# The PLATZ Transcription Factor GL6 Affects Grain Length and Number in Rice<sup>1[OPEN]</sup>

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Grain size is one of the key determinants of grain yield. Although a number of genes that control grain size in rice (*Oryza sativa*) have been identified, the overall regulatory networks behind this process remain poorly understood. Here, we report the mapbased cloning and functional characterization of the quantitative trait locus *GL6*, which encodes a plant-specific plant AT-rich sequence- and zinc-binding transcription factor that regulates rice grain length and spikelet number. *GL6* positively controls grain length by promoting cell proliferation in young panicles and grains. The null *gl6* mutant possesses short grains, whereas overexpression of *GL6* results in large grains and decreased grain number per panicle. We demonstrate that GL6 participates in RNA polymerase III transcription machinery by interacting with RNA polymerase III subunit C53 and transcription factor class C1 to regulate the expression of genes involved in rice grain development. Our findings reveal a further player involved in the regulation of rice grain size that may be exploited in future rice breeding.

Rice (*Oryza sativa*) is one of the three major cereal crops in the world, and the most important staple food in Asia. Rice has served as a model monocot plant for molecular genetic dissection since its reference genome sequence was generated in 2005 (International Rice Genome Sequencing Project, 2005). The exploitation of rice genetics to increase grain yield and improve plant architecture are the focus of current rice-breeding programs (Huang et al., 2009).

Grain size, one of the most important determinate factors of grain yield, is specified by grain length,

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width, and thickness. In recent years, a number of genes and quantitative trait loci (QTLs) that control grain size have been identified and functionally characterized in rice. These grain size genes have been found to be involved in signaling pathways mediated by guanine nucleotide-binding proteins (G-proteins), proteasomal degradation, phytohormones, protein kinases, and transcriptional factors that control cell division and/or cell expansion during seed development (Zuo and Li, 2014). GS3, a major gene controlling grain length and weight, encodes a putative  $G\gamma$  protein that functions in G-protein signaling (Fan et al., 2006; Mao et al., 2010). GS3 suppresses the interaction between  $G\beta$  protein RGB1 (G protein  $\beta$  subunit 1) and two other  $G\gamma$  proteins, namelyDEP1 (dense and erect panicle1) and GGC2 (G protein  $\gamma$  subunit type C2), to antagonistically regulate grain size (Sun et al., 2018). These  $G\beta\gamma$  subunits are also colocalized in the nucleus and interact directly with the downstream effector OsMADS1 (MADS-box gene 1) to regulate rice grain length and vield (Liu et al., 2018). GW2 (grain width 2) encodes a RING-type E3 ubiquitin ligase that functions in protein degradation via the ubiquitin-proteasome pathway and negatively regulates cell division of the spikelet hull, thus affecting grain width (Song et al., 2007). GLW7 (grain length and weight on chromosome 7), a target gene of OsmiR156 (microRNA 156), encodes the plantspecific transcription factor OsSPL13 that positively regulates grain length and yield. The large-grain allele of GLW7 in tropical japonica varieties was identified to

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B.H. designed studies and contributed to the original concept of the project; A.W. and Q.H. performed most of the experiments; Q.Z. and Y.Li contributed to genome data analysis; Q.F., C.Z., D.F., Q.T., and Y.Lu performed the genome sequencing; L.S., X.H., J.L., D.L., J.Z., Y.S., J.M., Y.X., Y.W., and Z.W provided field genetic analysis, and A.W., Q.H., and B.H. analyzed whole data and wrote the paper. <sup>[OPEN]</sup>Articles can be viewed without a subscription.

be introgressed from *indica* varieties under artificial selection (Si et al., 2016). Copy number variation at the GL7 locus affects the expression of two linked genes to regulate grain length (Wang et al., 2015b). GW8, encoding the further SQUAMOSA PROMOTER BINDING PROTEIN (SBP)-domain transcription factor OsSPL16, positively regulates grain width and directly binds to the GW7/GL7 promoter to repress its expression (Wang et al., 2012, 2015a). Some genes involved in auxin response and brassinosteroid signaling, such as TGW6 and GW5, also influence the grain size and yield in rice (Ishimaru et al., 2013; Liu et al., 2017). Most of these genes regulate grain size by increasing or decreasing cell number (Qi et al., 2012; Zhang et al., 2012; Liu et al., 2015; Song et al., 2015; Yu et al., 2018; Zhao et al., 2018a), and a few genes control grain length by regulating cell size or length (Wang et al., 2015b; Si et al., 2016). In the case of GS2 and GS5, they affect grain shape by controlling both cell number and size (Li et al., 2011; Che et al., 2015; Hu et al., 2015).

These findings have enriched our knowledge of the regulatory mechanisms behind grain size in rice. However, how these genes are integrated into signaling pathways, as well as into the regulatory networks behind grain development, and the cross talk between them remain poorly understood. Therefore, identification and molecular characterization of new QTLs/ genes involved in grain size will help to comprehensively describe regulatory networks and serve for future improvement of rice yield (Duan et al., 2015).

Here, we report the identification and characterization of the rice grain length QTL *GL6*. *GL6* encodes a plant AT-rich sequence- and zinc-binding (PLATZ) protein and is preferentially expressed in young panicles. A mutant version of GL6 that carries a premature stop codon results in short grains via impaired RNA polymerase III (RNAPIII)-mediated transcription. We demonstrate that *GL6* functions in a further molecular mechanism that modulates grain length and grain weight.

# RESULTS

# Map-Based Cloning of GL6

In our previous work, we mapped a major QTL, *GL6*, which explained 20.5% of the phenotypic variation (*R*<sup>2</sup>) in grain length within a set of 271 lines derived from a cross between a cultivated rice variety, *O. sativa* ssp. *indica* Guangluai4 (GLA4) and a wild rice accession, *Oryza rufipogon* W1943 (W1943; Huang et al., 2012). The *GL6* locus was initially mapped to the region between the recombination bins Bin\_1024 (53.3 cM) and Bin\_1028 (55.5 cM) on chromosome 6.

To fine map this locus, one backcross inbred line, BIL219, with shorter grain length, harboring the *GL6* locus and carrying several other segments from W1943 (Supplemental Fig. S1), was selected and backcrossed to GLA4. We then carried out high-resolution mapping using 2,181 BC<sub>1</sub> $F_5$  individuals, and the locus was finally delimited to a 6.1-kb region between the two markers GL4350 and GL4411 (Fig. 1, B–D).

According to the Rice Annotation Project annotation (http://rapdb.dna.affrc.go.jp/), this region contains only one candidate gene, Os06g0666100, which encodes a PLATZ protein. Compared to GLA4, the coding region of GL6 from BIL219 contained a C-to-T substitution in the third exon at nucleotide 352 that introduced a stop codon resulting in premature termination of translation (Fig. 1E; Supplemental Fig. S2). Interestingly, the original donor parent W1943 represented a T/C heterozygote at the premature stop codon allele; however, all other singlenucleotide polymorphisms (SNPs) were identical to those in BIL219. Thus, we considered that BIL219 inherited the T allele from the generation of progeny. In addition, we isolated three genotypes of GL6 in a filial generation by self-pollinating the heterozygous W1943, and found that the grain size of W1943 was related to different *GL6* genotypes (Supplemental Fig. S3). Thus, we hypothesized that this premature stop codon might cause reduction or loss of function of GL6, thus affecting rice grain length.

# Characterization and Validation of *GL6* Function in Regulating Grain Length

To further investigate the function of *GL6*, we developed a near isogenic line (NIL), NIL-*gl6*, containing a 17-kb "T-type" W1943 chromosomal region at the *GL6* locus in the GLA4 genetic background (Fig. 1F). Compared with GLA4, NIL-*gl6* showed shorter grains (a 10.4% decrease) and lower 1,000-grain weight (a 14% decrease). However, the width of the grains was not affected (Fig. 1, G and H).

In order to verify the function of the candidate gene GL6, we generated a genetic complementation construct in which the GL6 locus from GLA4, containing the entire gene region, the 9277-bp promoter region, and the 728-bp downstream sequence, was introduced into NIL-gl6. In comparison with NIL-gl6, the transgenic genetic complementation lines NIL-gl6-CP1 and NIL-gl6-CP2 showed obvious increases in grain length and weight, even greater than in GLA4, alongside a slight increase in grain width (Fig. 2, A and B; Supplemental Fig. S4). In addition, we carried out gene mutation of GL6 by using the clustered regularly interspaced short palindromic repeats (CRISPR)/ Cas9 genome editing system (Ma et al., 2015) in GLA4 (Fig. 2C) and a *japonica* variety, Nipponbare (Supplemental Fig. S5A). In GLA4, the resulting GL6 loss-of-function CRISPR-Cas9-derived mutants, GLA4gl6<sup>CRISPR</sup>1 and GLA4-gl6<sup>CRISPR</sup>2, displayed reduced grain length and weight when compared with the wild-type plants (Fig. 2, D and E; Supplemental Fig. S5, B-E). Altogether, these results demonstrate that Os06g0666100 is the causative gene for the QTL GL6

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Figure 1. Map-based cloning of GL6. A, Location of GL6 on rice chromosome 6, with 271 chromosome segment substitution lines. B, High-resolution mapping of the GL6 region was performed with 2,181 BC<sub>1</sub>F<sub>5</sub> lines. The number of recombinants between molecular markers is indicated below the linkage map. C, GL6 was narrowed down to a 6.1-kb genomic region. D, Grain length is shown for each recombinant line. Values are given as the mean  $\pm$  sp. E, Gene structure and allelic variations of GL6 between GLA4 and BIL219. Colored boxes and black lines represent exons and introns, respectively. The start and stop codons are indicated above the gene, and the coding region is blue. F, Chromosome maps of NILgl6. NIL-gl6 contained the T-type W1943 allele at GL6 in the 17-kb region on chromosome 6, shown as a red bar. G, Mature paddy grain (upper) and brown rice (lower) morphology of GLA4 and NIL-gl6. Scale bar = 5 mm. H, Comparison of grain length, width, and 1,000grain weight between GLA4 and NIL-gl6. Values are given as the mean  $\pm$  sp. \*\*\*P < 0.001, significant difference determined by Student's t test.

and functions in the positive regulation of rice grain length and weight.

# GL6 Influences Cell Proliferation to Regulate Grain Length

Since cell division and cell expansion are responsible for altering grain size and grain size is restricted by the size of the spikelet hull, we compared the epidermal cells of GLA4 and NIL-*gl6* by scanning electron microscope. Observations of the outer glume surface showed that total cell number along the longitudinal axis in NIL-*gl6* was reduced by 17.38% compared with that in GLA4, without any significant difference in single cell length (Fig. 3, A–D). Similarly, total cell number of the outer epidermal cells was also decreased by 20.72% in GLA4-*gl6*<sup>CRISPR</sup>1 and increased by 16.54% in NIL-*gl6*-CP1 (Fig. 3, A–D). The CRISPR/Cas9-derived *gl6* mutant in Nipponbare, NIP-*gl6*<sup>CRISPR</sup>1, also showed a reduction in cell number (Supplemental

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**Figure 2.** Validation that *GL6* controls grain length. A, Grain morphology of GLA4, NIL-*gl6*, and two independent complemented transgenic lines (NIL-*gl6*-CP1 and NIL-*gl6*-CP2), Scale bar = 5 mm. B, Comparisons of grain length, width, and 1,000-grain weight among the lines shown in A. Lowercase letters indicate significant differences (P < 0.05) as determined by Duncan's multiple range test. C, Sequence of the CRISPR mutant alleles. The wild-type sequence is shown at the top, with the target sites underlined in black and the protospacer adjacent motif sequence highlighted in blue. Deletions are shown as red dashes. D, Grain phenotype of GLA4 and two independent CRISPR transgenic lines (GLA4-*gl6*<sup>CRISPR</sup>1 and GLA4-*gl6*<sup>CRISPR</sup>2). Scale bar = 5 mm. E, Comparisons of grain length, width, and 1,000-grain weight among the lines shown in D. Values are given as the mean  $\pm$  sp. \*\*\* Significant difference (P < 0.001, Student's t test).



**Figure 3.** The effect of *GL6* on cell number contributes to grain length. A and B, Scanning electron micrographs of the whole grain (A) and outer glume surfaces (B) in GLA4, NIL-*gl6*, GLA4-*gl6*<sup>CRISPR</sup>1, and NIL-*gl6*-CP1. Scale bars = 1 mm (A) and 100  $\mu$ m (B). C and D, Comparisons of single cell length (C) and total cell number (D) among the lines shown in A. E, Comparisons of relative expression levels of *GL6* among the lines shown in A. Values are given as the mean ± sp. Lowercase letters indicate significant differences (P < 0.05) as determined by Duncan's multiple-range test. F, Relative expression levels of 26 cell cycle-related genes in young panicles (0–1 cm) of GLA4 and NIL-*gl6*. *OsUBQ5* was used as the control and the values of expression levels in GLA4 were set to 1. Values are given as the mean ± sp. Student's *t* test significant difference: \*P < 0.05, \*\*P < 0.01 and \*\*\* P < 0.001; ns, not significant.

Fig. S5, F–I). Consistent with this, comparative measurement of GL6 expression level among these plants showed that NIL-gl6-CP1 exhibited elevated GL6 transcripts, whereas decreased GL6 expression levels observed in GLA4-gl6<sup>CRISPR</sup>1 and NIP-gl6<sup>CRISPR</sup>1 (Fig. 3E; Supplemental Fig. S5J). Together, these results show that higher expression level of GL6 contributes to increased cell numbers for the spikelet hull, thus resulting in large grains. As expected, expression levels of 12 cell cycle-related genes, including CAK1A, CYCD4, CYCT1, E2F2, and H1, were significantly down-regulated in NILgl6 (Fig. 3F), indicating that the reduced cell number in NIL-gl6 might result from decreased expression of genes that promote cell proliferation. Hence, these results suggest that GL6 positively regulates grain length by altering cell division instead of cell expansion to control cell number of the glume during spikelet development.

# GL6 Negatively Regulates Grain Number per Panicle

In addition to the grain size and weight, we also compared other agronomic traits among GLA4, NILgl6, and transgenic lines in field trials. We found that larger seed had fewer grains per panicle (Fig. 4A). Compared to that in GLA4 (107.25  $\pm$  15.16), the number of grains per panicle was increased by 32.9% and 33.4% in NIL-gl6 (141.53 ± 19.89) and GLA4- $gl6^{CRISPR}$ 1  $(143.10 \pm 20.82)$ , respectively. However, the complemented transgenic line NIL-gl6-CP1 showed a dramatically reduced grain number (84.16 ± 14.74; Fig. 4E). We then measured the panicle length, primary branches, and secondary branches in these lines (Fig. 4, B-D). NIL-gl6 and GLA4-gl6<sup>CRISPR</sup>1 exhibited more secondary branches, whereas NIL-gl6-CP1 had fewer secondary branches, compared to GLA4 (Fig. 4D). The increased grain number was mainly attributed to the increased number of secondary branches. Other panicle phenotypes, such as panicle length and primary branches, showed only a slight effect on grain number. Moreover, we observed that the complemented line showed significantly increased tiller number per plant compared to other lines (Fig. 4F). Taking all these effects into account, the grain yield per plant was lower in NIL-*gl6,* GLA4-gl6<sup>CRISPR</sup>1, and NIL-gl6-CP1 than in GLA4 (Fig. 4G). These results reveal that GL6 influences both panicle and spikelet development and that a balance between grain number and grain size may exist that determines grain yield.

# Spatial Expression Pattern of GL6

The expression pattern of *GL6* was detected by reverse transcription quantitative PCR (RT-qPCR) analysis. *GL6* expression was found in all organs and tissues examined; higher expression levels were observed in young panicles (<1 cm), but these gradually decreased during panicle development (Fig. 5A). Furthermore, we investigated the specific temporal and spatial expression pattern of *GL6* by RNA in situ hybridization (Fig. 5,

B–M). *GL6* transcripts were initially detected when the secondary branch primordia were formed (Fig. 5B). During subsequent growth, GL6 transcripts were observed in abundance in both the lemma and palea primordia (Fig. 5, C, E, and F). With the development of floral organs, lemma, palea, and stamen primordia all exhibited a strong expression signal of GL6 at stage spikelet development 6 (Sp6; Fig. 5G). Subsequently, the expression of GL6 was restricted to the stamen and carpel primordia, and gradually decreased in the primordia of lemma and palea (Fig. 5, H and I). Finally, after floral organ differentiation, GL6 expression signals disappeared during late-stage Sp8 (Fig. 5J). No signals were detected with the sense probe (Supplemental Fig. S6A). By contrast, compared to expression of the positive control HISTONE H4, we could barely detect GL6 expression in NIL-gl6 at any stage (Fig. 5, K-M; Supplemental Fig. S6B).

# GL6 Interacts with OsRPC53 and OsTFC1

RNAPIII is a multisubunit complex eukaryotic RNA polymerase that transcribes tRNA genes, 5S ribosomal RNA (5S rRNA), RNase P, and other noncoding RNAs to regulate RNA and protein synthesis for multiple cellular developmental processes (Abascal-Palacios et al., 2018). The maize PLATZ protein FL3 was revealed to interact with RPC53 (RNA polymerase III subunit C53) and TFC1 (transcription factor class C1) of the RNAPIII complex to modulate the RNAPIII transcription machinery (Li et al., 2017). Amino acid sequence analysis showed that GL6 shared high sequence similarity with other PLATZ proteins, and phylogenetic analysis revealed that GL6 belonged to the PLATZ family (Supplemental Figs. S7 and S8). Considering the conserved PLATZ domain and potential functional conservation among the PLATZ family, we wondered whether GL6 could affect the function of RNAPIII in rice. We tested the interaction between GL6 and three rice homologs of RNAPIII transcription machinery, namely OsRPC53, OsBRF1 (TFIIIB B-related factor 1), and OsTFC1, respectively. Yeast two-hybrid assays indicated that GL6 interacts with OsRPC53 and OsTFC1, but not OsBRF1 (Fig. 6A). Further in vivo interactions were validated using a bimolecular luciferase complementation (BiLC) assay, and we found that the interaction between GL6 and OsRPC53 was much stronger than that between GL6 and OsTFC1 (Fig. 6B). Additionally, RNAPIIIdependent transcripts of tRNAs and 5S rRNA were decreased in NIL-gl6 compared with GLA4. These results suggest that GL6 might participate in mediating RNAPIII to coordinate ribosome biogenesis (Fig. 6C).

# RNA-Seq Analysis of the *GL6* Downstream Regulatory Network

To further explore the regulatory mechanism of *GL6*, we carried out RNA sequencing (RNA-seq) analysis of



**Figure 4.** *GL6* affects panicle morphology. A, Panicles of GLA4, NIL-gl6, GLA4-gl6<sup>CRISPR</sup>1, and NIL-gl6-CP1. Scale bar = 5 cm. B to G, Comparisons of panicle length (B), primary branches (C), secondary branches (D), spikelets per panicle (E), tiller number per plant (F), and grain yield per plant (G), among the lines shown in A. Values are given as the mean  $\pm$  sp. Lowercase letters indicate significant differences (P < 0.05) as determined by Duncan's multiple-range test.

Figure 5. The spatial expression pattern analysis of GL6 in rice. A, Relative expression levels of GL6 mRNA in vegetative tissues and developmental panicles. The abundance of GL6 transcripts was normalized to that of OsUBQ5 (ubiquitin 5). Values are given as the mean  $\pm$  sp. B to J, In situ analysis of GL6 in GLA4 during different inflorescence stages and flower development processes. GL6 transcripts at stages In5 (B), In6 (C), In7 (D), Sp4 (E), Sp5 (F), Sp6 (G), Sp7 (H), Sp8e (I), and Sp8e (J). K to M, In situ analysis of GL6 in NIL-gl6 at stages In7 (K), Sp6 (L), and Sp7 (M). ca, carpel; fm, floral meristem; le, lemma; lo, lodicule; ov, ovule; pa, palea; pb, primary branch; sb, secondary branch; sp, spikelet; st, stamen; Sp8e, early-stage Sp8; Sp8l, late-stage Sp8. Scale bars =  $50 \ \mu m$ .



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young panicles from GLA4 and NIL-gl6 to investigate the downstream gene regulatory networks. Principlecomponents analysis of the gene expression data showed close clustering of biological replicates and clear differentiation of separate samples (Supplemental Fig. S9A). A total of 2,711 differentially expressed genes (DEGs) were detected (P < 0.05), of which 52.6% (1,426) up-regulated genes and 47.4% (1,285) down-regulated genes were found in NIL-gl6 relative to GAL4 (Supplemental Table S1). Gene Ontology (GO) analysis of these DEGs showed significant enrichment (false discovery rate < 0.05) of the biological process associated with transcription, transport, translation, and hormone stimulus (Supplemental Fig. S9B). Similarly, molecular functional categories associated with DNA binding, transferase activity, transcription regulator activity, protein binding, and signal transducer activity were also highly enriched (Supplemental Fig. S9, C and D). In addition, we found that OsMADS1 (a known gene controlling grain size; Liu et al., 2018), eIF1 (a protein translation factor), and *TFIIS* (a transcription elongation factor) were down-regulated in NIL-gl6, and further RT-qPCR analyses verified the differential expression of these genes (Supplemental Fig. S9E). These results support the hypothesis that GL6 functions as a transcription factor to regulate downstream gene expression.

Figure 6. GL6 interacts with RPC53 and TFC1 to regulate tRNA transcripts. A, Yeast two-hybrid assay, showing how GL6 interacts with OsRPC53, OsTFC1, and itself. AD, activation domain; BD, binding domain; 10-fold serial dilutions show the gradients. B, BiLC assay showing that GL6 interacts with OsRPC53 and OsTFC1 in vivo. C, Relative expression levels of tRNAs and 5S rRNA in young panicles (0-1 cm) of GLA4 and NIL-gl6. OsUBQ5 was used as the control and the values of expression levels in GLA4 were set to 1. Values are given as the mean  $\pm$  sd. Student's t test significant difference,\*P < 0.05, \*\*P<0.01 and \*\*\*P<0.001; ns, not significant.

Intriguingly, we found that a series of genes including *TAW1*, *OsMADS22*, *OsMADS47*, and *OsMADS55*, which function in regulatory pathways that control inflorescence architecture by delaying the developmental meristem phase transition from inflorescence meristem to spikelet meristem (Yoshida et al., 2013), were significantly up-regulated in NIL-gl6, and these expression changes were confirmed by RT-qPCR (Supplemental Fig. S9F). High expression of these five genes resulted in more secondary branches and spikelets, which was consistent with the increased grain number in NIL-gl6 compared with GLA4. These results imply that the null gl6 mutation might affect grain number by extending the activity of the inflorescence meristem to produce more seeds in NIL-gl6.

# Natural Variation in the GL6 Gene

To investigate the variation of *GL6* in diverse rice germplasms, 67 accessions from the rice pan-genome dataset (Zhao et al., 2018b) were selected alongside additional *GL6* sequences of 52 wild rice (*O. rufipogon*) accessions (Huang et al., 2012). The resulting 119 rice accessions consisted of 54 cultivated rice and 65 wild rice accessions (Supplemental Table S2). Although the null *gl6* allele was from wild rice W1943, comparisons

of nucleotide diversity and neutrality testing between cultivars and wild rice revealed that *GL6* was not a locus targeted by human selection during domestication (Supplemental Table S3).

We then analyzed the *GL6* open reading frame (ORF) to mine other natural variations in the *GL6* gene, and a total of 20 polymorphic sites were observed, including 16 SNPs and four insertions or deletions (InDels). Among these variations, 13 SNPs were synonymous mutations, whereas no other variety except W1943 carried the identical SNP (premature stop codon), T allele, or T/C allele. The remaining four InDels and two missense mutation SNPs were located in the last

exon of *GL6*. A total of 19 haplotypes based on these variations were generated and named H1 to H19 (Fig. 7A; Supplemental Table S4). Most of the *japonica* varieties showed no differences from the Nipponbare genome, belonging to the H1 haplotype, whereas the *indica* varieties were distributed among several haplo-types. Interestingly, 60% of the wild rice and most *aus* accessions, such Kasalath, shared one identical insertion (+595) in the last *GL6* exon, which resulted in an insertion of three to six histidines. The geological distribution of these accessions originated mainly from India, indicating that this insertion was fixed in this region (Fig. 7B). However, whether the Kasalath allele



**Figure 7.** Natural variation in the *GL6* ORF. A, Haplotype network of *GL6* ORF among 119 rice accessions. Each circle represented one of the 19 different haplotypes and circle size was proportional to haplotype frequency. Different colors refer to different rice subpopulations. The red dashed square encloses rice lines with the same insertion (+595), and the purple dashed square refers to W1943. B, Comparisons of distribution among accessions with the insertion (+595).

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and other InDels have effects on the determination of grain size needs more investigation in the future.

# DISCUSSION

PLATZ family proteins are a class of plant-specific transcription factors with widespread distribution in dicots, monocots, mosses, and algae. All PLATZ family members share a conserved PLATZ domain that is  $\sim$ 82 amino acids in length and consists of two noncanonical zinc finger domains (Nagano et al., 2001; Wang et al., 2018). The first PLATZ gene was isolated from pea (*Pisum sativum*), which binds to the A/T-rich sequence to negatively regulate the enhancer element of the pea plastocyanin gene (petE; Nagano et al., 2001). At present, only a few PLATZ genes have been isolated. Arabidopsis (Arabidopsis thaliana) AtPLATZ1 and AtPLATZ2 were confirmed to enhance seed desiccation tolerance (González-Morales et al., 2016). The maize (Zea mays) PLATZ protein FL3 interacts with RNA polymerase III for biogenesis of tRNA and 5S rRNA to regulate endosperm storage filling (Li et al., 2017). Moreover, AtORE15, an Arabidopsis ortholog of GL6, was identified recently as being involved in the regulation of leaf growth and suppression of senescence via cooperation with the growth regulating factor/growth regulating factorinteracting factor regulatory pathway (Kim et al., 2018).

There are 15 PLATZ genes in rice, yet there has been little functional characterization of rice PLATZ genes. In this study, we first identified the rice PLATZ gene *GL6*, which affects grain length and yield in rice. *GL6* positively regulates cell division to increase cell numbers of the spikelet hull, resulting in larger grains.

RNAPIII synthesizes various small noncoding RNAs that are essential for general biological activities. Dysregulation of the RNAPIII machinery results in multiple reported instances of reduced cellular and organismal growth (Dauwerse et al., 2011; Soprano et al., 2013, 2017; Borck et al., 2015; Johnson et al., 2016). However, the regulation of RNAPIII transcriptional activity remains poorly understood in plants. Our data show that GL6 interacts with two critical RNAPIII subunits, RPC53 and TFC1, to regulate tRNAs and 5S rRNA, indicating that rice GL6 might have a similar function as that of the maize PLATZ protein FL3.

Seed size is crucial for evolutionary fitness in plants. Considering rice yield potential, there is a balance between panicle architecture and grain size. Larger grains are often associated with reduced grain number, and smaller grain size indicates that more seeds are produced (Guo et al., 2018). For example, the rare allele *gw2* increases grain size and weight but also reduces grain number per panicle (Song et al., 2007). Similarly, in our study, we observed that NIL-*gl6* and GLA4-*gl6*<sup>CRISPR</sup> lines that had shorter grains showed increased grain number per panicle compared to GLA4, consistent with the observed dramatic reduction in grain number per panicle in NIL-CP lines that had large grains. These phenotypes demonstrate the negative correlation between grain size and grain number due to a trade-off between inflorescence and spikelet development.

In conclusion, we identified a new QTL, *GL6*, which functions in the RNAPIII transcription machinery to affect rice grain length and number. Although the detailed mechanism by which GL6 regulates expression of tRNAs and 5S rRNA is still unclear, our data provided here indeed contribute toward understanding this process. The extended regulatory pathway surrounding GL6 remains to be investigated in further detail. Moreover, the exploitation of *GL6* may be a potential approach to manipulate the molecular balance between grain size and number for the development of elite rice varieties with improved grain productivity.

# MATERIALS AND METHODS

#### Plant Materials and Trait Measurement

BIL219 (small grain) and GLA4 (large grain) were used as two parents for QTL mapping, and Nipponbare was used for transgenic confirmation. All rice plants were cultivated under field conditions with transplant spacing of  $20 \times 20$  cm in Shanghai and Hainan, China. The measurement of grain yield-related field agronomic traits was conducted with edge lines excluded. Plant height, panicle morphology, and grain number were obtained from the main culm. Grain yield per plant and tiller number were measured from the whole plant. Fully filled dry grains were used for measuring grain length, width, and weight by image analysis method provided with SC-E software (Hangzhou Wanshen Detection Technology Co., Ltd.). All trait measurements were repeated at least 3 times.

#### Fine Mapping of GL6

The backcross inbred line BIL219 was selected to carry out the backcross with GLA4 to generate the  $BC_1F_2$  population, which contained 288 lines. Then we performed marker assisted selection on each substitution segment to purify the genetic background. Further fine mapping using 2,181  $BC_1F_5$  lines narrowed the *GL6* locus down to a 6.1-kb region between markers GL4350 and GL4411.

#### Primers

All primers used in this study are listed in Supplemental Table S5.

#### **Transgene Constructs and Plant Transformation**

The entire 14365-bp *GL6* genomic region was digested with *SacI* from GLA4 bacterial artificial chromosome clone osigba0159g02 and then inserted into the binary vector pCAMBIA1301 to generate the complementation construct. The gene editing constructs of *GL6* via CRISPR/Cas9 were designed as previously described (Ma et al., 2015). All these constructs were introduced into *Agrobacterium tumefaciens* strain EHA105 and subsequently transferred into Nipponbare, GLA4, and NIL-*gl6* by *Agrobacterium*-mediated transformation. More than 10 independent transgenic lines were generated, respectively. All analyzed phenotypes were measured in the T2 generation of transgenic plants.

#### Scanning Electron Microscopy

Mature seeds were first cleaned ultrasonically several times to remove epidermal hairs and dust. The samples were then dried in a critical point drier and coated with gold sputter. For glume cell observation, the outer surfaces of the spikelet glumes were observed by scanning electron microscope (Hitachi). Cell size and cell number were calculated along the longitudinal axis.

#### **Phylogenetic Analysis**

Protein sequences of PLATZ family members in rice and other organisms were obtained by BLAST from the National Center for Biotechnology Information database. Multiple sequence alignments of protein were performed using the ClustalW program. The phylogenetic tree of aligned sequence was constructed by MEGA7 using a neighbor-joining tree with 1000 bootstrapped replicates.

#### Neutrality Test

Multiple sequences of *GL6* genomic DNA were aligned with ClustalW. Nucleotide diversity and Tajima's *D* test were calculated and performed using DnaSP version 6.12.03 (Rozas et al., 2017).

#### Haplotype Network

Multiple sequences of *GL6* ORF were aligned with ClustalW. Haplotypefrequency data were processed with DnaSP version 6.12.03 and visualized Median-joining networks were generated by PopART with some modifications (each continuous InDel was considered as one site; Leigh et al., 2015).

# Yeast Two-Hybrid Assays

For the two-hybrid assay, the full-length coding region of *GL6* was amplified and fused in frame with the GAL4 DNA-binding domain via cloning into *pGBKT7* DNA-BD vector as the bait plasmid (Clontech). The entire coding regions of *GL6*, *OsRPC53*, *OsBRF1*, and *OsTFC1* were introduced into the *pGADT7* AD vector (Clontech) prey vector. The resulting constructs were then transformed into yeast strain AH109. The cotransformants were diluted (1, 1/10, 1/100, 1/1000) and spotted on control medium (SD/-Trp/-Leu) and selective medium (SD/-Trp/-Leu/-His/-Ade) and incubated at 30°C for 3 d.

# **BiLC Assay**

The coding-region sequence of *GL6* was cloned to the N-terminal luciferase (Luc) fusion vector JW771-NLUC, and *OsRPC53*, *OsBRF1*, and *OsTFC1* were cloned respectively to the C-terminal Luc fusion vector JW772-CLUC. The BiLC assay procedure was performed as previously described (Gou et al., 2011). *A. tumefaciens* [strain GV3101 (pSoup-p19)] transformants containing the testing split LUC fusion constructs were cotransfected into *Nicotiana benthamiana* leaves via infiltration, and LUC activity was captured using a cooled CCD-image system (Tanon 5200) with injecting 0.94 mm luciferin (cat. no. 122799; PerkinElmer) after growing for 48 h under 16 h light/d.

#### **RNA Extraction and RT-qPCR Analysis**

Total RNA was extracted using the Trizol reagent (Invitrogen) according to the manufacturer's instructions. Then 500 ng of total RNA was used to synthesize first-strand cDNA using ReverTra Ace qPCR RT Master Mix with gDNA Remover (Toyobo). Real-time PCR was performed on the Applied Biosystems QuantStudio 5 PCR system with diluted cDNA was used as a template using THUNDERBIRD SYBR qPCR Master Mix (Toyobo). Rice gene *UBQ5* (Os01g0328400) was used as the internal control to normalize all data. Each set of experiments was repeated three times.

#### **RNA In Situ Hybridization**

Fresh young panicles of GLA4 and NIL-gl6 were collected and fixed in formaldehyde-alcohol-acetic acid solution at 4°C overnight, dehydrated by series ethanol procedures, and embedded in paraplast. The tissues were sliced into 8- $\mu$ m sections with a microtome (Leica). The gene-specific region of *GL6* was then amplified from FL-cDNA and used to generate digoxigenin-labeled RNA probes (Roche). In situ hybridization was performed as described (Luo et al., 1996).

# **RNA-Seq Analysis**

Total RNA was extracted from young GLA4 and NIL-gl6 inflorescences (<1 cm) with two biological replicates using the Trizol reagent (Invitrogen) according to the manufacturer's instructions. After treatment with DNase, the mRNA was purified using the NEBNext Poly(A) mRNA Magnetic Isolation Module (E7490; New England Biolabs). Libraries were synthesized using the NEBNext Ultra II RNA Library Prep Kit for Illumina (E7770; New England

Biolabs) and sequenced on an Illumina HiSeq2500. A total of 269 million 150-bp paired-end reads were generated, yielding 40.34 Gb raw reads. After trimming of Illumina adaptors and low-quality reads, the filtered clean reads were aligned to the MSU version 7 genome assembly (http://rice.plantbiology.msu. edu) using HISAT2 (Kim et al., 2015). The aligned read files were sorted and indexed by SAMtools (Li et al., 2009) and reads of each sample were then used to calculate raw counts for each gene and transcript using the function SummarizeOverlaps within the GenomicAlignments package. The sample-to-sample distances were presented by principal-components analysis, which was based on the transcript count of each sample. DESeq2 software packages (bioconductor.org/) were used to detect DEGs with the threshold of genes with Benjamini-Hochberg-adjusted *p*-values <0.05 and absolute log2 fold change >0.6. GO enrichment analyses were performed using agriGO V2.0 (Tian et al., 2017).

# **Statistical Analysis**

Statistical analyses were carried out using Excel 2010 with two-tailed Student's t test for comparison of two groups and R package "agricolae" with Duncan's multiple-range tests for multiple mean comparisons.

#### Accession Numbers

Sequence data from this article can be found in the GenBank/EMBL data libraries under accession numbers MK959365 (*GL6* cDNA from W1943), XM\_015781518.2 (*OsRPC53*), XM\_015766861.2 (*OsTFC1*), and XM\_015785297.2 (*OsBRF1*).

# Supplemental Data

The following supplemental materials are available.

- Supplemental Figure S1. Graphical genotype and phenotype of BIL219.
- Supplemental Figure S2. Sequence alignment of *GL6* in GLA4 and NILgl6.
- Supplemental Figure S3. Relationship between different *GL6*-type and grain length in W1943.
- Supplemental Figure S4. GL6 positively regulates grain length.
- **Supplemental Figure S5.** Suppression of *GL6* in Nipponbare results in smaller grains.

Supplemental Figure S6. RNA in situ hybridization analysis.

- Supplemental Figure S7. Amino acid sequence alignments of GL6 and its homologs in various species.
- Supplemental Figure S8. Phylogenetic analysis of GL6 and other related PLATZ proteins.
- Supplemental Figure S9. GO enrichment analysis of RNA-seq DEGs.
- Supplemental Table S1. The DEGs and GO enrichment in GLA4 and NILgl6.

Supplemental Table S2. The list of 119 rice accessions in the collection.

- Supplemental Table S3. Nucleotide diversity and Tajima's D test.
- **Supplemental Table S4.** Sequence variation and distribution of *GL6* ORF haplotypes among 119 rice germplasms.
- Supplemental Table S5. List of primers used in this study.

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