control  $W_{100}$  (a), 75% field capacity  $W_{75}$  (b), and 50% filed capacity  $W_{50}$  (b) of the six stay-green sorghum introgression lines evaluated with their parents under  $W_{100}$ ,  $W_{75}$ , and  $W_{50}$  of soil field capacity. Asterisks indicate significant difference from Tabat (TAB) (P < 0.05, Fisher's PLSD test).



FIGURE 3: Relative number of green leaves/plants (GN) at one week before drought (WD), mid-grain filling (GF), and maturity (M) under control  $W_{100}$  (a), 75% field capacity  $W_{75}$  (b), and 50% filed capacity  $W_{50}$  (c) of the six stay-green sorghum introgression lines evaluated with their parents under  $W_{100}$ ,  $W_{75}$ , and  $W_{50}$  of soil field capacity. Asterisks indicate significant difference from Tabat (TAB) (P < 0.05, Fisher's PLSD test).

were comparable to TAB, but BIL Stg2 showed a substantial increase in TR (Figure 5(a)).

3.3. The Effect of the Introgressed QTLs under Stress Conditions at  $W_{75}$  and  $W_{50}$ . Overall, the performance of the stay-green BILs was better than that of the recurrent parent TAB under both  $W_{75}$  and  $W_{50}$  (Table 1, Figures 1–5).

All the BILs had higher GY than TAB under both  $W_{75}$  and  $W_{50}$  except BILs Stg2 and Stg4 (Table 1). BIL Stg2 had lower GY than TAB under both  $W_{75}$  and  $W_{50}$ , whereas BIL Stg4 had comparable GY to that of TAB under  $W_{75}$ . TAB and BIL Stg2 showed the highest reduction in GY from  $W_{100}$  to  $W_{75}$  and  $W_{50}$ , whereas B35 and Stg3 showed the lowest reduction. These lines showed rather substantially higher GY under  $W_{75}$  than that under  $W_{100}$ . This may explain the ability of the stay-green genotypes to increase translocation efficiency under a specific level of drought.

The other Stg lines were intermediate between their parents in their reduction (Table 1). The stay-green QTLs

improved the GY of TAB under drought by different magnitudes, the GY of TAB improved by 25.1% with Stg1+4 to 54.6% with Stg3 under  $W_{75}$ , and by 87.5% with Stg1 to 186.1% with Stg3 under  $W_{50}$  (Table 1).

To show the degree of drought tolerance conferred by each of the stay-green QTLs, we calculated the YSI at both  $W_{75}$  and  $W_{50}$ . TAB showed 1.3 and 1.6 YSI at  $W_{75}$  and  $W_{50}$ , respectively, and BIL Stg2 showed 2.9 and 2.3 at  $W_{75}$  and  $W_{50}$ . These were classified as sensitive (YSI > 1.00). BIL Stg3 showed -0.36 and -0.08, BIL Stg1+4 showed 0.55 and 0.56, and BIL Stg1+2+4 showed 0.22 and 0.50, under  $W_{75}$  and  $W_{50}$ , respectively. These were classified as highly tolerant (YSI  $\leq$ 0.50). BIL Stg4 showed 0.73 and 0.67 and was regarded as moderately tolerant (0.50 < YSI  $\leq$  1.00). BIL Stg1 was tolerant at  $W_{75}$  (0.49) but moderately tolerant at  $W_{50}$  (0.80).

In BM, BILs Stg1, Stg2, Stg4, and Stg1+4 had higher BM than TAB under both  $W_{75}$  and  $W_{50}$ . BIL Stg3 which had the highest GY under  $W_{50}$  had lower BM than TAB and was the least in the ranking of the genotypes at  $W_{50}$  (Table 1). Except for BIL Stg2, all of the BILs and their parents showed slight



FIGURE 4: Photosynthesis rate (PR) at one week before drought (WD), mid-grain filling (GF), and late-grain filling (LGF) under control  $W_{100}$ (a), 75% field capacity  $W_{75}$  (b), and 50% filed capacity  $W_{50}$  (c) of the six stay-green sorghum introgression lines evaluated with their parents under  $W_{100}$ ,  $W_{75}$ , and  $W_{50}$  of soil field capacity. Asterisks indicate significant difference from Tabat (TAB) (P < 0.05, Fisher's PLSD test).

or no reduction in BM from  $W_{100}$  to  $W_{75}$  and  $W_{50}$ . BIL Stg2 did not exhibit any reduction and had the highest BM at both  $W_{75}$  and  $W_{50}$  and was the top in the ranking of the genotypes (Table 1).

In HI, BILs Stg3 and Stg1+2+4 had higher HI than TAB under both  $W_{75}$  and  $W_{50}$ , whereas BILs Stg2 and Stg1+4 had lower HI than TAB under both treatments (Table 1). Among the BILs, Stg3 was the highest with no reduction from  $W_{100}$  to  $W_{75}$  and  $W_{50}$ , whereas Stg2 was the lowest with the highest reduction from  $W_{100}$  to  $W_{75}$  and  $W_{50}$  (Table 1).

In PH, BILs Stg3 and Stg1+2+4 had higher PH than TAB at both  $W_{75}$  and  $W_{50}$  (Table 1). At  $W_{75}$ , also Stg4 and Stg1+4 had higher PH than TAB. The highest reduction in PH was exhibited by Stg1, whereas Stg1+2+4 did not exhibit any reduction compared to TAB. Stg2 and Stg3 showed a slight reduction (Table 1).

In GLA, under both  $W_{75}$  and  $W_{50}$  at all stages, Stg1, Stg2, and Stg1+4 had higher GLA than TAB (Figures 1(b) and 1(c)). B35 did not show any reduction, whereas the reduction in TAB, Stg1 and Stg2, was higher than that in the other lines. From  $W_{100}$  to  $W_{50}$  the reduction in TAB and

B35 was higher than that in the other lines. All the lines showed a reduction of GLA from WD to GF with different magnitudes (Figure 1(b)). Line Stg1+4 showed the lowest reduction from WD to GF. From GF to M the reduction was higher than that from WD to GF (Figures 1(b) and 1(c)).

Generally, RCC of TAB reduced with the progress of the drought and was low at  $W_{50}$ , whereas BILs showed improved RCC than TAB (Figure 2). At WD, all the BILs were comparable to TAB, except BIL Stg3 under  $W_{75}$  and BIL Stg2 under  $W_{50}$  (Figures 2(a) and 2(b)). At GF, all the BILs were comparable to TAB under  $W_{75}$ , whereas they had higher RCC than TAB under  $W_{50}$  (Figures 2(a) and 2(b)). At M, under  $W_{75}$  all BILs except Stg1 and Stg3 had higher RCC than TAB, whereas under  $W_{50}$ , except Stg1+4, all the BILs had higher RCC than TAB. B35 had the highest RCC and maintained more than 75% of its RCC from WD to M at both  $W_{75}$  and  $W_{50}$ , whereas TAB maintained only 20% of its RCC and showed the highest rate of reduction (Figure 2). On the other hand, BILs showed different reduction magnitudes and maintained higher RCC than TAB under the  $W_{50}$  and



FIGURE 5: Transpiration rate (TR) at one week before drought (WD), mid-grain filling (GF), and late-grain filling (LGF) under control  $W_{100}$  (a), 75% field capacity  $W_{75}$  (b), and 50% filed capacity  $W_{50}$  (c) of the six stay-green sorghum introgression lines evaluated with their parents under  $W_{100}$ ,  $W_{75}$ , and  $W_{50}$  of soil field capacity. Asterisks indicate significant difference from Tabat (TAB) (P < 0.05, Fisher's PLSD test).

 $W_{75}$  treatments, which indicate their ability to retain more chlorophyll content than TAB.

In GN, all the BILs performed better than TAB (Figure 3). At WD, only BIL Stg2 was higher than TAB under both  $W_{75}$  and  $W_{50}$ , whereas at GF and M, all the BILs were higher than TAB, except Stg4 and Stg1+2+4 at M under  $W_{75}$  (Figures 3(b) and 3(c)). Similar to the RCC, all BILs showed less reduction rates compared to TAB, and they were able to partially maintain their GN at M under  $W_{50}$  when TAB was completely dry (Figure 3(c)).

In PR, all the BILs and their parent PR decreased with the progress of the drought from  $W_{75}$  to  $W_{50}$  and from WD to GF and LGF (Figure 4). At WD, the BILs showed comparable PR to that of TAB under both  $W_{75}$  and  $W_{50}$ , except that Stg4 and Stg1+4 under  $W_{75}$  had higher PR than TAB (Figure 4(b)). At GF, under  $W_{75}$ , all the BILs had higher PR than TAB, whereas under  $W_{50}$ , Stg2 and Stg1+2+4 showed higher PR than TAB (Figures 4(b) and 4(c)). At LGF, some BILs had comparable PR to that of TAB and others showed less PR than that of TAB. At GF the lowest reduction from  $W_{100}$  to  $W_{75}$  and from  $W_{100}$  to  $W_{50}$  was observed in line Stg1+2+4

indicating that QTL pyramiding is essential to maintain stable photosynthesis under drought conditions (Figure 4).

As in the PR, the TR decreased with the progress of the drought from  $W_{75}$  to  $W_{50}$  and from WD to GF and LGF (Figure 5), but interestingly BILs Stg3 and Stg1+2+4 showed the ability to maintain stable TR with the progress of the drought. However, their TR was lower than TAB under  $W_{100}$  without drought (Figure 5). At WD, BILs Stg1, Stg2, and Stg1+4 showed higher TR than TAB under both  $W_{75}$  and  $W_{50}$ . At GF, BIL Stg1+2+4 showed higher TR than TAB under both  $W_{75}$  and  $W_{50}$ , and BILs Stg2 and Stg3 were higher than TAB under than TAB under  $W_{50}$ . At LGF, under  $W_{75}$  Stg3 had substantially higher TR than TAB, whereas, under  $W_{50}$ , Stg3 and Stg1+4 had higher TR than TAB (Figure 5).

3.4. Principal Component Analysis. Principal component analysis (PCA) (Figure 6) illustrates the differences between the three water regimes in traits association. Sum of PC1 and PC2 explained 79.8, 63.4, and 62.4% of the total variation in  $W_{100}$ ,  $W_{75}$ , and  $W_{50}$ , respectively (Figures 6(a), 6(b), and 6(c)). Under all treatments, PC1 showed high coordination



FIGURE 6: PCA analysis of stay-green and yield traits measured in six stay-green sorghum introgression lines and their parents under  $W_{100}$ (a),  $W_{75}$  (b), and  $W_{50}$  (c). Traits included in the PCA are grain yield (GY), biomass (BM), harvest index (HI), plant height (PH), yield susceptibility index (YSI), relative chlorophyll content in SPAD units (RCC), relative number of green leaves/plants (GN), green leaf area (GLA), transpiration rate (TR), and photosynthesis rate (PR). PR and TR were measured at one week before drought (WD), mid-grain filling (GF), and late-grain filling (LGF) whereas RCC, GN, and GLA were measured at WD, GF, and maturity (M).

with GY and HI, whereas PC2 showed high coordination with BM and YSI. PC2 had a negative correlation with GY. Thus, it was called stress susceptibility component. This component separated genotypes with high and low GY in different environments.

Under the  $W_{100}$  condition with no drought, GY correlated with the HI, PH, PR, and TR at WD, GF, and LGF. The staygreen traits showed negative and no association with the GY (Figure 6(a)), although YSI of  $W_{75}$  and  $W_{50}$  were strongly correlated. Under  $W_{75}$ , the GY was associated with HI, PH, and PR at WD and GF, and TR at GF and LGF. There was an association between the GY and RCC at GF. GY correlated negatively with BM and YSI and the stay-green traits (GN and GLA) at all stages (Figure 6(b)). Under  $W_{50}$ , the GY was associated with HI, PH, and PR at WD and LGF, TR at LGF, and the RCC at WD and GF (Figure 6(c)). On the other hand, GY was negatively associated with the BM and YSI. There was a close association between BM and stay-green trait (GN and GLA) under all drought treatments.

Selection of genotypes with high PC1 and low PC2 indicates the suitable genotypes for both stress and non-stress environments [42, 43]. Based on the PCA we classified the genotypes into four groups according to their GY under non-stress and stress conditions: genotypes with high GY under

both stress and non-stress conditions (Group A), genotypes with high GY only under non-stress conditions (Group B), genotypes with high GY only under stress conditions (Group C), and at last genotypes with low GY under both conditions (Group D). Thus, Stg1+2+4 and Stg3 with rather higher PC1 and lower PC2 are superior genotypes under both stressed and non-stressed conditions (Figures 6(a), 6(b), and 6(c)). These genotypes had stable performance in the circumstances of low sensitivity to drought stress. Therefore, they belong to Group A. Stg2 could be known as Group D. This genotype is drought sensitive and had low GY and HI under drought conditions.

3.5. Regression Analysis. We applied regression analysis to provide more information on the significance of the important associations identified by the PCA analysis. The results indicated that HI contribution to the GY in the staygreen BILs was significant under the three water regimes  $(W_{100}, W_{75}, \text{ and } W_{50})$  (Figure S1). The PH had a significant association with the GY only under  $W_{100}$  without drought (Figure S2). Under  $W_{75}$ , PR was the most contributing factor  $(R^2=0.85^{**})$ , whereas under  $W_{50}$ , TR at GF was the most important factor contributing to GY  $(R^2=0.69^*)$  (Figure S3). GLA at M was strongly correlated with GN  $(R^2=79^{**}, R^2=0.87^{**})$  and  $R^2=0.87^{**})$  under  $W_{100}$ ,  $W_{75}$ , and  $W_{50}$ , respectively (Figure S4).

#### 4. Discussion

Stay-green is positively correlated with sorghum yield under post-flowering drought stress [10]. Although a positive correlation between stay-green and GY has been demonstrated in earlier studies, the physiological and molecular basis of the stay-green trait remains unclear [30]. Our results showed that stay-green QTLs affected a number of traits under terminal drought conditions in sorghum, and the significance and magnitude of the effects depended critically on the drought severity.

4.1. The Drought Threshold for the Stay-Green Lines. We performed this study at three levels of soil FC, W<sub>100</sub>, W<sub>75</sub>, and W<sub>50</sub>. Our results indicated that the performance of the stay-green lines reduced with the reduction in soil moisture content in all traits. W<sub>50</sub> was the most effective treatment in classifying and reducing the performance of the staygreen lines, and the most affected traits were the PR at LGF and GLA (Tables 1 and 2). From  $W_{100}$  to  $W_{50},$  the GY of TAB reduced by 70%, whereas that of the BILs reduced by only 36%. This finding shows clearly the effectiveness of the stay-green QTLs in enhancing the adaptation of TAB to the post-flowering drought. Interestingly, we observed clear expression and response of the stay-green trait under the moderate drought (W75) in terms of GLA and GN at all stages and RCC at maturity. An earlier report of Mahalakshmi and Bidinger [44] indicated that moderate but prolonged terminal drought stress during GF is the ideal environment for evaluating the stay-green trait. However, in this study we could identify that drought as 75% of the soil FC could

induce the expression of the stay-green and, hence, it could be enough to evaluate genotypes for stay-green traits.

4.2. The Impact of the Stay-Green QTLs under Non-Stress Conditions. Understanding the effect of the stay-green QTLs in the modification of plant performance under adequate soil moisture is essential as drought is fluctuating from year to year and place to place, and thus elasticity in crop performance is essential to have good yield with less or adequate water. Our results indicated that the single stay-green QTLs have no negative impact on the GY of TAB. However, the combination of Stg1+4 showed a decrease in TAB GY under W<sub>100</sub> conditions (Table 1). Other stay-green QTLs showed different impacts in TAB background; Stg1, Stg2, and Stg1+4 increased the GLA (Figure 1) and Stg3 and Stg1+2+4 decreased the TR at WD and LGF but not at the GF (Figure 5). The PCA analysis indicated that HI contributes to the GY (Figure 6(a)). Thus, we attribute the lower GY of Stg1+4 to its low HI. Borrell [30] reported that stay-green QTLs have no consistent yield penalty under irrigated conditions without drought. We attribute this contradiction of findings to the following: (1) Borrell [30] studied the effect of single QTLs only and did not study the effect of multiple QTLs, (2) there is a difference between the genetic backgrounds used in this study and that employed by Borrell [30], and (3) there is linkage drag as these lines still at BC<sub>3</sub>. Vadez [9] showed that the stay-green QTLs, Stg1 and Stg3, decreased the tillering and leaf area in S35 background, whereas there was no such effect in R16 background. Thus, they concluded that the impact of the stay-green QTLs depends on their interaction with the genetic background. Therefore, it is important for the breeding programs to consider this interaction to assure good performance and yield in the wet periods.

4.3. Association between the Studied Traits and GY under Normal and Drought Conditions. Understanding the association between the traits and the GY under different soil moisture contents is a prerequisite to decide which traits should be focused on in the breeding programs to increase the GY and also to understand how the stay-green contributes to increasing or stabilization of GY under drought conditions. Our results indicated that HI, PH, TR, and PR are the major traits contributing to the GY at all soil moisture levels (Table 1, Figure 6). Interestingly, the TR contribution to the GY differs among the treatments; it was important at the  $W_{100}$  condition and  $W_{75}$  at GF and LGF, whereas it was important at LGF at  $\mathrm{W}_{50}.$  The contribution of the PR varied with the variation in the soil moisture content. The contribution was high at the  $W_{100}$  and  $W_{75}$ , whereas it was less at  $W_{50}$  (Figures 6(a), 6(b), and 6(c)). At the  $W_{75}$  and W<sub>50</sub>, PR at LGF was more important than at WD and GF for higher GY. These findings indicated clearly that the response of the stay-green lines to the post-flowering drought depends on the drought severity level. These findings are consistent with the findings of Vadez [9]. They concluded that variation in GY of stay-green QTLs in different genetic backgrounds was due to HI and transpiration efficiency. The difference between our study and Vadez [9] is that they estimated

the transpiration efficiency by measuring the water supply and consumption, and we estimated the TR of the plants leaves by direct measurement of the leaf activity under two levels of drought at three different developmental stages. Thus, we were able to understand the change in the plant behavior with the progress of drought and development of the plant. Our findings and that of Vadez [9] indicate that the stay-green genotypes stabilize their GY under drought by manipulation of their behavior of water uptake and utilization, photosynthesis performance, and increasing the mobilization of the photo-assimilate to the grains (high HI). This manipulation of physiological performance or behavior depends on the degree of drought severity and the genetic background.

In this study, PH was associated with GY at the three water regimes (Figures 6(a), 6(b), and 6(c)). The association was high and significant under  $W_{100}$  compared to that at  $W_{75}$  and  $W_{50}$  (Figure S2). This could be explained by the findings of Sabadin [45] that PH QTLs were colocalized with the GY and stay-green QTLs. On the other hand, taking into consideration that also HI association with GY was higher under  $W_{100}$  compared to that under  $W_{75}$  and  $W_{50}$  (Figure S1), we can suggest that stay-green BILs are less reliant on the stem reserve under drought and operate another mechanism to maintain or stabilize their GY under drought. In addition, high GY could be reasonably predicted from PH ( $R^2$ =0.92<sup>\*\*\*</sup>) under non-stress environment.

HI was correlated with RCC at WD in  $W_{100}$  and  $W_{75}$  but not in  $W_{50}$ , where it was correlated with PR and RCC at GF and M (Figure 6). These results implied that remobilized reserves explained at least more than 60% of the variation observed in the GY of stay-green BILs tested under normal and moderate or severe drought condition. Ongom [46] suggested that plants with high remobilization could perform well under post-flowering drought. Under severe drought, PR and RCC at GF and M could be good indicators for high GY. Furthermore, PR had a positive association with GLA only under  $W_{50}$ , and this may explain its role under severe drought and could be explained by the findings of Swain [47]; that is, variation in photosynthesis is associated with leaf protein content.

The GLA and RCC were correlated at WD, GF, and M under  $W_{100}$ ,  $W_{75}$ , and  $W_{50}$  and were not correlated with the GY (Figure 6). These findings indicate that higher leaf senescence is due to higher translocation of food reserve from leaves to grains for better grain filling and increased GY as reported by Reddy [25]. This finding also explains the importance of the high HI trait in the stabilization of the GY under the drought conditions. This is typically the case of the BIL Stg3 as it had more HI and GY than the other BILs, but it had low GLA and RCC compared to the other BILs. Furthermore, GLA at M and GF stages were significant and positively correlated with GN under all soil moisture levels ( $W_{100}$ ,  $W_{75}$ , and  $W_{50}$ ) indicating that GN can be used as easy/fast indicator for stay-green trait instead of GLA.

4.4. Stay-Green QTLs Contribution to the GY Stabilization under Drought. As a result of their low rate of leaf senesce,

all the stay-green single QTLs improved or increased the GN under both the moderate and the severe drought (Figure 3), especially at M. Under  $W_{75}$ , Stg3 and Stg4 did not affect the GLA (Figure 1), whereas at  $W_{50}$  all the Stg QTLs increased/maintained high GLA. These results indicate that the impact of the Stg QTLs depends on the degree of the drought severity. Our results indicated that Stg2 and Stg1+4 had the lowest reduction in GN and GLA, and similar results were reported by Jordan [48].

The stay-green trait is positively correlated with GY in field conditions under terminal drought [6, 8, 10, 47]. The QTLs contribution in the GY varied with a variation in the soil moisture content or, in other words, with the severity of the drought. At the  $W_{100}$  condition, none of the QTLs increased the GY in TAB, and Stg1+4 decreased the GY, whereas at  $W_{75}$  and  $W_{50}$  all the stay-green QTLs increased the GY. Stg3 and the combination of Stg1+2+4 were the most efficient QTLs in terms of GY performance. These lines decreased the TR under W<sub>100</sub> condition, whereas under drought both possessed higher TR. Interestingly the high TR in these two lines was related to specific stages depending on the drought severity. At W75 Stg3 increased the TR at WD and LGF, whereas at  $\mathrm{W}_{50}$  the TR was high at the GF and LGF. Stg1+2+4 increased the TR at GF under both  $W_{75}$  and  $W_{50}$ (Figure 5). In addition, these two lines possessed high HI and increased PR at GF and LGF (Table 1, Figure 5). Based on these results we attribute the high GY of Stg3 and Stg1+2+4 to their high HI, PR, and TR. Stg3 is found to be positively important for improving GY under post-flowering drought stress (Table 1), and similar results were reported by Reddy [25]. Also, Sabadin [45] pointed out the colocalization of Stg3 and GY QTL and suggested the potential of indirect selection based on stay-green to improve sorghum GY under drought.

Stg2 showed very low HI and had a great reduction in GY under W<sub>75</sub> and W<sub>50</sub> compared to the other Stgs (Table 1), although it has a lower reduction in GN (Table 1). This result explains that variation in GY reduction depends on the stress severity since the variation in GY/panicle was found to be a function of terminal drought [39]. Moreover, introducing Stg QTL into highly senescent background could affect the sink source relationship as reported by earlier studies of Kassahun [49]. Our results confirmed previous reports that Stg2 is an important QTL for maintaining higher GLA contributing to slow senescence (Figure 4). This QTL was also reported to contribute to higher GLA at WD and M [50] and to %GLA at 45 days after flowering [15] in different genetic backgrounds. Reddy [25] indicated that the expression of Stg2 QTL was consistent and formed an important QTL for marker-assisted improvement of post-rainy sorghum lines for terminal drought tolerance. We attributed the low GY observed for Stg2 in our study to the presence of linkage drag and the difference of the genetic background and the environments.

4.5. Impact of the QTL Pyramiding. In this study, we compared the effect of the single QTLs and two combinations of double and triple QTLs on the adaptation to post-flowering drought stress. The performance in terms of GY of the single QTL Stg3 under both drought treatments  $W_{\rm 75}$  and  $W_{50}$  was similar or better than that of the double or triple QTLs (Table 1). However, the effect of the QTL pyramiding was evident in the tolerance of the BILs; using the YSI we classified Stg1 and Stg4 as moderately tolerant but when these QTLs were combined in Stg1+4 the tolerance increased, and Stg1+4 was classified as tolerant. Interestingly, when the sensitive QTL Stg2 was coupled with Stg1+4 the combination Stg1+2+4 was classified as tolerant. On the other hand, Stg2 had lower GY and HI under drought, but, when combined with Stg1+4 (the combination Stg1+2+4), it increased the GY and HI under drought (Table 1). These findings indicate that QTL pyramiding can enhance the adaptation to postflowering drought. However, this effect needs to be investigated across different backgrounds and environments, as the stay-green QTLs effect on improving adaptation to postflowering drought was found to be dependent on the genetic background and the environment [9, 51]. In addition, in this study, the single QTL Stg3 had comparable GY to that of the combination Stg1+2+4 which suggests that QTL pyramiding might not always be necessary depending on the environment and genetic background.

4.6. The Putative Model of Stay-Green Adaptation to Post-Flowering Drought. Using stay-green introgression lines in the background of RTX7000, Borrell [5, 30] concluded that stay-green genotypes adapt to the post-flowering drought through decreased tillering and the size of upper leaves, which reduced canopy size at flowering. This reduction in transpirational leaf area reduced pre-flowering water demand, thereby increasing water availability during GF and, ultimately, GY. Recently, Borrell [30] using the same introgression lines reported that tiller leaf area rather than transpiration efficiency, or transpiration per leaf area, was the main driver of weekly transpiration and the reduced pre-flowering water use in stay-green lines. In this study, we did not observe any reduction in the GLA of the staygreen lines, and stay-green QTLs Stgl, Sg2, and Stg1+4 increased the GLA of the senescent parent TAB before flowering under both control and drought conditions. We attribute this contradiction in findings to the difference in the genetic backgrounds used, especially that Vadez [9] showed that the stay-green QTLs, Stg1, and Stg3 decreased the tillering and leaf area in S35 background, whereas there was no such effect in R16. This contradiction demonstrates that (1) the stay-green plants adapt to the post-flowering drought through other different mechanisms and not only GLA reduction and water saving before flowering and (2) the stay-green effect depends largely on the genetic background.

In these earlier reports, plants water utilization behavior was evaluated by measuring the plant water consumption and transpiration efficiency. In this study, we measured the actual leaf TR at three different stages under two different levels of post-flowering drought severities. This enabled us to examine in more detail the behavior of the Stg QTLs in the modification of the plant behavior under drought. Thus, based on our results and the other reports we can suggest that stay-green genotypes adapt to post-flowering drought by reducing the transpirational leaf area and the TR per leaf that reduce pre-flowering water demand, thereby increasing water availability during grain filling and utilizing the conserved water depending on the drought severity and the genetic background.

## 5. Conclusion

In conclusion, our results clearly showed that the staygreen QTLs enhance post-flowering drought response to a level up to 50% of the soil FC and the stay-green trait is expressed under the moderate drought (W75). The staygreen QTLs help to increase or stabilize the GY under drought through efficient water utilization (TR) depending on the drought severity coupled with the high rate of photoassimilates translocation (high HI). QTL pyramiding could increase the drought tolerance but might not always be necessary to stabilize and increase the GY under postflowering drought. The understanding of the physiological mechanisms associated with drought severity, senescence, and photosynthetic efficiency and the connection between QTL expression/interaction with genetic background and physiological response to drought could be the key to remove the plateau of productivity associated with sorghum adaptation to unfavorable environmental conditions.

#### **Data Availability**

The data used to support the findings of this study are included within the article.

## **Conflicts of Interest**

The authors declare no conflicts of interest.

# **Authors' Contributions**

Nasrein Mohamed Kamal proposed the research, designed and performed the experiment, analyzed the data, and drafted the manuscript. Yasir Serag Alnor Gorafi performed the experiment and drafted the manuscript. Hisashi Tsujimoto designed the experiment and revised the manuscript. Abdelbagi Mukhtar Ali Ghanim, principal investigator of the project, supervised the research and revised the manuscript.

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#### **Supplementary Materials**

Figure S1. Relationship between grain yield (GY) and harvest index (HI) for six stay-green introgression lines and their parents evaluated under  $W_{100}$  (a),  $W_{75}$  (b) and  $W_{50}$  (c) of soil field capacity. \* denote significant differences at  $P \leq 0.05$ . Figure S2. Relationship between grain yield (GY) and plant height (PH) for six stay-green introgression lines and their parents evaluated under  $W_{100}$  (a),  $W_{75}$  (b) and  $W_{50}$  (c) of soil field capacity. \* \* \* denote significant differences at  $P \leq$ 0.001, ns denote not significant. Figure S3. Relationship between photosynthesis rate (PR) at grain filling and grain yield (GY) under W<sub>100</sub> (a), transpiration rate (TR) at grain filling and grain yield (GY) under W<sub>50</sub> (b) for six staygreen introgression lines and their parents. \*, \*\* denote significant differences at  $P \le 0.05$  and  $P \le 0.01$ , respectively. Figure S4. Relationship between green leaf area at grain filling and maturity (GLA.GF and GLA.M, respectively) and relative number of green leaves/plant under  $W_{100}$  (a),  $W_{75}$  (b) and  $W_{50}$  (c) for six stay-green introgression lines and their parents. \*, \* \*, \* \* \* denote significant differences at  $P \le 0.05$ ,  $P \le 0.01$ ,  $P \le 0.001$ , respectively. (Supplementary Materials)

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