

The endosperm-specific transcription factor TaNAC019 regulates glutenin and starch accumulation and its elite allele improves wheat grain quality

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Abstract

In wheat (*Triticum aestivum* L.), breeding efforts have focused intensively on improving grain yield and quality. For quality, the content and composition of seed storage proteins (SSPs) determine the elasticity of wheat dough and flour processing quality. Moreover, starch levels in seeds are associated with yield. However, little is known about the mechanisms that coordinate SSP and starch accumulation in wheat. In this study, we explored the role of the endosperm-specific NAC transcription factor TaNAC019 in coordinating SSP and starch accumulation. TaNAC019 binds to the promoters of *TaGlu-1* loci, encoding high molecular weight glutenin (HMW-GS), and of starch metabolism genes. Triple knock-out mutants of all three *TaNAC019* homoeologs exhibited reduced transcript levels for all SSP types and genes involved in starch metabolism, leading to lower gluten and starch contents, and in flour processing quality parameters. TaNAC019 directly activated the expression of HMW-GS genes by binding to a specific motif in their promoters and interacting with the *TaGlu-1* regulator TaGAMYB. TaNAC019 also indirectly regulated the expression of *TaSPA*, an ortholog of maize *Opaque2* that activates SSP accumulation. Therefore, TaNAC019 regulation of starch- and SSP-related genes has key roles in wheat grain quality. Finally, we identified an elite allele (*TaNAC019-BI*) associated with flour processing quality, providing a candidate gene for breeding wheat with improved quality.

IN A NUTSHELL

Background: Wheat flour is used to produce a wide range of products, such as leavened and unleavened breads, steamed buns, noodles, pancakes, cakes, and cookies. In wheat (*Triticum aestivum* L.) breeding, efforts have focused on improving both grain yield and quality. The content and composition of seed storage proteins (SSPs) are the target for quality improvements, as they determine the elasticity of wheat dough and flour processing quality. Starch levels in seeds are themselves associated with yield. However, improving flour quality generally comes at the expense of yield, thus hindering the development of high-yielding wheat cultivars with good flour quality.

Question: We wanted to know which regulators coordinate SSP and starch accumulation in wheat grain. We explored the role of the endosperm-specific NAC transcription factor TaNAC019 in coordinating SSP and starch accumulation.

Findings: We determined that TaNAC019 binds to the promoters of *TaGlu-1* loci, encoding high molecular weight glutenins (HMW-GS), as well as starch metabolism genes. The genetic inactivation of *TaNAC019* by genome editing resulted in reduced transcript levels for all SSP types and genes involved in starch metabolism, leading to lower gluten and starch contents, as well as lower flour processing quality parameters. TaNAC019 directly activated the expression of HMW-GS genes by binding to a specific motif in their promoters and interacting with the *TaGlu-1* regulator TaGAMYb. TaNAC019 also indirectly regulated the expression of TaSPA, an ortholog of maize Opaque2 that activates SSP accumulation. Therefore, TaNAC019 regulation of starch- and SSP-related genes has key roles in wheat grain quality. Finally, we identified an elite allele (*TaNAC019-BI*) associated with flour processing quality, providing a promising candidate gene for breeding wheat with improved quality.

Next steps: Wheat breeders aim to improve grain yield and quality. Our work demonstrates that TaNAC019 is an excellent candidate for developing wheat cultivars with both high yield and good quality. The *TaNAC019-BI* allele represents a genetic marker that may be used in marker-assisted breeding programs.

Introduction

Bread wheat (*Triticum aestivum* L.) is one of the most widely cultivated and consumed crops worldwide. Wheat flour is used to produce a wide range of products, such as leavened and unleavened breads, steamed buns, noodles, pancakes, cakes, and cookies (Veraverbeke and Delcour, 2002). The end-use value of wheat flour mainly depends on the properties of seed storage proteins (SSPs), especially gluten proteins, which give wheat dough products their unique viscoelastic properties (Payne et al., 1987; Biesiekierski, 2017).

Gluten is a complex mixture of two types of proteins: polymeric glutenins and monomeric gliadins. Glutenins are subdivided into high molecular weight (HMW-GSs) and low molecular weight (LMW-GSs) glutenin subunits, both of which confer elasticity to dough. Although HMW-GSs account for only about 10% of gluten proteins, the variation in their composition explains up to 70% of the genetic variation in dough processing quality (Payne et al., 1987; Eagles et al., 2002; Liu et al., 2005). Compared with HMW-GSs, higher levels of LMW-GSs are present in the grain of bread wheat; the LMW-GS content is associated with dough resistance and extensibility (Dong et al., 2010). Gliadin content is also associated with dough extensibility (Shewry et al., 2002). The quantitative effects of individual HMW-GSs (Ragupathy et al., 2008; Li et al., 2015; Cho et al., 2017), as well as the ratios of HMW-GSs/LMW-GSs and HMW-GSs/gliadins, are relevant for different uses of dough (Veraverbeke and Delcour, 2002; Rasheed et al., 2014).

Wheat HMW-GSs are encoded by *TaGlu-1* loci on the long arms of homoeologous chromosomes 1A, 1B, and 1D. Each locus contains two tightly linked genes encoding X- and Y-type subunits, i.e. *TaGlu-1Ax*, *TaGlu-1Ay*, *TaGlu-1Bx*,

TaGlu-1By, *TaGlu-1Dx*, and *TaGlu-1Dy* (Galili and Feldman, 1985). LMW-GSs are encoded by *Glu-3* loci (i.e. *Glu-A3*, *Glu-B3*, and *Glu-D3*) located on the short arm of homoeologous group 1 chromosomes, and each locus contains multiple gene copies (Singh and Shepherd, 1988; Zhang et al., 2011; Lee et al., 2016). Gliadins are encoded by genes in two genomic regions: the *Gli-1* loci, located on the short arms of group 1 chromosomes, encoding γ -, δ -, ω -gliadins, and the *Gli-2* loci, located on the short arms of homoeologous group 6 chromosomes, encoding α -gliadins (Singh and Shepherd, 1988; Huo et al., 2018).

Conserved cis-regulatory motifs have been identified in the promoters of genes encoding SSPs (Makai et al., 2015), and a number of transcription factors (TFs) involved in SSP regulation have been characterized. Wheat storage protein activator (TaSPA) binds to a general control nondepressible 4 (GCN4)-like binding motif in the promoters of genes encoding glutenins and gliadins and activates their transcription (Albani et al., 1997; Ravel et al., 2014), whereas SPA heterodimerizing protein (SHP) suppresses *glutenin* gene expression (Boudet et al., 2019). TaFUSCA3, a B3-Superfamily TF that also interacts with TaSPA, activates *TaGlu-1Bx7* by binding to the RY-box motif in its promoter (Sun et al., 2017). The DNA-binding with one finger (Dof)-type prolamins-box binding factor (PBF) binds to the prolamins-box (P-box) in the promoter regions of wheat α -gliadin and LMW-GS genes (Dong et al., 2007) and binds to the *TaGlu-1By8* and *-1Dx2* loci, leading to promoter hypomethylation (Zhu et al., 2018). Finally, the GA-dependent MYB TF TaGAMYb activates HMW-GS gene expression by directly binding to the *TaGlu* promoter and by recruiting the GCN5-like histone acetyltransferase (HAT; Guo et al.,

2015). Despite the identification of these TFs, little is known about their functional interactions or the intricate mechanisms of the SSP regulatory pathway.

High baking quality is associated with the protein composition of wheat grain, but negatively correlated with high yield, which is itself associated with starch content (Rharrabti et al., 2001; Zorb et al., 2018). Although TFs that regulate the expression of genes encoding wheat starch biosynthetic enzymes have been reported (Kang et al., 2013; Kumar et al., 2018), how TFs regulate starch biosynthesis remains unknown. In addition, little is known, in any plant species, about how TFs simultaneously regulate SSP and starch accumulation, thus coordinating the balance between quality and yield.

A recent report indicated that ZmNAC128 and ZmNAC130, which belong to the NAC (NAM [No Apical Meristem]–ATAF [Arabidopsis Transcription Activation Factor]–CUC [Cup-shaped Cotyledons]) superfamily, act redundantly to modulate starch and protein accumulation during maize (*Zea mays*) endosperm development (Zhang et al., 2019). The NAC domain superfamily is one of the largest groups of plant-specific TFs. NAC TFs are involved in various developmental and growth processes, although they are mainly known for their roles in plant responses to biotic and abiotic stress (Olsen et al., 2005; Mathew and Agarwal, 2018). Rice (*Oryza sativa*) OsNAP (NAC-LIKE, ACTIVATED BY AP3/PI) and wheat NAM-B1 play indirect roles in regulating protein content in grains, by accelerating leaf senescence and nitrogen accumulation in aerial parts (Uauy et al., 2006; Liang et al., 2014). In addition, NAC genes that are specifically expressed in barley (*Hordeum vulgare*) and rice seeds may directly regulate seed size, but little is known about the underlying mechanism (Christiansen et al., 2011; Mathew et al., 2016).

In this study, we provide evidence that TaNAC019, a seed-specific NAC TF, regulates both SSP and starch accumulation in wheat grain. We explored the mechanism underlying TaNAC019-mediated transcriptional regulation of gene expression, in which TaNAC019 directly binds to and activates the expression of its target genes, including HMW-GS genes, the starch synthase gene *TaSSIIa*, and the sucrose synthase gene *TaSuSy1*. TaNAC019 also affects *TaSPA* expression, and physically interacts and cooperates with TaGAMYB to activate *TaGlu* expression. Finally, we identified a specific natural genetic variant of *TaNAC019* (*TaNAC019-B1*) that is associated with higher wheat processing quality, thus providing a promising candidate gene for wheat quality improvement in breeding programs.

Results

Identification of the endosperm-specific nuclear protein TaNAC019, which binds to the promoters of *glutenin* genes

With the aim of identifying TFs that regulate the expression of *glutenin* genes, we performed a yeast one-hybrid assay using the *TaGlu-1Dy* promoter as bait against a wheat cDNA

expression library. Among the 347 positive clones, three matched TraesCS3A02G077900, as revealed by Basic Local Alignment Search Tool for DNA (BLASTN) analysis; this gene encodes a member of the wheat NAC TF family. Phylogenetic analyses of wheat NAC TF family members indicated that TraesCS3A02G077900 is an ortholog of the seed-specific barley gene *HvNAC019* (Christiansen et al., 2011; Borrill et al., 2017; Guerin et al., 2019), whose function is still unknown. However, reciprocal BLAST analysis (Borrill et al., 2017; Guerin et al., 2019) indicated that TraesCS3A02G077900 is not an ortholog of the endosperm-specific rice NAC TF genes, Os01g01470 or Os01g29840, which are associated with seed size (Mathew et al., 2016), or their maize orthologs, ZmNAC128 and ZmNAC130, which directly regulate protein and starch accumulation (Zhang et al., 2019). Accordingly, we named TraesCS3A02G077900 as *TaNAC019*.

Bread wheat is an allohexaploid species whose genes usually have homoeologs in all three subgenomes (Glover et al., 2016). Among transcripts sharing high sequence similarity with TraesCS3A02G077900, the two loci TraesCS3B02G092800 and TraesCS3D02G078500 mapped to chromosome 3 of subgenomes B and D, respectively. The NAC proteins encoded by these three genes grouped within the same phylogenetic cluster, along with most of the wheat seed-specific NACs identified to date (Guerin et al., 2019; Supplemental Figure S1). We confirmed the genomic locations of the three genes using a set of Chinese Spring (CS) nullisomic–tetrasomic lines and homoeolog-specific primers (Supplemental Figure S2). The three homoeologous sequences were named *TaNAC019-A*, *TaNAC019-B*, and *TaNAC019-D*.

Although most NAC TFs are soluble proteins found in the nucleus, several NACs are membrane-bound TFs or localize to the cytoplasm (Mathew and Agarwal, 2018). Analysis of protein fusion constructs between TaNAC019 homoeologs and the green fluorescent protein (GFP), transiently expressed in *Nicotiana benthamiana* leaves, indicated that all three homoeologs are localized to the nucleus and cytoplasm (Supplemental Figure S3).

Reverse transcription-quantitative PCR (RT-qPCR) analysis showed that the *TaNAC019* homoeologs are expressed almost exclusively in the endosperm and that *TaNAC019-B* is expressed at higher levels than *TaNAC019-A* or *TaNAC019-D* (Figure 1, A). During kernel development, *TaNAC019-A/B/D* transcripts were undetectable until 12 days after pollination (DAP), and the highest transcript levels were detected at ~25 DAP (Figure 1, B). The *glutenin* gene transcripts started to accumulate before *TaNAC019* transcripts and reached their highest peak at 25 DAP, as measured using a primer combination that detects the total mRNA levels of the four *TaGlu* genes expressed in the wheat cultivar Nongda3338 (i.e. *TaGlu-1Bx/1By* and *TaGlu-1Dx/1Dy*; *TaGlu-1Ax/1Ay* loci are not expressed in this cultivar; Supplemental Figure S4 and Figure 1, B).

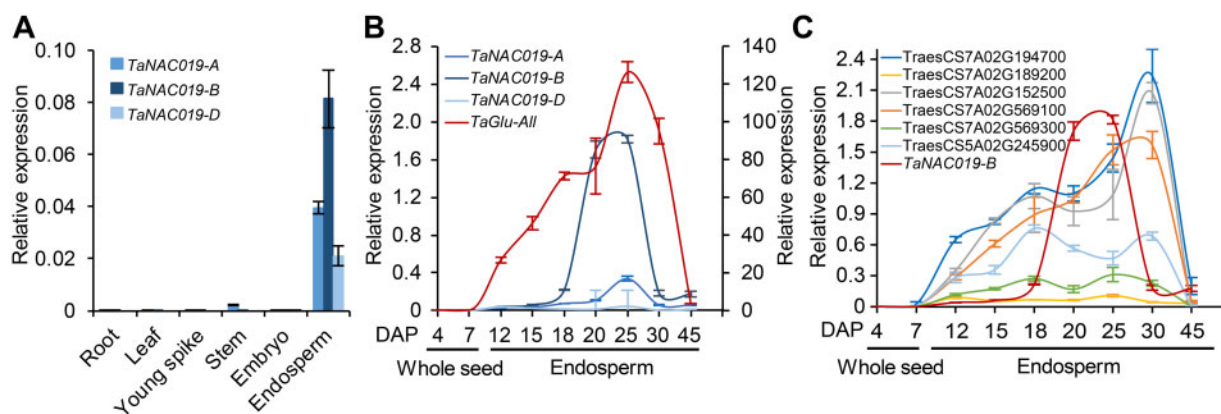


Figure 1 TaNAC019 expression in wheat seeds. A, Relative transcript levels of the three *TaNAC019* homoeologs in the indicated tissues, as determined by RT-qPCR. Data were normalized to wheat *TaACTIN*, shown as mean \pm SD ($n = 3$). B, Relative transcript levels of the three *TaNAC019* homoeologs (y -axis on the left) and *TaGlu-All* (y -axis on the right), estimated using total RNA extracted from whole seeds from 4 to 7 DAP and endosperm (from 12 to 45 DAP). *TaGlu-All* transcript levels were measured using a primer combination allowing mRNA amplification of four *TaGlu* genes (i.e. *TaGlu-1Bx*, *TaGlu-1By*, *TaGlu-1Dx*, and *TaGlu-1Dy*). C, As described in B; primer combinations specific for other wheat seed specific NACs (Supplemental Figure S1) were employed. For clarity, only data obtained for the homoeolog located on the A subgenome for each locus are shown, as mean \pm SD ($n = 3$).

Immunoblot analysis with an antibody that recognizes the three TaNAC019 proteins validated the expression patterns observed by RT-qPCR (Supplemental Figure S5, A and C). These results suggest that TaNAC019 is positively associated with the expression of *TaGlu* genes. A comparison of *TaNAC019* to other wheat seed-specific NAC transcripts (Supplemental Figure S1) indicated that they have different expression patterns during seed development, with most other transcripts accumulating to higher levels during two distinct developmental stages, while *TaNAC019* reached its highest expression level from 20 to 25 DAP and *TraesCS7A02G0569100* was expressed early and reached its highest expression level from 25 to 30 DAP (Figure 1, C). These results suggest that different seed-specific TaNACs might have distinct, or not fully overlapping, functions.

TaNAC019 regulates glutenin accumulation, gluten formation, and wheat processing quality

To investigate the role of TaNAC019, we generated knock-out mutants in the wheat cultivar Fielder background via CRISPR (clustered regularly interspaced short palindromic repeats)/Cas9 (CRISPR-associated protein 9) gene editing. To simultaneously knock out all three *TaNAC019* homoeologs, we designed a single guide RNA (sgRNA) that targets a conserved region within the first exon of these three genes (Supplemental Figure S6). We obtained two independent lines (*nac019-1* and *nac019-2*) with frameshift mutations leading to premature termination in all three *TaNAC019* homoeologs. TaNAC019 protein was undetectable in lines *nac019-1* and *nac019-2*, as revealed by immunoblot analysis (Supplemental Figure S5, B).

Since TaNAC019 is a putative regulator of *TaGlu-1Dy*, we performed reverse-phase high-performance liquid chromatography (RP-HPLC) to measure HMW-GS contents in mature seeds. The levels of all subunits including TaGlu-1Bx, TaGlu-1By, TaGlu-1Dx, and TaGlu-1Dy were reduced in the

nac019 mutants compared with the wild type (Figure 2, A). This difference was confirmed by protein gel analysis, followed by silver staining (Supplemental Figure S7). The LMW-GS levels were also reduced in the *nac019* mutants (Figure 2, B), whereas gliadin levels in the mutants were similar to those in the wild type (Figure 2, C). In addition, the loss of TaNAC019 function was associated with a decrease in dry gluten contents (Figure 2, D) and SDS sedimentation volume (Figure 2, E), an indicator of processing and end-product quality. Starch levels and accumulation of starch granules were also reduced in the *nac019* mutants (Figure 2, K and L). These results indicate that TaNAC019 regulates both glutenin and starch accumulation.

The *nac019* mutants had smaller kernels than the wild type. Indeed, grain width and length (Figure 2, F and G), as well as thousand-grain weight (TGW; Figure 2, H) were lower in the mutants. We measured the cell size and cell number of developing endosperm from *nac019* mutants and wild type, and the results indicated that cell size decreased while cell number increased in *nac019* mutants when compared with the wild type (Figure 2, I and J and Supplemental Figure S8). Conversely, and in agreement with the seed-specific expression of *TaNAC019*, no visible phenotypic differences were detected in *nac019* mutant mature plants relative to wild type, including plant height, tiller number, spike number, and flowering time (Supplemental Figure S8). These results indicate that TaNAC019 regulates glutenin and starch accumulation, gluten formation, and seed size and weight, thus influencing the processing quality of wheat flour.

TaNAC019 regulates the expression of various genes, including genes involved in SSP and starch accumulation

We identified the downstream targets of TaNAC019 by transcriptome deep sequencing (RNA-seq), using RNA extracted

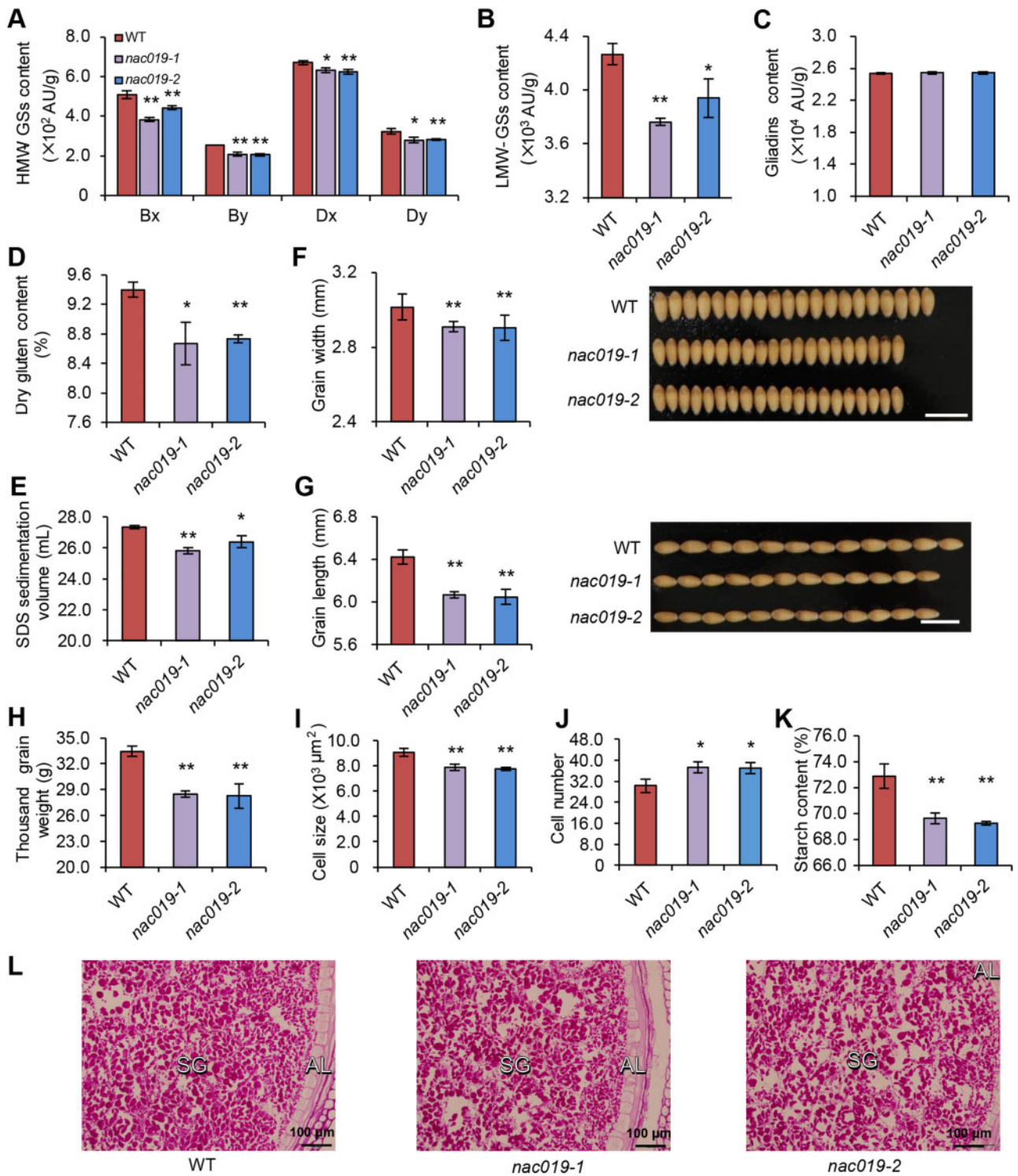


Figure 2 Seed phenotypes of wheat *nac019* knock-out lines. A–D, Results from RP-HPLC analysis of HMW-GS (A), LMW-GS (B), and gliadin (C) contents and from measuring dry gluten content (D) and SDS sedimentation volume (E). F–K, Seed size-related phenotypes of *nac019* mutants: grain width (F), grain length (G), TGW (H), cell size (I), cell number (J), and starch content (K). Three repeats were carried out for each sample to measure the total amounts of HMW-GSs, LMW-GSs, gliadins, dry gluten content, SDS sedimentation values, and starch content. For kernel weight, grain width, and grain length, nine replicates were used, each with approximately 25 g seeds. The single and double asterisks represent significant differences compared with WT and determined by Student's *t* test at $P < 0.05$ and $P < 0.01$, respectively. L, Starch granules analysis of *nac019* mutants and wild type. SG, starch granules; AL, aleurone. Three seeds were used for histological analysis for starch granules staining.

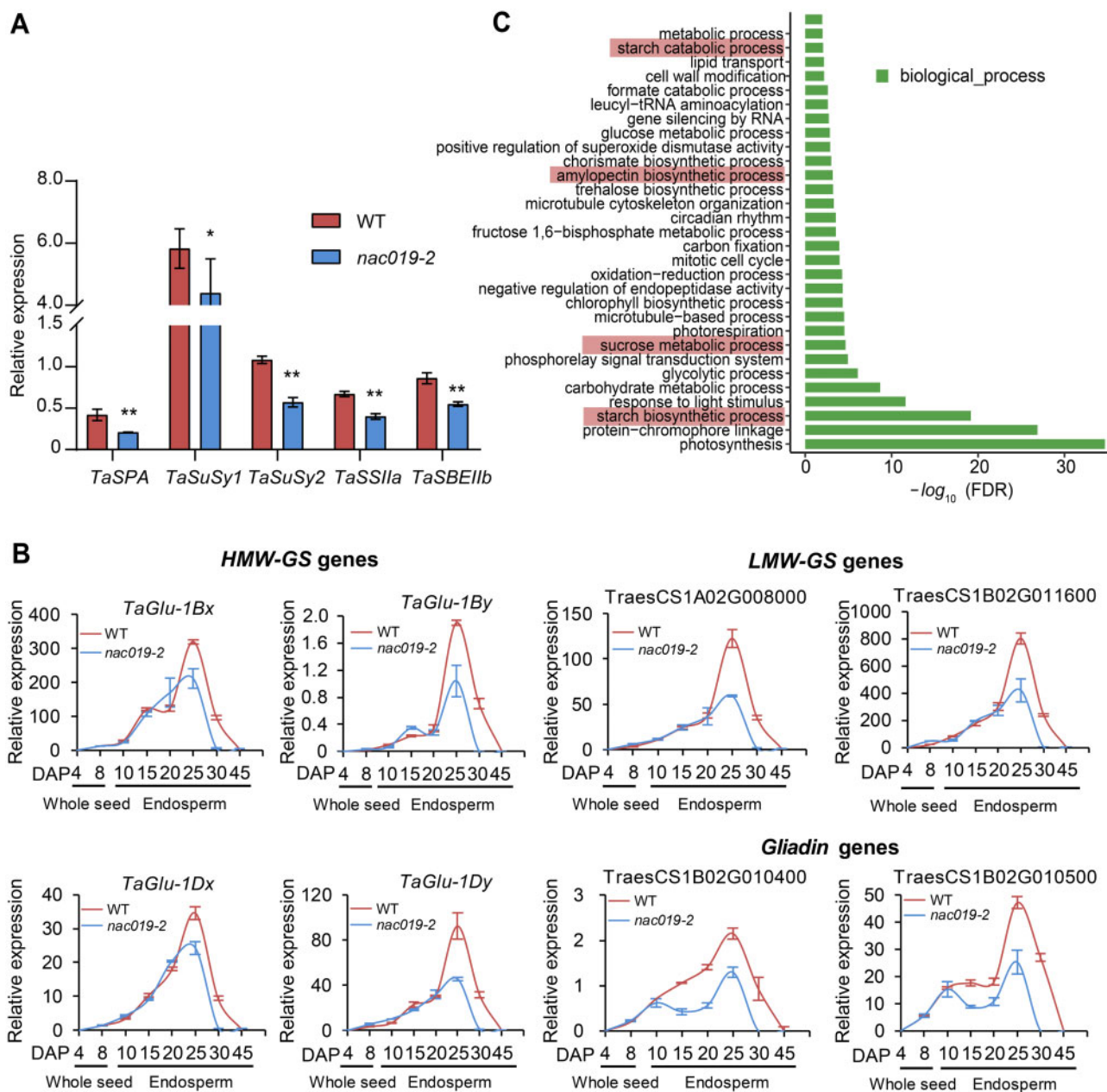


Figure 3 Genes and processes regulated by TaNAC019. A, RT-qPCR analysis of *TaSPA*, *TaSBEIIb* (starch branching enzyme IIb), *TaSuSy1* (sucrose synthase), *TaSuSy2* (sucrose synthase), and *TaSSIIa* (starch synthase 2a) expression in *nac019-2* and wild-type endosperm at 25 DAP. RT-qPCR data were normalized to wheat *TaACTIN*. The data from three replicates are shown as \pm sd. The single and double asterisks represent significant differences determined by Student's *t* test at $P < 0.05$ and $P < 0.01$, respectively. B, Expression of the indicated HMW-GS, LMW-GS, and gliadin genes during seed development, analyzed in *nac019-2* and wild type. RT-qPCR data were normalized to wheat *TaACTIN*, and each bar in the graph corresponds to the mean value of three data points from one representative biological replicate. C, GO analysis of downregulated genes in the *nac019-2* transcriptome compared with wild type. *P*-values were adjusted by the Benjamini–Hochberg correction, and only significant categories (FDR < 0.05) are displayed. The red rectangles indicate the pathways related to starch metabolism.

from wild-type Fielder and *nac019-2* endosperms at 25 DAP. Among the 36,456 genes detected by RNA-seq, a total of 2,260 genes were downregulated and 3,580 were upregulated in *nac019-2* compared with wild-type Fielder (Supplemental Data Set S1). Among the downregulated genes, we identified the three TaNAC019 homoeologs, indicating that the CRISPR/Cas9 gene editing affected not only their protein biosynthesis, but also their mRNA production or stability.

Importantly, *TaSPA* was also downregulated in the *nac019-2* mutant (Figure 3, A and Supplemental Data Set S1), indicating that although TaNAC019 binds directly to *TaGlu-1Dy* in the yeast one-hybrid system, it might also regulate SSP accumulation indirectly by modulating *TaSPA* expression. Indeed, all four *TaGlu-1* loci expressed in Fielder (*TaGlu-1Bx*, *TaGlu-1By*, *TaGlu-1Dx*, and *TaGlu-1Dy*), as well as all 17 genes encoding LMW-GS and 66 of 78 genes encoding gliadins,

were downregulated in *nac019-2* (Supplemental Data Set S2).

Since RP-HPLC indicated that the levels of HMWs and LMW proteins were reduced in mature *nac019* seeds, whereas the levels of gliadins were not (Figure 2, B–D), we investigated the effects of the *nac019-2* mutation on SSP gene expression during seed development. Specifically, we used RT-qPCR to compare the transcript levels of the four *TaGlu-1* genes, two LMW-GS genes, and two *gliadin* genes in *nac019-2* versus the wild type. In agreement with the RNA-seq data, all of these genes were downregulated in 25 DAP endosperm from *nac019-2* compared with the wild type (Figure 3, B). However, while expression of the genes encoding HMW-GS and LMW-GS started to decrease after 20 DAP, *gliadin* gene expression started to decrease earlier during seed development. Overall, these results indicate that TaNAC019 activates expression of glutenin- and gliadin-encoding genes during specific stages of endosperm development. However, in mature seeds of the *nac019* knock-out mutants, the amounts of some SSPs, especially gliadins, were restored to wild-type levels, possibly due to the previously documented phenomenon of compensation between glutenin and gliadin levels that occurs when the level of one of these types of SSPs is altered (Scossa et al., 2008; Chen et al., 2019).

Gene ontology (GO) analysis revealed that the genes downregulated in *nac019* are highly enriched for starch and amylopectin biosynthetic processes, as well as sucrose metabolic and starch catabolic processes (Figure 3, C). In agreement with the reduced starch levels in *nac019*, the downregulated genes included 45 genes related to starch metabolism (Supplemental Figure S9 and Supplemental Data Set S3). The downregulation of four genes that are key factors related to sucrose and starch metabolism, including *TaSBE11b* (encoding starch branching enzyme IIb), *TaSuSy1* (encoding sucrose synthase), *TaSuSy2* (also encoding sucrose synthase), and *TaSSIIa* (encoding starch synthase IIa) was confirmed by RT-qPCR (Figure 3, A). Genes upregulated in *nac019* were enriched in GO terms associated with oxidation–reduction processes, chitin catabolic process, and regulation of transcription (Supplemental Figure S10). Together, these results indicate that TaNAC019 concomitantly regulates both SSP and starch accumulation.

TaNAC019 binds to a specific motif in its target genes and activates their expression

To explore the mechanism of TaNAC019-mediated regulation, we investigated the TaNAC019 binding motif in its target genes. We used 60 fragments covering a 1.6-kbp region upstream of the transcriptional start site (TSS) of *TaGlu-1Dy* to detect the DNA-binding ability of TaNAC019 by electrophoretic mobility shift assays (EMSAs). Four fragments were bound by a bacterially expressed glutathione S-transferase (GST)-TaNAC019-B recombinant protein, with regions P11 and P43 showing the strongest binding (Supplemental

Figure S11). We validated the binding of TaNAC019-B to the P11 region by yeast one-hybrid assay (Supplemental Figure S12). To identify the precise TaNAC019-binding site, we used a series of oligonucleotides spanning the P11 region harboring mutations as competitor probes in EMSA. The probes M3, M7, and M9 failed to prevent TaNAC019 binding, indicating that the nucleotides mutated in these probes are required for the binding of TaNAC019 to its targets (Figure 4, A). We then performed EMSAs using competitor probes with single-nucleotide mutations covering the region between M3 and M9. Comparison to the wild-type probe revealed that the binding sequence was [AT]NNNNN NNNNN[ATC][CG]A[CA]GN[ACT]A (Figure 4, B). To verify the specificity of this motif, we generated a P11 fragment harboring mutations in the predicted TaNAC019-binding site (MP11). TaNAC019-B recombinant protein failed to bind to this fragment in both yeast one-hybrid assays (Supplemental Figure S12) and EMSAs (Supplemental Figure S13).

We searched for the TaNAC019-binding sequence in the promoter regions (2.5 kbp upstream of the TSS) of the genes that were differentially expressed in *nac019-2* versus wild type. The TaNAC019-binding motif was found in 37.1% and 39.1% of down- and upregulated genes, respectively (Supplemental Data Set S1). Importantly, the *TaSPA* promoter lacked this motif. The TaNAC019-binding site was detected in all four HMW-GS genes expressed in Fielder, whereas only 3 out of 17 LMW-GS and 18 out of 69 *gliadin* genes contained this sequence (Supplemental Data Set S2). In addition, among the 48 downregulated genes involved in starch metabolism, 27 contained the TaNAC019-binding motif in their promoter regions (Supplemental Data Set S3). The binding of TaNAC019 to the *TaGlu-1Bx*, *TaGlu-1By*, *TaGlu-1Dx*, *TaSuSy1*, and *TaSSIIa* promoters was confirmed by EMSA (Supplemental Figure S13). We then assessed the transactivation of four *TaGlu* genes, *TaSuSy1*, and *TaSSIIa* by TaNAC019 in a dual luciferase transcriptional activity assay, which established that TaNAC019 activates the expression of these genes in vivo (Figure 4, C).

Overall, these findings suggest that TaNAC019 binds to and directly activates the expression of a subset of its targets, whereas other targets might be regulated via indirect mechanisms. Alternatively, TaNAC019 might bind to a subset of its targets at a different motif from that found in *TaGlu-1Dy*, or perhaps it binds to the same motif but located more than 2.5 kbp away from the TSS.

TaNAC019 and TaGAMyB physically interact and additively cooperate to activate *TaGlu* expression

NAC TF activity sometimes requires interactions with paralogous proteins (Mendes et al., 2013; Mathew et al., 2016) or other TFs (Xu et al., 2013). We therefore assessed whether TaNAC019 homoeologs can interact with one another using the yeast two-hybrid system. However, none of the proteins interacted with each other in yeast two-hybrid assays (Supplemental Figure S14). Subsequently, we tested the

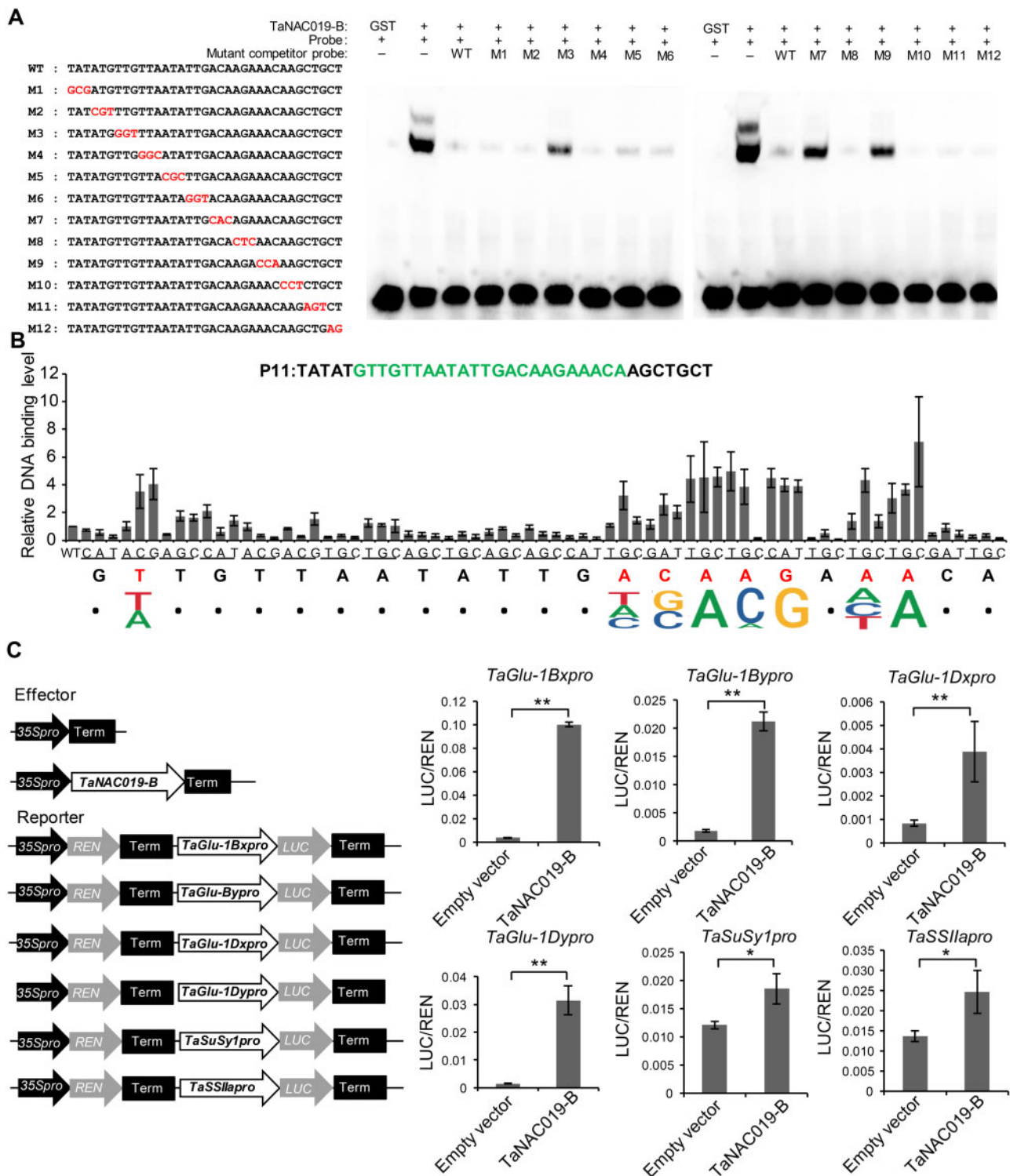


Figure 4 TaNAC019 binds to a specific motif in the *TaGlu* promoters. A, A three-nucleotide mutation series spanning the P11 oligonucleotide sequence was generated (highlighted in red) and used as competitors of the P11 probe in EMSAs with TaNAC019-B-GST recombinant protein. “+” and “-” indicate the presence and absence, respectively, of the corresponding probes and proteins. B, Mononucleotide mutations were introduced into the P11 core region (highlighted in green) and used as competitor probes in EMSAs. Each nucleotide of the core region was converted into the three other nucleotides. The average values of TaNAC019-B-binding levels from three independent replicates are shown on the y-axis. Nucleotides required for TaNAC019-B binding are indicated in red and the deduced binding motif is shown along the x-axis. C, Dual luciferase transcriptional activity assay to assess the ability of TaNAC019-B to transactivate target gene expression. Schematic diagrams of the effector and reporter plasmids. On the y-axis, LUC/REN indicates the ratio of the signal detected for firefly luciferase (LUC) versus *Renilla reniformis* luciferase (REN) activity. Error bars represent standard deviation for three replicates. The single and double asterisks indicate statistically significant differences estimated by Student’s *t* test at $P < 0.05$ and $P < 0.01$, respectively.

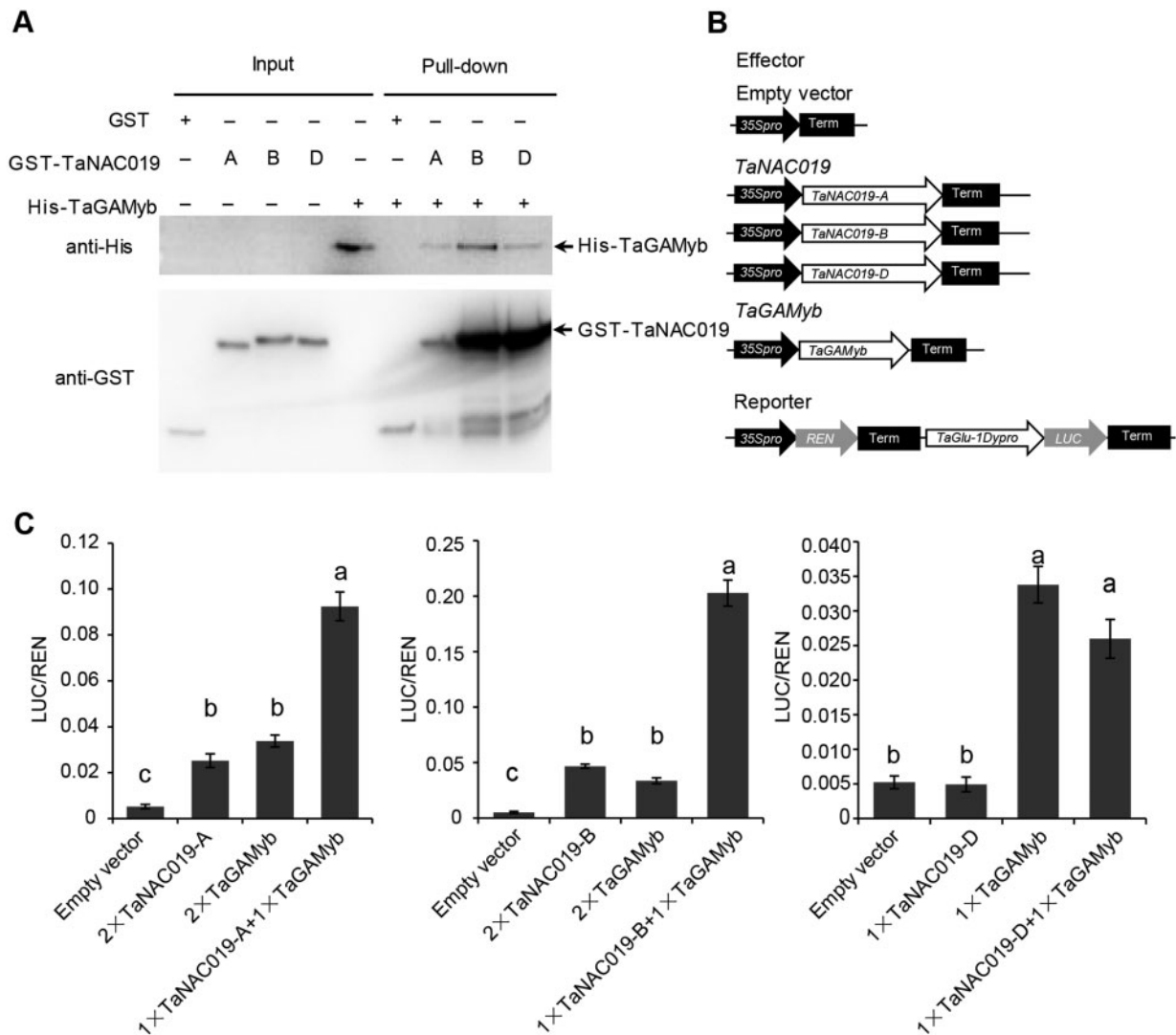


Figure 5 TaNAC019 and TaGAMyb physically interact and additively cooperate to drive *TaGlu-1Dy* promoter expression. A, Pull-down experiments to verify the interactions between the three TaNAC019 homoeologs and TaGAMyb. B and C, Dual luciferase transcriptional activity assays to assess the ability of TaNAC019 homoeologs and TaGAMyb to transactivate *TaGlu-1Dy* promoter expression. B, Schematic diagrams of the effector and reporter plasmids. C, On the y-axis, LUC/REN indicates the ratio of the signals detected for firefly luciferase (LUC) and *R. reniformis* luciferase (REN) activity. Error bars represent standard deviation for six replicates. The same letters above each bar indicate that the means did not differ significantly by one-way ANOVA and Tukey's multiple comparison test at $P < 0.05$. "2×" and "1×" indicate that the final concentration of Agrobacteria containing the corresponding construct was OD_{600} 0.2 and OD_{600} 0.1, respectively.

interaction between each TaNAC019 homoeolog and several TFs involved in SSP regulation, such as TaSPA, TaGAMyb, and the TaGCN5-like protein HAT. TaNAC019 interacted with TaGAMyb, but not with TaGCN5 or TaSPA in this assay (Supplemental Figure S14). The physical interaction between TaNAC019-A/B/D and TaGAMyb was confirmed in a pull-down assay using recombinant proteins fused with an epitope tag (Figure 5, A).

To investigate the biological significance of the interaction between TaNAC019 and TaGAMyb, we assessed their ability to transactivate the *TaGlu-1Dy* promoter in a dual luciferase transcriptional activity assay (Figure 5, B). TaNAC019-A and TaNAC019-B, but not TaNAC019-D, activated the *TaGlu-*

1Dy promoter (Figure 5, C). TaGAMyb also activated the promoter when expressed alone (Figure 5, C). Interestingly, co-expression of TaNAC019-A or TaNAC019-B with TaGAMyb led to a significant increase in *TaGlu-1Dy* promoter activation compared with the expression of the single effector constructs. Conversely, co-expressing TaNAC019-D and TaGAMyb activated the *TaGlu-1Dy* promoter to a level similar to that observed when TaGAMyb was expressed alone (Figure 5, C). Collectively, these results indicate that the three TaNAC019 homoeologs physically interact with TaGAMyb and that two of these TFs, TaNAC019-A and TaNAC019-B, cooperate and have additive effects with TaGAMyb in driving *TaGlu-1Dy* expression.

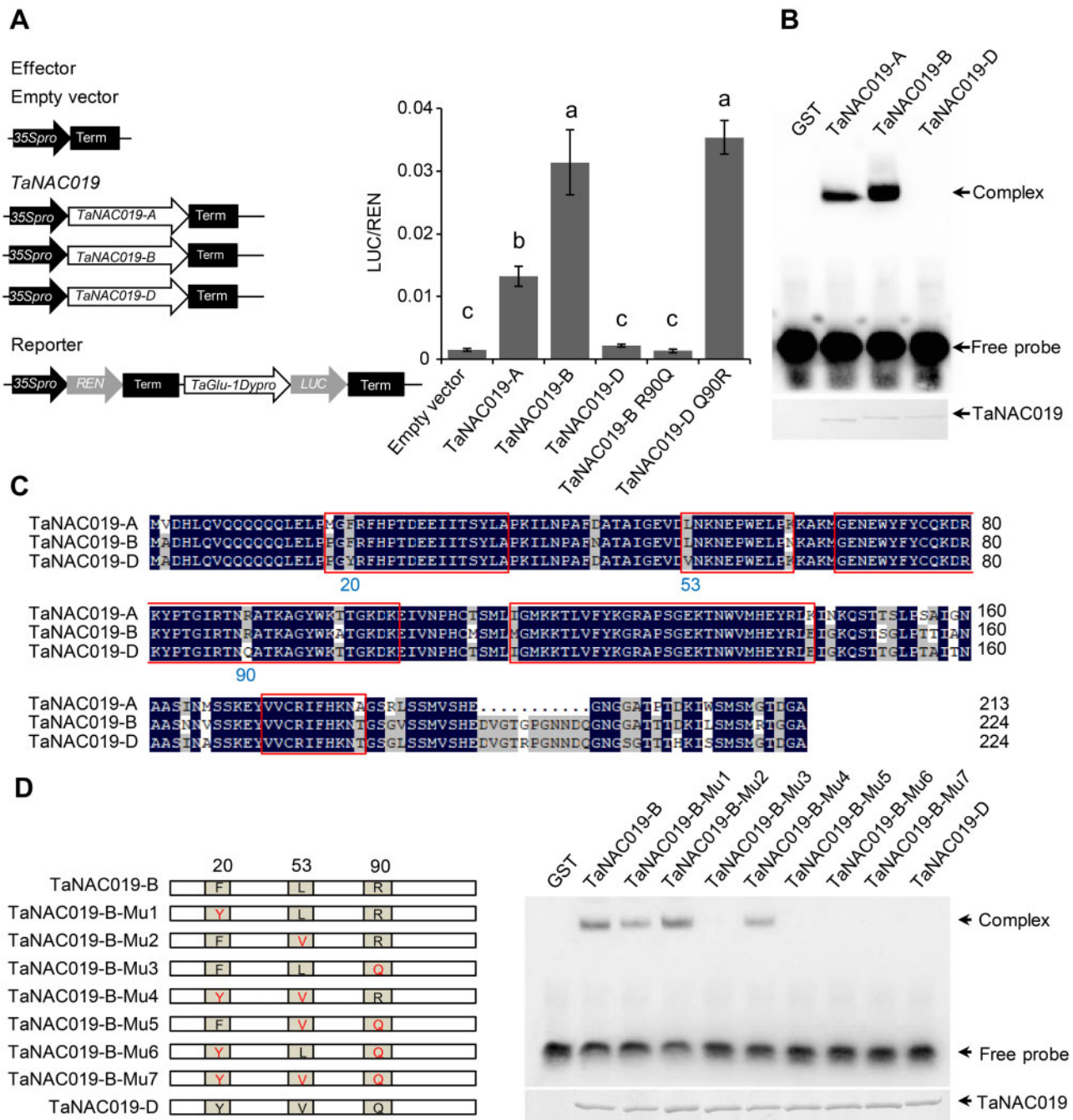


Figure 6 The three TaNAC019 homoeologs show functional divergence. A, Dual luciferase transcriptional activity assays to compare the ability of the three TaNAC019 homoeologs to transactivate *TaGlu-1Dy* promoter expression. R90Q and Q90R indicate the R to Q mutation at amino acid 90 of TaNAC019-B and the Q to R mutation at amino acid 90 of TaNAC019-D, respectively. On the y-axis, LUC/REN indicates the ratio of the signals detected for firefly luciferase (LUC) and *R. reniformis* luciferase (REN) activity. Error bars represent standard deviation for six replicates. The same letters above each bar indicate that the means did not differ significantly by one-way ANOVA and Tukey's multiple comparison test at $P < 0.05$. B, EMSAs to compare *TaGlu-1Dy* promoter binding between the three TaNAC019 homoeologs. C, Amino acid sequence alignment of the three TaNAC019 homoeologs. The core NAC DNA-binding domains are highlighted with red rectangles. The amino acids where TaNAC019-D differs from TaNAC019-A and TaNAC019-B at positions 20, 53, and 90 are labeled. D, The TaNAC019-B sequence was mutated to generate proteins carrying different combinations of the three amino acid residues characterizing TaNAC019-D, and each GST-fused recombinant protein was used in EMSAs to assess the binding of these proteins to the *TaGlu-1Dy* promoter.

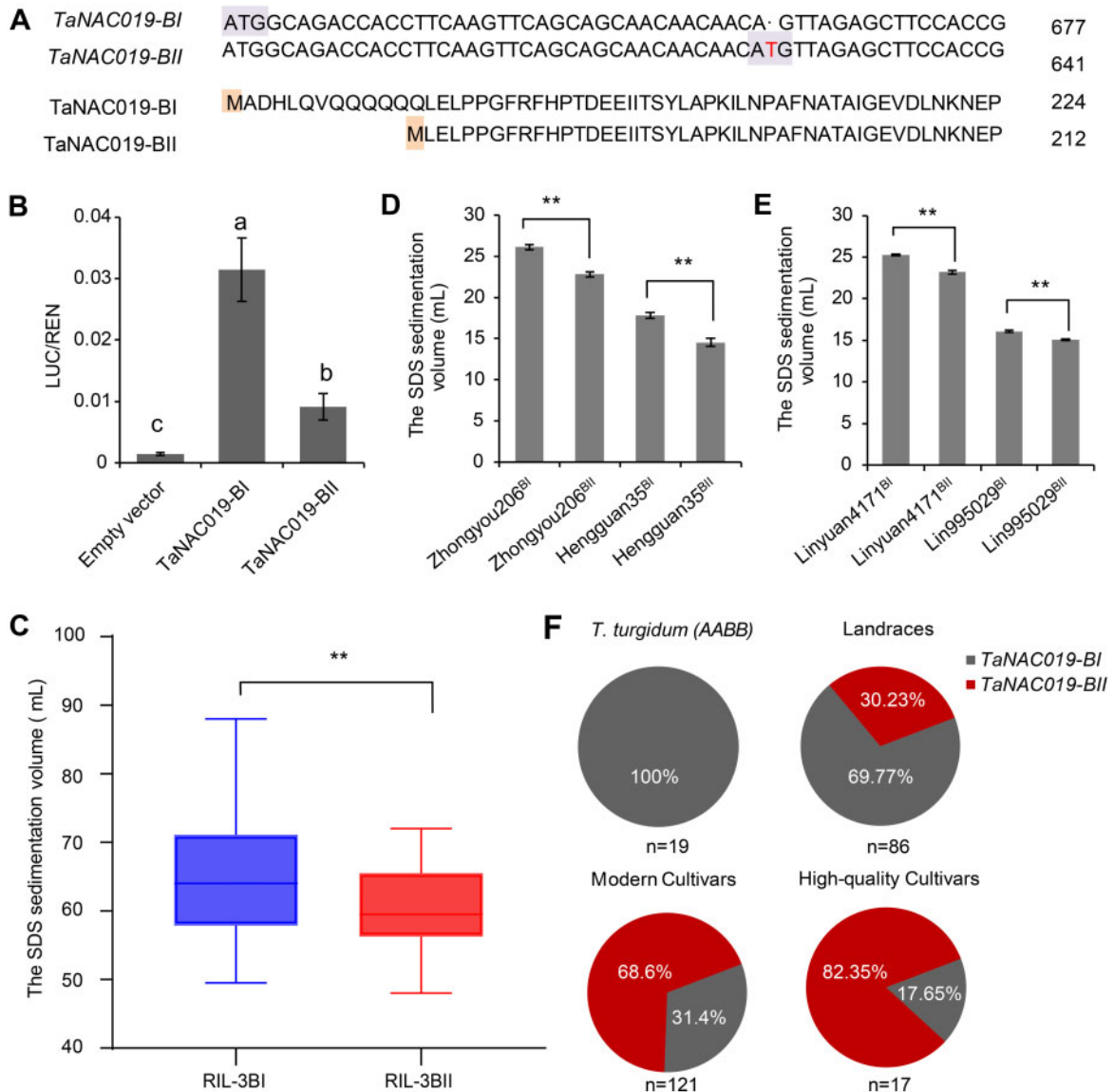


Figure 7 Natural variation of *TaNAC019-B*. **A**, Nucleotide and amino acid sequence alignments of *TaNAC019-BI* and *TaNAC019-BII* haplotypes indicating the detected polymorphism. The T insertion located 39 nucleotides downstream of the translation start site of *TaNAC019-BII* is shown in red. **B**, Dual luciferase transcriptional activity assay (performed as described in Figures 5, C and 6, A) to compare the efficiency of *TaNAC019-BI* and *TaNAC019-BII* in activating *TaGlu-1Dy* promoter. On the y-axis, LUC/REN indicates the ratio of the signals detected for firefly luciferase (LUC) and *R. reniformis* luciferase (REN) activity. Error bars represent standard deviation for six replicates. The different letters above each bar indicate that means differ significantly by one-way ANOVA and Tukey's multiple comparison test at $P < 0.05$. **C**, Comparison of SDS sedimentation volume between RILs containing *TaNAC019-BI* and *TaNAC019-BII*, generated from Jimai44 (*TaNAC019-BI*) and Jimai229 (*TaNAC019-BII*). Lines in the box plots indicate the median. The 10th/90th percentiles of outliers are shown. The double asterisks represent significant differences, as determined by Student's *t* test at $P < 0.01$. **D**, SDS sedimentation volume, measured in backcrossed lines in the Zhongyou206 and Hengguan35 genetic backgrounds containing *TaNAC019-BII* or *TaNAC019-BI*. Double asterisks indicate statistically significant differences estimated by Student's *t* test at $P < 0.01$. **E**, As described in D, except that the SDS sedimentation volume was measured in backcrossed lines in the Linyuan4171 and Lin995029 genetic backgrounds containing *TaNAC019-BII* or *TaNAC019-BI*. **F**, Distribution of *TaNAC019-BII* and *TaNAC019-BI* haplotypes in tetraploid wheat, landraces, elite modern cultivars, and cultivars with high bread-baking quality. *n* indicates the sample size.

The three *TaNAC019* homoeologs exhibit functional divergence

The combination and interactions of the three subgenomes of allohexaploid wheat have given rise to phenotypic variation, which is sometimes associated with the generation of functional divergence among homoeologous genes (Feldman et al., 2012). To test for possible functional divergence

among *TaNAC019-A*, *TaNAC019-B*, and *TaNAC019-D*, we performed a dual luciferase transcriptional activity assay to compare their ability to activate *TaGlu-1Dy* expression. LUC activity was 9-fold and 21-fold higher in the presence of *TaNAC019-A* and *TaNAC019-B*, respectively, compared with the empty vector control, whereas no significant changes were detected in the presence of *TaNAC019-D* (Figure 6, A),

which was consistent with results of Figure 5, C. Similar results were obtained when we analyzed the activation of the *TaGlu-1By* and *TaGlu-1Dx* promoters (Supplemental Figure S15). We also compared the ability of the three TaNAC019 homoeologs to bind to the *TaGlu-1Dy* promoter. Consistent with the results obtained from our transactivation analysis, TaNAC019-B showed a stronger binding affinity to the *TaGlu-1Dy* promoter than TaNAC019-A, whereas TaNAC019-D did not bind to this promoter (Figure 6, B).

A comparison of the amino acid sequences of TaNAC019 homoeologs revealed a high level of similarity (92.9%). However, the NAC domain of TaNAC019-D differs from TaNAC019-A and TaNAC019-B in three amino acid residues located at positions 20, 53, and 90 (Figure 6, C). We introduced these three mutations into TaNAC019-B and used the resulting GST-fused recombinant proteins in EMSAs. The R90RQ mutation was sufficient to impair the ability of the protein to bind to the *TaGlu-1Dy* promoter (Figure 6, D). The R90Q mutation also affected the transactivation ability of TaNAC019-B in a dual luciferase assay, whereas the Q90R mutation in TaNAC019-D restored the transactivation activity of this protein (Figure 6, A). Overall, these results indicated that the abilities of the three homoeologs to bind to and activate the *TaGlu-1Dy* promoter differ, with TaNAC019-D showing no activity.

The elite allele *TaNAC019-BI* improves wheat processing quality

Finally, we used single-nucleotide polymorphism (SNP) analysis to investigate the genetic variation of the three homoeologs in a natural population of 226 wheat cultivars from China and foreign countries, including 19 tetraploid wheat (AABB) accessions, 86 landraces corresponding to the original pool of diversity in China, 121 elite modern cultivars registered after the Green Revolution, and 17 cultivars with high bread-baking quality. This analysis uncovered two *TaNAC019-B* haplotypes in this population (Supplemental Data Set S4). The first haplotype, named *TaNAC019-BI*, encoded a 224 amino acid protein (Figure 7, A). The second haplotype, named *TaNAC019-BII*, carried a single T insertion 39 nucleotides downstream of the translational start site. This insertion led to a premature translation termination, but also generated a new translational start site, thus producing a shorter protein lacking 12 N-terminal amino acids (a glutamine-rich activation domain; Escher et al., 2000). Dual luciferase transcriptional activity assays showed that TaNAC019-BI activates the *TaGlu-1Dy* promoter more strongly than TaNAC019-BII does (Figure 7, B).

To compare the contributions of *TaNAC019-BI* and *TaNAC019-BII* to wheat processing quality, we examined recombinant inbred lines (RILs) generated from the parents Jimai44 (carrying *TaNAC019-BI*) and Jimai229 (carrying *TaNAC019-BII*). RILs containing *TaNAC019-BI* had better processing quality than RILs containing *TaNAC019-BII*, as revealed by higher SDS sedimentation volumes (Figure 7, C). To investigate the value of *TaNAC019-BI* for quality

improvement during wheat breeding, we generated backcrossed lines containing *TaNAC019-BII* in the genetic background of two cultivars (Zhongyou206 and Hengguan35) originally harboring the *TaNAC019-BI* variant (Supplemental Figure S16). The SDS sedimentation volume was significantly lower in backcrossed lines (BC₃F₃) Zhongyou206^{BII} and Hengguan35^{BII} compared with Zhongyou206^{BI} and Hengguan35^{BI} (Figure 7, D). Conversely, the production of backcrossed lines (BC₃F₃) with *TaNAC019-BI* in the genetic background of two cultivars (Linyuan4171 and Lin995029) originally containing *TaNAC019-BII* led to an increase in the SDS sedimentation volume (Figure 7, E). Moreover, kernel weight increased in association with *TaNAC019-BII*, but only in the Hengguan35 genetic background (Supplemental Figure S17). These results suggest that the *TaNAC019-BI* allelic variant is associated with higher wheat quality compared with *TaNAC019-BII*.

All tetraploid ancestors tested here contained the *TaNAC019-BI* haplotype (Figure 7, F), indicating that this haplotype originated from tetraploid wheat. Curiously, 69.8% of the 86 landraces contained *TaNAC019-BI*, but only 31.4% of elite modern cultivars and 17.65% of wheat cultivars with high bread-making quality had this haplotype (Figure 7, F). These observations suggest that *TaNAC019-BII* was selected during the modern wheat breeding process. Collectively, these findings identify *TaNAC019-BI* as a novel marker for wheat processing quality that could be used in breeding programs.

Discussion

Mechanism of TaNAC019-mediated regulation of SSP and starch accumulation in wheat seeds

In this study, we showed that *nac019* knock-out mutants of all three *TaNAC019* homoeologs exhibited the following characteristics: (i) reduced transcript levels of almost all genes encoding SSPs (HMW-GS, LMW-GS, and gliadins) and TaSPA, a well-known SSP activator (Albani et al., 1997; Ravel et al., 2014), in 25 DAP endosperm, resulting in decreased levels of HMW-GS and LMW-GS but not gliadin in mature seeds; (ii) reduced transcript levels of many genes involved in starch metabolism in 25 DAP endosperm, resulting in lower starch contents in mature seeds; and (iii) reduced mature seed size and weight, as well as reduced parameters associated with processing quality. In addition, TaNAC019 bound to a specific motif in the *TaGlu-Dy* promoters, and activated their expression; it also physically interacted with and additively cooperated with TaGAMYb for *TaGlu-Dy* transactivation. Overall, these findings indicate that TaNAC019 plays an important role in the concomitant regulation of SSP and starch contents.

We propose a model describing how TaNAC019 regulates SSP and starch levels in wheat grain (Figure 8). According to this model, TaNAC019 regulates the expression of SSP genes in two ways. First, it directly binds to and activates SSP promoters. Second, it activates TaSPA expression, which might occur indirectly, as no TaNAC019-binding site was detected

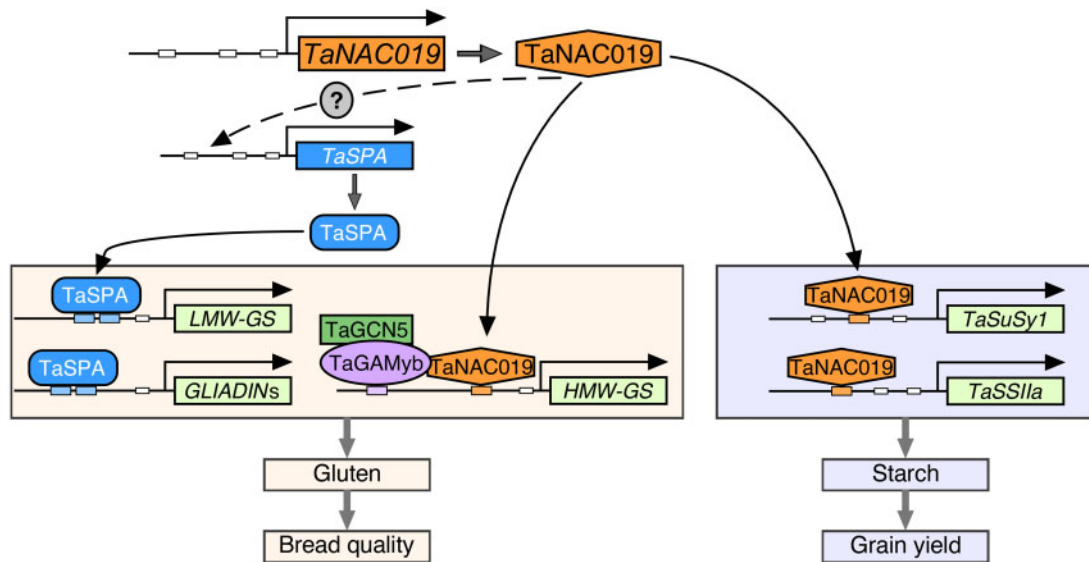


Figure 8 Model of the role of TaNAC019 in regulating SSP and starch accumulation in wheat seeds. See the text for details. Dotted arrows indicate indirect activation, while solid arrows represent direct activation due to the presence of a TaNAC019-binding site within the promoter. The question mark indicates the possible TaNAC019-mediated regulation of still unknown regulatory factors. TaNAC019 might regulate other genes involved in cell division or other processes controlling cell number, which were not shown in this model.

in the 2.5-kbp sequence upstream of the TSS of *TaSPA*. TaNAC019 binds to the promoters and activates the expression of *HMW-GS* genes via additive cooperation with TaGAMyb. Since TaGAMyb recruits the TaGCN5 HAT (Guo et al., 2015), it is likely that chromatin-related regulatory mechanisms are also involved in this process. In addition, SSP genes are expressed earlier than *TaNAC019* genes, suggesting that other regulatory factors regulate SSP genes during early seed developmental stage, but TaNAC019 is required during later stage. TaNAC019-mediated activation of genes encoding LMW-GS and gliadin might primarily occur indirectly through the activation of *TaSPA* and/or other still unknown regulatory factors. Indeed, the TaNAC019-binding site identified in the *TaGlu-1* promoters was only found in a few genes encoding LMW-GS and gliadin. Notably, in mature seeds, the LMW-GS and HMW-GS protein contents were lower in *nac019* compared with wild type, whereas similar levels of gliadin were detected in these seeds. Perhaps this was due to compensation, which occurs when the protein level of one of the two SSP types is altered (Galili et al., 1986; Dumur et al., 2004; Gil-Humanes et al., 2012; Chen et al., 2019). Since *gliadin* transcript levels were lower in *nac019* versus wild type even during late endosperm development, i.e. after the peak of *gliadin* gene expression, the restoration of gliadin proteins to wild-type levels in the *nac019* mutants appears to be related to a post-transcriptional mechanism. A similar phenomenon was observed in the maize *reas1-ref* mutant, where the expression of *zein* genes was downregulated, but their translation level was increased significantly (Qi et al., 2016). TaNAC019-mediated regulation of starch accumulation likely occurs via a similar mechanism (Figure 8). TaNAC019 was able to directly bind the *TaSuSy1* and *TaSSIIa* promoters and activate their expression in vivo (Figure 4, C and Supplemental

Figure S13). Since the TFs and mechanisms involved in regulating wheat starch metabolism have not yet been clarified (Kang et al., 2013; Kumar et al., 2018), it is unclear whether, in addition to binding to and activating various starch metabolism genes, TaNAC019 also modulates the expression of specific regulatory factors. In addition, transcriptome analysis indicated that TaNAC019 is not only involved in SSP and starch gene regulation; it may also regulate genes involved in cell division or other processes controlling cell number.

A recent study reported that overexpression of *TaNAC019-A* reduces seed size and starch content (Liu et al., 2020), which is similar to the phenotype of the *nac019* mutant. *TaNAC019* is specifically expressed in the endosperm; therefore, the reduced seed size and starch content might be a secondary effect of *TaNAC019* overexpression outside of the seed. Moreover, overexpression of a TF perturbs the stoichiometry of TF-containing complexes, which can affect the regulation of their target genes (Levine and Tjian, 2003). By contrast, the precise elimination of a TF through gene editing represents a more specific strategy relative to overexpression. In addition, Liu et al. (2020) reported that *TaNAC019-A* repressed the expression of *ADP-Glucose Pyrophosphorylase Small subunit 1* (*TaAGPS1*), which was downregulated in the overexpression line. We found that *TaAGPS1* was also downregulated in the *nac019* mutant (Supplemental Figure S9). TaNAC019 may act by both activating and repressing the expression of different genes and intra- and intermolecular interactions are relevant for NAC-mediated gene activation or repression (Xu et al., 2013). These observations might also indicate functional divergence among the three homoeologs, considering that TaNAC019-B showed stronger binding affinity for the *TaGlu-1Dy* promoter than TaNAC019-A (Figure 6, A and B), and TaNAC019-A might play an important role in *TaAGPS1* regulation.

Finally, there are six additional NAC TF genes in wheat with seed-specific expression, whose functions are currently unknown. Since SSP and starch production are two major processes that occur during seed formation, it is reasonable to assume that other seed-specific NACs are involved in their regulation. We found that the mRNA expression patterns of these NACs differ, with specific transcripts accumulating at distinct developmental stages, suggesting that they have distinct and/or stage-specific activity. A comprehensive functional characterization of each seed-specific NAC will provide definitive information about their biological roles. However, these NACs probably form an elaborate network that functions during different stages of seed development and might interact at the genetic or physical levels to optimize SSP and starch levels.

Peculiarities of TaNAC019-mediated regulation

A recent study (Zhang et al., 2019) provided evidence that the NAC paralogs ZmNAC128 and ZmNAC130, which are specifically expressed in maize seed, regulate both SSP and starch accumulation. These maize NACs are the orthologs of two rice NACs, Os01g01470 and Os01g29840. In previous phylogenetic analyses (Borrill et al., 2017; Guerin et al., 2019) and in our data (Supplemental Figure S1), a reciprocal BLAST strategy was used to identify wheat NAC orthologs across rice and barley. In this strategy, if the reciprocal BLAST analysis did not identify the same pair of genes in both directions, the genes in question were ruled out as orthologs. Although some rice orthologs of wheat NACs were found using this strategy, no *TaNAC019* orthologs were identified in rice. Hence, although ZmNAC128/ZmNAC130 and TaNAC019 play similar roles in regulating SSP and starch accumulation, they are not orthologs. However, it is worth noting that ZmNAC128/ZmNAC130 and Os01g01470/Os01g29840 do not have any other orthologs among seed-specific NACs in wheat. This finding suggests that seed-specific rice/maize NACs differentiated from wheat NACs, likely reflecting the evolutionary distance between these species. In this context, it is relevant that wheat contains specific SSP classes, such as HMWs, which may require specific regulatory factors not present in maize or rice.

Aside from possible evolutionarily related differentiation, ZmNAC128/ZmNAC130 and TaNAC019 also differ in the number of their SSP and starch-related targets. ZmNAC128/ZmNAC130 regulate the expression of some but not all SSP genes (Zhang et al., 2019), whereas TaNAC019 modulates the expression of genes for all SSP types. In addition, ZmNAC128/ZmNAC130 activate the expression of seven genes involved in starch metabolism (Zhang et al., 2019). Six of these genes have wheat orthologs that are also downregulated in *nac019-2*, but 10 additional starch-related genes were downregulated in this mutant (Supplemental Figure S9). In conclusion, it appears that TaNAC019 has a broader effect than ZmNAC128/ZmNAC130 on the expression of genes associated with SSP and starch production.

Another TaNAC019 feature is the presence of three homoeologs exhibiting functional divergence, which

represents an intrinsic peculiarity of allohexaploid bread wheat, where different subgenome combinations and interactions have given rise to phenotypic variation (Feldman et al., 2012). We found that one of the three TaNAC019 homoeologs, TaNAC019-D, can interact with TaGAMYb but does not bind to or activate *TaGlu* promoters. The very low transcript levels of *TaNAC019-D* suggest that it might be a pseudogene without any residual relevant function. However, this notion contrasts with the finding that the R90Q mutation, which impairs the binding capacity of TaNAC019-D, is fully conserved in all hexaploid wheat accessions (Supplemental Data Set S4). The TaNAC019-D sequence in the seven diploid goat grass (*Aegilops tauschii*) accessions, which are among the D genome donors for hexaploid wheat, also contains the Q residue, suggesting that this mutation originated in *A. tauschii* (Supplemental Data Set S4). Similar to other NACs (Mendes et al., 2013; Mathew et al., 2016), TaNAC019 homoeologs, including TaNAC019-D, interact with other regulatory factors such as TaGAMYb. The interaction between TaNAC019-D and TaGAMYb did not affect the TaGAMYb-related activation of *TaGlu* in dual luciferase transactivation assays. However, TaNAC019-D might act as a modulator of gene expression with other TFs.

In this study, we focused on the roles of TaNAC019 proteins as activators of SSPs and starch accumulation. However, transcriptome analysis revealed a similar number of down- and upregulated genes in *nac019-2* versus the wild type, and a similar percentage of TaNAC019-binding sites was found in the two groups of genes. Indeed, a NAC can act both as an activator and repressor, and the conformational interactions between the N-terminal NAC domain, harboring a NAC repression domain (NARD), and the C-terminal transcriptional activation region (TAR) of NAC proteins are crucial for their activating or repressive activities (Xu et al., 2013). Accordingly, the capacity of TaNAC019 to function as a transcriptional repressor or activator might depend on its conformational assembly at the binding site. Different assemblies could occur depending on interactions with other proteins, as well as the promoter context, which could provide different stabilization contacts for the DNA–protein interaction.

TaNAC019 is a new candidate gene for improving processing quality in wheat

Two TFs were shown to be involved in the concomitant regulation of SSP and starch accumulation in rice and maize seeds (Wang et al., 2013; Zhang et al., 2019). TaNAC019 is the first wheat TF with this function identified to date in wheat. Starch content is related to yield, while SSP content and composition are associated with quality (Rharrabti et al., 2001; Zorb et al., 2018). Therefore, the finding that TaNAC019 regulates both starch and SSP contents and affects the HMW/gliadin and LMW/gliadin ratios in mature seeds makes *TaNAC019* an excellent candidate for developing wheat cultivars with both high yields and good quality. Overexpression of *TaNAC019-A* led to reduced starch

content, kernel weight, and kernel width (Liu et al., 2020), indicating that specific levels of *TaNAC019* might be required for yield and quality improvement. Biotechnological approaches may be used to precisely control *TaNAC019* expression in seeds. In addition, our analysis of the genetic variability of *TaNAC019* in natural populations provided evidence that a specific haplotype, *TaNAC019-BI*, is positively associated with wheat processing quality. Hence, *TaNAC019-BI* represents a genetic marker that may be used in breeding programs.

Importantly, only 30% of modern cultivars carry a *TaNAC019-BI* allele, whereas this allele was detected in 70% of the landraces examined, suggesting that this allele has not yet been well exploited by breeding programs. During the Green Revolution, bread wheat underwent a dramatic transformation, resulting in a significant reduction in genetic diversity, while landrace germplasms remained largely unexploited (Balfourier et al., 2019). As demonstrated by the *Reduced height (Rht)* dwarfing and *Fusarium head blight 1 (Fhb1)* disease resistance genes (Hedden, 2003; Su et al., 2019), these landrace accessions are a valuable reservoir of novel alleles. Perhaps the *TaNAC019-BI* haplotype from landraces could be exploited, providing another example of how this germplasm could contribute to breeding programs.

Materials and methods

Plant materials

The names and geographical origins of all wheat (*T. aestivum*) cultivars, landraces and *Triticum turgidum* (AABB) accessions used in this study are listed in [Supplemental Data Set S4](#). All wheat plants were grown in the experimental field of China Agricultural University in Beijing (39°57'N, 116°17'E) and in a greenhouse at a relative humidity of 75% and 26/20°C day/night temperatures, with a light intensity of 3,000 lux (Master GreenPower CG T 400W E40; Philips). For gene expression analysis, root, leaf, and stem tissues were harvested from greenhouse-grown plants 10 days after seed germination, and young spikes (2 cm), embryos at 15 DAP, whole seeds at the watery ripe stage (4 and 7 DAP), and endosperm at the milky stage (12, 15, and 18 DAP), soft dough stage (20 and 25 DAP), and hard dough stage (30 and 45 DAP) were harvested from plants grown in the field. The endosperm and embryos were manually obtained by dissecting grains. For each sample, developing seeds from three different plants within one planting plot were mixed to represent one biological replicate and at least three biological replicates from different plot or different year (2017–2019) were used. All samples were immediately frozen in liquid nitrogen and stored at –80°C.

Backcrossed lines were obtained as shown in [Supplemental Figure S15](#). The restriction endonuclease *PciI* (NEB, R0655L) was used to distinguish between *TaNAC019-BI* and *TaNAC019-BII* by the Cleaved Amplified Polymorphic Sequence (CAPS) method, which was also used for genotyping during backcrossed lines generation. The RIL population

was generated from two parents (Jimai44 and Jimai229) with selfing for five generations.

Yeast one-hybrid screening and assay

For yeast one-hybrid screening a cDNA expression library derived from wheat seeds mixed from various pollination days in the pGADT7 vector was used as prey. The promoter fragment for *TaGlu-1Dy* was amplified and cloned into the pAbAi vector. This construct was linearized by the restriction enzyme *BstBI* and then integrated into the genome of the Y1HGold yeast strain to obtain a bait-specific reporter strain. Yeast one-hybrid screening was performed following the manufacturer's instructions (Takara, 630491).

For the detection of *TaNAC019* binding to a specific *TaGlu-1Dy* promoter region, the 35 bp P11 sequence and the corresponding MP11-mutated sequence as three repeats were synthesized and inserted into the pAbAi vector, respectively (BGI, China). The construct was linearized by *BstBI* digestion and transformed into Y1HGold cells to generate the Y1H bait strain. Full-length *TaNAC019-BI* cDNA was inserted into the pGADT7 vector and transformed into the Y1H bait strain. The transformants were selected on synthetic dropout –Ura and –Leu medium containing 125 mM AbA.

Phylogenetic analysis

We obtained protein sequences with a NAM domain (PF02365) in barley (122), wheat (454), and rice (94) from Ensembl Plants with the BioMart tool from Phytosome ([Supplemental File S2](#)). Maize sequences were reported by Zhang et al. (2019; [Supplemental File S2](#)). We aligned the amino acid sequences corresponding to the NAM domain with ClustalW and constructed a phylogenetic tree using the neighbor-joining method in MEGA 7 software with default parameters (Kumar et al., 2016). The evolutionary distances were calculated using the Poisson model. The phylogeny test was computed using bootstrap method with 10,000 replications ([Supplemental File S3](#)).

Subcellular localization

The full-length open-reading frames (ORFs) of *TaNAC019-A/B/D* were amplified using specific primers and Tks Gflex DNA polymerase (TaKaRa, R060A) and cloned upstream of the 5'-end of the GFP gene (*GFP*) in the pCAMBIA1300 vector containing the *mannopine synthase* promoter (*maspro*). The resulting construct *maspro:TaNAC019-GFP* and the *maspro:GFP* control were separately introduced into young leaves of 4-week-old *N. benthamiana* plants by *Agrobacterium (Agrobacterium tumefaciens)*-mediated transient infiltration (Liu et al., 2010). The infiltrated plants were cultured at 22°C for 48 h, and GFP signals observed on a confocal microscope (Eclipse TE2000; Nikon).

Production of *nac019* knock-out mutants

For genome editing via CRISPR/Cas9, a sgRNA was designed based on the first exon sequence of *TaNAC019* using the E-CRISP Design website (<http://www.e-crisp.org/E-CRISP/design>)

crispr.html). Two reverse complementary sgRNA sequences with *Bsal* cohesive ends were synthesized. Double-stranded oligonucleotides were annealed by slowly cooling from 100°C to room temperature and inserted into the expression cassette of the pBUE411 vector (Xing et al., 2014). The resulting plasmid was transformed into wheat cultivar Fielder via *Agrobacterium*-mediated (strain EHA105) transformation (Ishida et al., 2015). Homozygous *nac019-1* and *nac019-2* mutants were obtained by selfing of T_0 plants for four generations. At each generation, sequences of all three homoeologs from representative transgenic seedlings were analyzed. The lines with homozygous editing at *TaNAC019-A/B/D* were selected for selfing. For example, lines homozygous for edited alleles at *TaNAC019-A* and *TaNAC019-B* were obtained in the T_2 generation of *nac019-1* mutant, and homozygous edited alleles at *TaNAC019-D* were obtained in the T_3 generation.

Phenotypic analysis

Wild-type and *nac019* seeds were planted in a block containing 10 rows (1.5 m long) at a spacing of 20 cm. Mature seeds from the plot were harvested and seeds from every 10 plants were mixed for phenotypic assessment. After adjusting the moisture content of seeds to 14%, the seeds were milled into flour using a Brabender Quadrumat Junior laboratory mill (Brabender, Duisburg, Germany). The contents of HMW-GSs, LMW-GSs, and gliadins were detected by RP-HPLC as previously described with some modifications (Zhang et al., 2014): the Agilent Technologies 1260 Infinity II RP-HPLC system and an Agilent ZORBAX 300SB-C18 column (150 mm × 4.6 mm, 5 μm) were used. The elution solvents were water containing 0.6 mL/L trifluoroacetic acid (solvent A) and acetonitrile containing 0.6 mL/L trifluoroacetic acid (solvent B). The elution gradient for glutenins was as follows: 0–2 min, 21% B; 2–52 min, from 21% to 53.5% B; 52–54 min, from 53.5% to 21% B; 54–60 min, 21% B. The elution gradient for gliadins was as follows: 0–2 min, 25% B; 2–27 min, from 25% to 50% B; 27–35 min, from 50% to 25% B; 35–50 min, 25% B. The column temperature was 60°C and the flow rate was 1 mL/min. The injection volume was 8 μL and the eluent was monitored at 210 nm. The total amounts of HMW-GSs, LMW-GSs, and gliadins were estimated by integrating the relevant RP-HPLC peaks present in the chromatograms. Three replications were performed for each sample. Dry gluten content was evaluated using the voluntary standard method of the National Standard of China 5506.4-2008 (GB/T5506.4-2008). SDS sedimentation values were determined using 2 g of flour to detect sedimentation volume for 5 min as described by Chen et al. (2019). Three repeats were carried out for each sample.

A Wanshen SC-G seed detector (Hangzhou Wanshen Detection Technology Co., Ltd.) was used to measure kernel weight, grain width, and grain length. Nine replicates were used, each with approximately 25 g seeds. Starch content was determined using Megazyme Total Starch Assay Kit (Megazyme; Catalog no.: KTSTA-50A) based on the use of

thermostable α -amylase and amyloglucosidase. Three repeats were carried out for each sample.

Histological analysis

Immature wild-type and *nac019* mutant seeds at 6, 15, and 20 DAP were sampled from spikes. Seeds were fixed in FAA solution (63% ethanol, 5% acetic acid, 2% formaldehyde) for 24 h at room temperature. After dehydration in a graded ethanol series, sections were slowly infiltrated with Steedman's wax, embedded in paraffin, sectioned to 4 μm thickness with a microtome, and dewaxed in absolute ethanol. Sections (4 μm) were stained with Toluidine Blue. Images were collected using a OLYMPUS BX53 Microscope. Endosperm cell number was manually calculated for three seeds and the differences between WT and the *nac019* mutants were determined by one-way ANOVA. Starch granules from three seeds of each line were stained by Periodic Acid–Schiff.

RT-qPCR

Total RNA was extracted from samples using TransZol Plant (TransGen Biotech, ET121-01), and first-strand cDNAs (20 μL) were synthesized from 1 μg starting total RNA using a reverse transcription kit (Vazyme Biotech, R223-01) according to the manufacturer's instructions. The first-strand cDNAs was diluted with H₂O to 80 μL. For RT-qPCR, the reaction mixture was composed of the 1 μL cDNA template, 0.2 mM primers, and 5 μL SYBR Green Mix (Vazyme Biotech, Q121-02/03) in a final volume of 10 μL. Amplification was performed using a CFX96 real-time system (BioRad). Differences in relative transcript levels were calculated using the $2^{-\Delta\Delta CT}$ method (https://assets.thermo-fisher.com/TFS-Assets/LSG/manuals/cms_040980.pdf) relative to wheat *TaACTIN* (TraesCS5B02G124100). RT-qPCR was performed as technical triplicates per sample. Three biological replicates were performed, with similar results; the results from one replicate are shown in the figures.

Electrophoretic mobility shift assay

TaNAC019 was expressed in *Escherichia coli* (BL21 DE3) by cloning the *TaNAC019* coding sequences into the pGEX6P-1 vector (Solarbio, P0300) to produce a GST fusion protein. Recombinant protein expression was induced with 0.1 mM isopropyl- β -D-thiogalactopyranoside (IPTG) in Luria Bertani (LB) buffer overnight at 16°C. The cells were harvested, sonicated in phosphate-buffered saline (PBS; 137 mM NaCl, 2.7 mM KCl, 10 mM Na₂HPO₄, 2 mM KH₂PO₄) containing 1 mM phenylmethanesulfonyl fluoride (PMSF) and 1/2 of a tablet of protease inhibitor cocktail (Roche) for 1 h, and centrifuged at 13,000 × g for 45 min. The supernatant was mixed with GST MAG Agarose Beads (Novagen, <http://www.novagen.com>) and rocked overnight at 4°C. The fusion protein was eluted from the beads with 50 mM Tris–HCl (pH 8.0) containing 10 mM reduced glutathione.

The double-stranded probes (labeled with biotin at their 5'-end) were annealed from complementary oligonucleotides by cooling from 100°C to room temperature in annealing

buffer. The sequences of the probes are listed in [Supplemental Data Set S5](#). The DNA-binding reactions were performed in a solution containing 20% binding buffer (100 mM Tris, 500 mM KCl, 10 mM DTT; pH 7.5), 2.5% glycerol, 0.2 mM EDTA, and 50 ng/ μ L poly(dI–dC). Following incubation at room temperature for 20 min, the samples were loaded onto a 6% native polyacrylamide gel. Transfer to a nylon membrane and detection of biotin-labeled DNA were performed according to the manufacturer's instructions (Thermo Scientific 20148 LightShift Chemiluminescent EMSA Kit). Three replicates were performed, with similar results; the results from one replicate are shown in the figures.

Yeast two-hybrid assay

The coding sequences of *TaSPA*, *TaNAC019-A*, *TaNAC019-B*, and *TaNAC019-D* were cloned into the pGADT7 vector, while *TaGAMyb*, *TaGCN5*, *TaNAC019-A*, and *TaNAC019-B* were cloned into the pGBKT7 vector. For the yeast two-hybrid assays, the plasmids were transformed into yeast strain AH109 using the lithium acetate method, and the cells were grown on minimal medium (–Leu –Trp) according to the manufacturer's instructions (Clontech). The positive colonies with T-antigen and P53 were plated onto minimal medium (–Leu –Trp –His –Ade) to test for possible interactions.

Dual luciferase transcriptional activity assay

Dual luciferase transcriptional activity assays were performed as previously described ([Guan et al., 2014](#)). Fragments corresponding to 1,600 bp upstream of the TSS of *TaGlu-1Bx*, *TaGlu-1By*, *TaGlu-1Dx*, *TaGlu-1Dy*, *TaSuSy1*, and *TaSSIIa* were PCR-amplified from CS genomic DNA and cloned into the pGreenII 0800-LUC vector as reporter plasmids ([Hellens et al., 2005](#)). The ORFs of *TaNAC019* and *TaGAMyb* were cloned into the pHB vector as effectors. Young leaves of 4-week-old *N. benthamiana* plants were co-infiltrated with *Agrobacterium* (strain GV3101) harboring different combinations of these plasmids. Firefly luciferase activity derived from the *TaGlu-1Bx* (*TaGlu-1Bxpro:LUC*), *TaGlu-1Dy* (*TaGlu-1Dypro:LUC*), *TaGlu-1By* (*TaGlu-1Bypro:LUC*), *TaGlu-1Dx* (*TaGlu-1Dxpro:LUC*), *TaSuSy1* (*TaSuSy1pro:LUC*), or *TaSSIIa* (*TaSSIIapro:LUC*) promoters and Renilla luciferase under the control of the 35S promoter (*35Spro:REN*) were quantified using the Dual-Luciferase Reporter Assay system (Promega) with a Synergy 2 Multi-Detection Microplate Reader (BioTek Instruments). Normalized data are presented as the ratio of luminescent signal intensity for reporter versus internal control reporter (*35Spro:REN*) from three independent biological samples.

TaNAC019 antibody production

The full-length *TaNAC019-B* sequence was used to produce a specific rabbit polyclonal antibody by Wuxi Pharma Tech Company (China).

Immunoblot analysis

Harvested seeds and endosperm were ground in liquid nitrogen, and proteins were extracted in 4 \times loading buffer (200 mM Tris–HCl pH 6.8, 40% [v/v] glycerol, 8% [w/v] SDS, and 20% [v/v] β -mercaptoethanol) at 100°C for 10 min. The solution was placed on ice for 2 min and centrifuged for 10 min at 12,000 rpm at room temperature. Protein extracts were separated by 10% (v/v) SDS-PAGE and transferred to a polyvinylidene fluoride membrane (cat. no. 162-0177; Bio-Rad) for 60 min at 300 mA under temperature-controlled conditions. The membrane was blocked using 5% (w/v) fat-free milk for 1 h and incubated overnight at 4°C in 1 \times Tris-buffered saline containing Tween 20 (TBST; 1 \times TBST with 0.1% [v/v] Tween 20) and anti-*TaNAC019-B* antibody (diluted 1:1,000). After washing three times with 1 \times TBST for 10 min each time, the membrane was incubated in 5% (w/v) fat-free milk in 1 \times TBST solution containing the secondary antibody (diluted 1:3,000) for 1 h at room temperature, followed by three washes (10 min each time) with 1 \times TBST. Two-component Reagent-clarity Western ECL Substrate (cat. no. 170-5060; Bio-Rad) was used for detection. The signals were detected using a Chemiluminescence Imaging system (ChemiScope 6000). Three biological replicates were performed, with similar results; the results from one replicate are shown in the figures.

Pull-down assay

To generate His-tagged *TaGAMyb* protein, the *TaGAMyb* ORF was amplified and cloned into the EcoRI/Sall sites of the pET-32a vector (Novagen). To generate GST-tagged *TaNAC019-A*, *TaNAC019-B*, and *TaNAC019-D*, the respective ORFs were amplified and cloned separately into the BamHI/EcoRI sites of the pGEX-6P-1 vector (Solarbio, P0300). Recombinant protein production was induced following the manufacturer's manual. For the GST pull-down assay, approximately 5 mL *E. coli* lysates containing GST or GST-*TaNAC019* fusion protein were mixed with 400 μ L immobilized glutathione agarose (Thermo Scientific) for 2 h at 4°C. The beads were washed five times with 1 mL PBS and incubated with His-tagged *TaGAMyb* fusion protein lysates for more than 2 h at 4°C. The protein complex-immobilized glutathione agarose conjugates were eluted in 100 μ L 4 \times loading buffer (200 mM of Tris–HCl at pH 6.8, 40% [v/v] glycerol, 8% [w/v] SDS, and 20% [v/v] β -mercaptoethanol) at 100°C for 10 min. The bound proteins were analyzed by 10% PAGE and immunoblot analysis was performed with anti-His or anti-GST antibodies (diluted 1:1,000; Abcam). Three replicates were performed, with similar results; the results from one replicate are shown in the figures.

Transcriptome deep sequencing (RNA-seq)

Three biological replicates were performed for each sample. RNA-seq libraries were prepared with a TruSeq RNA Sample Preparation Kit v2 (Illumina) according to the manufacturer's instructions and sequenced to generate 150-bp paired-end reads on the HiSeq platform. Approximately 72–

102 million reads were generated per sample. Raw reads obtained from Illumina sequencing were processed and filtered using Trimmomatic (Bolger et al., 2014) with “SLIDINGWINDOW:4:20 MINLEN:40” to generate high-quality reads. The RNA-seq reads were aligned to the CS reference genome (International Wheat Genome Sequencing Consortium RefSeq v1.1) downloaded from Ensembl Plants using STAR with default parameters (Dobin et al., 2013). FeatureCounts was used to count the reads mapped to each gene (Liao et al., 2014). The read counts were normalized to FPKM values to show the relative gene expression levels. After using a gene expression criterion of FPKM value ≥ 1 in at least one sample, DESeq2 was used for differential gene expression analysis (Love et al., 2014). The genes showing an absolute value of \log_2 (fold-change) > 0.8 and adjusted *P*-value (false discovery rate) of < 0.05 were considered to be differentially expressed genes.

Significantly enriched GO categories of up- and downregulated genes (adjusted *P*-value [false discovery rate] < 0.05) were identified using ClusterProfiler (Yu et al., 2012). FIMO with default parameters was used to identify TaNAC019-binding motifs within 2.5 kbp upstream of the TSS of each gene (Grant et al., 2011).

Statistical analysis

Quantitative data, including three biological and at least three technical replicates, are presented in the form of mean \pm SD. Means of two samples were compared using Student's two-tailed *t* tests. Analysis of variance (one-way ANOVA) was conducted using Graphpad Prism 8.0.2 software with default parameters. Significant differences were determined by one-way ANOVA or the Student's *t* test (Supplemental File S1): **P* < 0.05 and ***P* < 0.01 . Tukey's multiple comparison test was used to determine exactly which group means were different.

Data availability

Accession numbers

The cDNA sequences of *TaNAC019-A*, *TaNAC019-BI*, *TaNAC019-BII*, and *TaNAC019-D* have been deposited in GenBank under accession numbers MN685185, MN685186, MN685187, and MN685188, respectively. RNA sequencing data are available at the National Center for Biotechnology Information Sequence Read Archive (<http://www.ncbi.nlm.nih.gov/sra>) under accession number PRJNA656089.

Supplemental data

Supplemental Figure S1. Phylogenetic analysis of wheat, maize, rice, and barley NACs.

Supplemental Figure S2. Genomic locations of *TaNAC019* homoeologs.

Supplemental Figure S3. Subcellular localization of *TaNAC019* homoeologs.

Supplemental Figure S4. *TaGlu* expression during wheat seed development.

Supplemental Figure S5. Characterization of anti-TaNAC019 antibody and analysis of TaNAC019 levels.

Supplemental Figure S6. Strategy used to produce the *nac019* knockout lines.

Supplemental Figure S7. HMW-GS from *nac019-1*, *nac019-2*, and WT in a silver-stained protein gel.

Supplemental Figure S8. Histological analysis of developing seeds of *nac019* and wild type and phenotypes of mature *nac019* and wild-type plants.

Supplemental Figure S9. Starch metabolism genes that are downregulated in *nac019-2* compared with the wild type.

Supplemental Figure S10. GO analysis of genes upregulated in the *nac019-2* transcriptome compared with the wild type.

Supplemental Figure S11. Analysis of the binding of TaNAC019-B to the *TaGlu-1Dy* promoter via EMSA.

Supplemental Figure S12. Analysis of the binding of TaNAC019-B to the *TaGlu-1Dy* promoter via yeast one-hybrid analysis.

Supplemental Figure S13. Validation of the TaNAC019-binding motif identified using EMSA assay.

Supplemental Figure S14. Yeast two-hybrid analysis of TaNAC019 protein interactions.

Supplemental Figure S15. Dual luciferase transcriptional activity assays to detect TaNAC019-induced activation of *TaGlu-1Dx* and *TaGlu-1Bx* promoters.

Supplemental Figure S16. Schematic representation of the method used to generate backcrossed lines.

Supplemental Figure S17. TGWs of the backcrossed lines.

Supplemental Data Set S1. List of differentially expressed transcripts in the *nac019* mutant and wild type.

Supplemental Data Set S2. Expression of *SSP* genes in the *nac019* and wild type.

Supplemental Data Set S3. Expression of genes related to starch biosynthesis in *nac019* and the wild type.

Supplemental Data Set S4. Names and geographical origins of all wheats used in this research, and haplotype analysis of TaNAC019-B and TaNAC019-D.

Supplemental Data Set S5. Probes sequences and location on the promoter of *TaGlu-1Dy* used for EMSA.

Supplemental Data Set S6. Primers used for genes expression and vectors construction.

Supplemental File S1. Summary of statistical analyses.

Supplemental File S2. Protein sequences for the phylogenetic tree shown in Supplemental Figure S1.

Supplemental File S3. Protein sequence alignment for the phylogenetic tree shown in Supplemental Figure S1.

Supplemental File S4. Newick format of the phylogenetic tree of Supplemental Figure S4.

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Conflict of interest statement. The authors declare that they have no competing interests.

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