

RESEARCH ARTICLE

Allelic Diversity of OsGW5.1 Regulating Grain Width in Rice

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ABSTRACT

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Rice is the staple food for more than half of the global population and rice production has to be increased by at least 50% during 2030 to meet the requirements of the rapidly increasing population. Rice yield is determined by several quantitative traits viz. number of productive tillers, panicle length, number of grains per panicle, grain size, and photosynthetic efficiency. Grain size is a major determinant of rice yield and it is primarily governed by grain length, grain width, and grain thickness. Several candidate genes modulating yield components have been discovered and put into breeding applications. OsGW5.1 (LOC_OsO5g25350), encoding for a receptor-like kinase is reported to alter grain width in rice. Advancements in genomics and re-sequencing of 3000 rice genotypes paved the way for measuring allelic diversity of putative candidate genes, identifying elite haplotypes, and developing functional markers for breeding applications. In this study, the allelic diversity of OsGW5.1 was analyzed in a set of 151 IRRI-3K panel lines. Eight non-synonymous SNPs were observed, and haplotype analysis identified four haplogroups of OsGW5.1. Differences in grain width of the different haplogroups were tested by Dunnet's test. Results revealed that haplotype 1 (H,) and haplotype 2 (H,) showed a significant difference in grain width. KIKUBA (haplogroup 1) was found to possess greater grain width (3.62 mm), whereas NS1515 (haplotype 2) possessed a slender grain type (1.88 mm). Identified superior donors possessing elite haplotypes of GW5.1 may be utilized in haplotype-based breeding to develop next-generation tailor-made high-yielding rice varieties with desired grain size.

Keywords: Rice; Grain Width; Allelic Diversity; Haplotype Analysis

INTRODUCTION

Rice, one of the most important staple food crop species, is consumed by half of the world's population, particularly in Asia. By 2050, the world's population is predicted to exceed 9.1 billion, which requires a simultaneous increase in rice production (Hibberd et al., 2008). Rice productivity has undergone two major leaps viz. first during 1960's through the introduction of semi-dwarfism and secondly during the 1980's through the introduction of hybrids. After that, no major breakthrough in rice yields was achieved. A number of issues, including water scarcity, soil salinity, disease, climate change, and reduced arable land area have indicated exacerbated food shortages over the next 50 years, demanding immediate improvement in crop production (Zhang, 2007). Any further increase in rice production has to overcome challenges viz. yield

plateau, shrinking natural (land and water) resources, and increased occurrence of abiotic stresses, pests, and diseases due to changing climate (Khush *et al.*, 2005). Designing high-yielding rice genotypes adapted to diverse environments depends on the exploitation of genetic resources through phenomics and well-knitted genomics-assisted breeding tools. This study made attempts to compile information on various yield genes in rice and survey the allelic diversity of key candidate genes. Literature survey enabled identification of 189 genes related to grain yield attributes and 63 genes regulating grain quality traits. Identification of favorable alleles associated with yield traits may accelerate haplotype-based breeding in rice.

An ideal rice cultivar should have high grain yield potential with improved grain shape, nutritional value,



disease resistance, and stress tolerance (Rosegrant and Cline, 2003). Rice grain size is a significant trait influencing rice yield and quality. It is also influenced by three geometrical dimensions including grain length, width and thickness with a wide range of variation (Li *et al.*, 2019). Domestication indicators include reduced nucleotide diversity and changed allele frequency in domestication loci. Thus, bridging the molecular study of gene function and domestication research requires a thorough understanding of the molecular basis for natural variation in domesticated features (Purugganan, 2019).

Re-sequencing 3K diverse rice germplasm lines enabled identifying and exploring allelic/ haplotype variations and thus harnessing genetic diversity (Abbai et al., 2019). Ultimately, this the way for identifying novel donors and novel alleles associated with the traits of interest, which can, in turn, be deployed in crop improvement (Varshney et al., 2018). In this context, the 3K RG re-sequencing project holds great promise for harnessing genetic diversity in rice (Li et al., 2014). The identified superior versions of candidate genes could be combined via the recently established, fast and robust 'haplotype assembly' concept (Bevan et al., 2017). For grain width, the gene GW5.1 was encoding a receptor-like kinase (LOC_OsO5g25350) was selected (Zhang et al., 2021). The mutant gw5.1 lines showed wider grain width and the lines which harbor the GW5.1

gene produced narrow grains (slender grain type). The allelic diversity of OsGW5.1 was measured using appropriate statistical tools and discussed results.

MATERIAL AND METHODS

A set of 151 diverse rice accessions from IRRI-3K Panel was assembled (Table 1) and evaluated during kharif 2020-21. The grain width of 151 accessions was measured using Vernier caliper. Ten grains per genotype were used for measurement of grain width. Measured data were subjected to descriptive statistical analysis using Minitab 19, and frequency distribution and box plot graphs were created (Allen, 2019).

Nucleotide polymorphisms existing in the GW5.1 (LOC_OsO5g25350) were visualized using SNP seek database and haplotype analysis was performed by downloading the allelic variants of the GW5.1 gene in PLINK format (Weeks, 2010). Haploview 4.2 software created haplotype groups and linkage disequilibrium blocks containing allelic variation, with settings such as a HW p-value (Hardy-Weinberg p-value) of 0.001, a minimum genotype percent of 75, and a minimum minor allele frequency of 0.001(Barrett *et al.*, 2005). Dunnett's test, performed with Minitab 19 Statistical Software, revealed significant differences between the generated haplotype groups (Allen, 2019).



Figure 1. a) Boxplot for grain width; b) Frequency distribution curve for grain width





Figure 2. a) Formation of haplotype groups based on allelic variation;b) Linkage Disequilibrium (LD) block of GW5.1 gene for 150 genotypes



Figure 3. Box plot curve for different haplotypes indicating significant difference by Dunnett's test

Table 1. List of genotypes used in the present study and their haplotype groups

S. No.	Accessions	Н
1	CX145	H1
2	CX94	H1
3	IRIS_313-8288	H1
4	IRIS_313-8265	H1
5	IRIS_313-10403	H1
6	IRIS_313-10423	H1
7	IRIS_313-7668	H1
8	IRIS_313-7684	H2
9	IRIS_313-8067	H2

10	IRIS_313-7650	H2
11	IRIS_313-7799	H1
12	IRIS_313-8703	H1
13	IRIS_313-8660	H1
14	IRIS_313-8697	H1
15	IRIS_313-9427	H1
16	IRIS_313-9574	H1
17	IRIS_313-9067	H1
18	IRIS_313-9503	H1
19	IRIS_313-9348	H1
20	IRIS_313-10113	H1
21	IRIS_313-10047	H1
22	IRIS_313-9302	H1
23	IRIS_313-10224	H3
24	IRIS_313-10167	H2
25	IRIS_313-9968	H1
26	IRIS_313-9709	H2
27	IRIS_313-9696	H1
28	IRIS_313-9605	H1
29	IRIS_313-8956	H2
30	IRIS_313-9325	H1
31	IRIS_313-9320	H1
32	IRIS_313-9841	H1
33	IRIS_313-10000	H1
34	IRIS_313-9160	H1
35	IRIS_313-10054	H2



36	IRIS_313-9469	H3
37	IRIS_313-10171	H1
38	IRIS_313-10333	H1
39	IRIS_313-10016	H1
40	IRIS_313-10260	H3
41	IRIS_313-10046	H1
42	IRIS_313-10300	H1
43	IRIS_313-10314	H1
44	IRIS_313-10396	H1
45	IRIS_313-10402	H1
46	IRIS_313-9227	H1
47	IRIS_313-8932	H1
48	IRIS_313-9116	H1
49	IRIS_313-9433	H1
50	IRIS_313-9986	H1
51	IRIS_313-8595	H1
52	IRIS_313-8638	H1
53	IRIS_313-8509	H2
54	IRIS_313-15902	H1
55	IRIS_313-10298	H1
56	IRIS_313-10353	H1
57	IRIS_313-11744	H1
58	IRIS_313-11377	H1
59	IRIS_313-11624	H1
60	IRIS_313-10664	H1
61	IRIS_313-11812	H1
62	IRIS_313-10924	H1
63	IRIS_313-11656	H1
64	IRIS_313-11852	H1
65	IRIS_313-11853	H3
66	IRIS_313-11692	H1
67	IRIS_313-11854	H1
68	IRIS_313-11671	H1
69	IRIS_313-11676	H1
70	IRIS_313-11721	H1
71	IRIS_313-10769	H1
72	IRIS_313-10966	H1
73	IRIS_313-11802	H1
74	IRIS_313-11505	H3
75	IRIS_313-11506	H1
76	IRIS_313-11867	H1
77	IRIS_313-11814	H1
78	IRIS_313-11543	H1
79	IRIS_313-11577	H1
80	IRIS_313-11234	H3

81	IRIS_313-10609	H1
82	IRIS_313-11869	H2
83	IRIS_313-11784	H1
84	IRIS_313-11760	H2
85	IRIS_313-11545	H2
86	IRIS_313-11811	H1
87	IRIS_313-11446	H1
88	IRIS_313-11638	H1
89	IRIS_313-11644	H1
90	IRIS_313-11762	H2
91	IRIS_313-11761	H2
92	IRIS_313-10458	H1
93	IRIS_313-10955	H2
94	IRIS_313-11820	H1
95	IRIS_313-10717	H1
96	IRIS_313-12296	H1
97	IRIS_313-10715	H1
98	IRIS_313-11521	H1
99	IRIS_313-11548	H1
100	IRIS_313-11507	H1
101	IRIS_313-11885	H3
102	IRIS_313-11126	H1
103	IRIS_313-10880	H1
104	IRIS_313-11256	H1
105	IRIS_313-11262	H1
106	IRIS_313-11263	H1
107	IRIS_313-11266	H1
108	IRIS_313-11271	H1
109	IRIS_313-11308	H1
110	IRIS_313-11310	H1
111	IRIS_313-11192	H1
112	IRIS_313-11241	H2
113	IRIS_313-11358	H1
114	IRIS_313-10986	H1
115	IRIS_313-10775	H1
116	IRIS_313-11327	H1
117	IRIS_313-10974	H1
118	IRIS_313-10754	H1
119	IRIS_313-11076	H1
120	IRIS_313-10555	H1
121	IRIS_313-11418	H2
122	IRIS_313-11242	H1
123	IRIS_313-11399	H1
124	IRIS_313-11141	H1
125	IRIS_313-11372	H1



126	IRIS_313-11181	H1
127	IRIS_313-11254	H1
128	IRIS_313-10433	H1
129	IRIS_313-10822	H1
130	IRIS_313-10753	H1
131	IRIS_313-10897	H1
132	IRIS_313-11793	H2
133	IRIS_313-11467	H1
134	IRIS_313-11899	H2
135	IRIS_313-11808	H1
136	IRIS_313-11954	H1
137	IRIS_313-8349	H1
138	IRIS_313-8493	H2
139	IRIS_313-9329	H1
140	IRIS_313-11758	H1
141	IRIS_313-12280	H2
142	IRIS_313-11509	H1
143	IRIS_313-11914	H2
144	IRIS_313-11225	H2
145	IRIS_313-11554	H1
146	IRIS_313-10441	H1
147	IRIS_313-10900	H1
148	IRIS_313-11251	H1
149	IRIS_313-10690	H1
150	IRIS_313-10628	H1
151	IRIS_313-8293	H4

Table 2. Descriptive statistics for OsGW5.1 among the study population

Statistics	Total population	
Range	1.88 - 3.62 mm	
Mean	2.65 mm	
Median	2.64 mm	
SE Mean	0.03	
CV (%)	13.21	

Table 3. Details of identified significant alleles

Marker number	SNP site	Position	HW pval	MAF	Alleles
1	10514718976	14718976	8.84	0.218	T:G
			E-34		
2	10514719513	14719513	1.15	0.022	A:T
			E-08		

3	10514720000	14720000	3.91	0.04	A:T
			E-09		
4	10514720008	14720008	4.38	0.011	G:A
			E-05		
5	10514720069	14720069	5.53	0.215	A:G
			E-35		
6	10514720549	14720549	1.53	0.182	G:A
			E-29		
7	10514720861	14720861	1.36	0.232	C:T
			E-28		
8	10514721040	14721040	1.88	0.03	A:G
			E-06		

Table 4. Statistical difference between haplotype groups

Statistics	H1 (121)	H2 (22)	H3 (7)
Range	1.93 -	1.88 -	2.23 -
	3.62 mm	2.83 mm	2.97 mm
Mean	2.7 mm	2.42 mm	2.66 mm
Median	2.68	2.42 mm	2.67 mm
SE Mean	0.03	0.05	0.09
CV (%)	13.12	11.11	9.47

Table 5. Dunnett's Simultaneous Tests for Level Mean-Control Mean

Difference	Mean	t value	p value
of levels	difference		
H2 - H1	-0.2747	-3.49	0.001
H3 - H1	-0.04	-0.31	0.942

RESULTS AND DISCUSSION

Descriptive statistics analysis defined the measures of variability among the 151 accessions. The coefficient of variation was found to be moderately high, indicating there is abundant variation in the population and thus, selecting suitable donors for the respective trait can be made with the genotypes available (Table 2). The range of grain width varied from 1.88 mm to 3.62 mm, with an average of 2.65 mm, according to the box plot curve (Fig1a). According to the histogram, the population has a normal distribution (Fig1b).

All the 151 germplasm lines were clustered into 4 haplogroups based on the sequence diversity of *OsGW5.1*. Eight non-synonymous SNPs identified in the *OsGW5.1* (LOC_OsO5g25350) were identified. The haplotype group *viz*. H1, H2, H3 and H4 were found to contain 121, 22, 7 and 1

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genotype, respectively. The H4 haplotype showed a heterozygous allele and it was not taken for further analysis. Eight SNPs (Table 3) from GW5.1 gene were employed to construct four haplotypes using haploview, as illustrated in Fig. 2a. The linkage disequilibrium block depicts the alleles at nearby positions that can occur on the same haplotype (Fig. 2b). In contrast to Mendel's law of independent assortment, linkage disequilibrium refers to the non-random association of alleles in a locus. The SNP sites 10514718976 and 10514720069 had the highest LD r2 value, indicating (Fig 2b-Values inside the blocks) a high correlation (98) between these two sites.

From Table 4, it was understood that haplotype H₁ contained maximum number of genotypes (121) followed by $H_2(22)$ and $H_3(7)$. H_4 was represented by only one genotype that showed heterozygous allele and was not included in the phenotypic analysis. The maximum grain width was observed in haplotype 1 (3.62 mm), followed by H_3 (2.97 mm) and H_2 (2.83 mm). From the Dunnett's test results, it was observed that there is a significant difference between H_2 - H_1 (p values <0.05) (Table 5 and Figure 3). KIKUBA was identified to possess a wider grain type (3.62 mm) observed in the H₁ haplotype group and NS1515 had a narrow grain type (slender, 1.88 mm) present in the haplotype group H₂. From this study, it was found that the genotype, NS1515 can be used as a donor for the development of rice genotypes with slender grain type. Similarly, Ramasamy et al. (2021) and Sundaramoorthy et al. (2022) identified superior haplotypes for An-1 and Gn1a genes, respectively.

CONCLUSION

The study resulted in unravelling the allelic variations in the GW5.1 gene with the help of 151 genotypes from the 3K RG panel, and their grouping based on haplotyping identified a superior haplotype H_2 which can be exploited in allele mining and plant breeding programmes. Thus, haplotype-based breeding is expected to aid in the development of high-quality rice varieties with desired grain width to fulfill the growing demands of the rice consuming population.

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Ethics statement

No specific permits were required for the described field studies because no human or animal subjects were involved in this research.

Originality and plagiarism

This is original research work and any work and/ or words of others, has been appropriately cited.

Consent for publication

All the authors agreed to publish the content.

Competing interests

There were no conflict of interest in the publication of this content.

Data availability

All the data of this manuscript are included in the MS. No separate external data source is required. If anything is required from the MS, certainly, this will be extended by communicating with the corresponding author through corresponding official mail; raveendrantnau@gmail.com; dsudhakar@ hotmail.com

Author contributions

Idea conceptualization - MR, Experiments - BA, Guidance – MR, DS, Writing original draft – BA, AP; Writing- reviewing & editing – BA, AP, DS, MR.

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