Cloning and Characterization of a Critical Regulator for Pre-harvest Sprouting in Wheat
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**ABSTRACT** Sprouting of grains in mature spikes before harvest is a major problem in wheat (*Triticum aestivum*) production worldwide. We cloned and characterized a gene underlying a wheat quantitative trait locus (QTL) on the short arm of chromosome 3A for pre-harvest sprouting (PHS) resistance in white wheat using comparative mapping and map-based cloning. This gene, designated *TaPHS1*, is a wheat homolog of a MOTHER OF FLOWERING TIME (*TaMFT*)-like gene. RNA interference-mediated knockdown of the gene confirmed that *TaPHS1* positively regulates PHS resistance. We discovered two causal mutations in *TaPHS1* that jointly altered PHS resistance in wheat. One GT-to-AT mutation generates a mis-splicing site, and the other A-to-T mutation creates a premature stop codon that results in a truncated non-functional transcript. Association analysis of a set of wheat cultivars validated the role of the two mutations on PHS resistance. The molecular characterization of *TaPHS1* is significant for expediting breeding for PHS resistance to protect grain yield and quality in wheat production.

WHEAT is a staple food crop for more than 40% of the world’s population and provides more than 20% of calories and proteins for humans (Gill et al., 2004). Pre-harvest sprouting (PHS) in wheat (physiologically mature grains germinating in spikes before harvest) causes significant loss in grain yield and quality, particularly in regions with prolonged wet weather during the harvest season. Direct annual losses caused by PHS approach US $1 billion worldwide (Black et al., 2006).

Resistance to PHS in wheat is a complex trait that is affected by both genotype and environment (Imtiaz, et al., 2008). Grain color and seed dormancy have long been regarded as two major factors affecting PHS resistance (Gfeller and Svejda, 1960; Bewley, 1997; Groos et al., 2002). White grain wheat is usually more susceptible to PHS than red grain wheat (Gale and Lenton, 1987; Groos et al., 2002; Himi et al., 2002). Demand for white grain wheat is increasing rapidly in many countries because of consumer preferences, higher flour yield, and better end-use quality; therefore, improving resistance to PHS in white wheat is imperative for successful production in environments where PHS occurs.

Seed dormancy is another important trait of PHS resistance (Bewley, 1997; Mares et al., 2005; Sussman and Phillips, 2009). Adequate seed dormancy can reduce or block PHS during harvest seasons, but dormancy breaks down during seed storage so seeds germinate uniformly after sowing. Several other factors have been proposed as potential contributors to overall PHS resistance in field conditions, including germination-inhibitory substances residing in chaff tissue (Derera and Bhatt, 1980; Gatford et al., 2002), physical barriers to water penetration in a spike, and spike morphology such as structure and erectness of wheat spikes, openness of florets, and tenacity of glumes (King and Richards, 1984). The degree to which these factors contribute to the levels of wheat PHS resistance remains unknown.
To date, PHS resistance genes have not been well characterized at the nucleotide sequence level in wheat, although several genes for seed dormancy have been reported in other species. *DELAY OF GERMINATION* 1 (*DOG1*) in *Arabidopsis* and *seed dormancy* 4 (*Sdr4*) in rice (Bentsink et al., 2006; Sugimotoa et al., 2010) were cloned. In wheat, quantitative trait loci (QTLs) for PHS resistance have been reported on most wheat chromosomes (Groos et al., 2002; Intiaz et al., 2008; Kulwal et al., 2012; Mori et al., 2005), and QTLs on chromosomes 2B (Munkvold et al., 2009), 3A (Liu et al. 2008), and 4A (Mares et al., 2005; Liu et al., 2011) have demonstrated major effects on PHS resistance. Recently, a *MOTHER OF FT AND TFL1* was identified to be involved in seed dormancy under low temperature (13°C) through a microarray study (Nakamura et al., 2011). Sequencing the gene from two cultivars identified a single nucleotide polymorphism (SNP) from the promoter region as the functional SNP that regulates seed dormancy in red wheat. However, wheat PHS usually occurs before harvest in a field at much higher temperatures, so it is not known if this gene is responsible for PHS resistance in wheat under natural conditions and whether the SNP causes the change in PHS resistance.

We previously mapped a major QTL (*Qphs.pseru-3AS*) for PHS resistance to the distal end of the short arm of chromosome 3A of a PHS-resistant white wheat cultivar Rio Blanco (Liu et al., 2008; Liu and Bai, 2010). In this study, we used comparative fine-mapping and map-based cloning to 1) determine the candidate gene underlining the QTL, 2) identify the causal variations in the candidate gene responsible for the change in PHS resistance in wheat, and 3) develop a diagnostic gene assay for marker-assisted selection to improve PHS resistance in wheat.

**Materials and Methods**

**Plant materials and PHS evaluation**

The major QTL for PHS resistance, *Qphs.pseru-3AS*, was previously mapped using two F$_6$ recombinant inbred lines (RILs) populations developed from the crosses of Rio Blanco/NW97S186 and Rio Blanco/NW97S078, where Rio Blanco carries *Qphs.pseru-3AS*, whereas NW97S186 and NW97S078 do not (Liu et al., 2008). To fine-map and clone this QTL, RIL#25, a F$_6$ RIL that segregated at *Qphs.pseru-3AS* locus, was selected from the Rio Blanco/NW97S186 population to develop recombinant near-isogenic lines (NILs) of *Qphs.pseru-3AS* using the heterogeneous inbred family method (Tuinstra et al., 1997) (Supplemental Figure S1A). This RIL segregated at the *Qphs.pseru-3AS* region represented by three closely linked markers, *Xbarc321*, *Xbarc57*, and *Xbarc12*, in the QTL region. A total of 1874 RIL#25-derived plants (equals to F$_2$ plants) were screened for recombination within the QTL region using the three SSR markers. The selected heterozygous plants with a recombination event were self-pollinated to produce homozygous recombinants (equals to F$_3$). A total of 56 homozygous NILs with recombination among the three markers were selected (Liu and Bai, 2010). Two contrasting homozygous NILs, 08F485 (resistant NIL) and 08F481 (susceptible NIL)
without recombination among the three markers, and the two parents, Rio Blanco and NW97S186 (Figure 1A), were used to screen polymorphic markers for further fine-mapping and gene expression analysis.

The wheat accessions used for association analysis included 60 white and 22 red wheat accessions. Among them, 61 were from U.S. Southern and Northern Hard Winter Wheat Regional Performance Nurseries, and 21 were kindly provided by the USDA National Small Grain Collection, Aberdeen, Idaho.

Wheat PHS resistance was evaluated in the greenhouses at Kansas State University, Manhattan, Kansas, as described previously (Liu et al., 2008). To evaluate sprouting rates, wheat spikes were harvested from each replication at physiological maturity as characterized by loss of green color on the spike. The harvested spikes were air-dried for 5 days in the greenhouse at 25 ± 5 °C and then stored in a freezer at –20 °C to maintain dormancy. Sprouting assays were conducted in a moist chamber for 7 days at 23 ± 2°C with 100% humidity (Liu et al., 2008). For association mapping, phenotyping experiments were repeated twice with two replications per experiment and five spikes from different plants per accession in each replication. For fine mapping, 3 to 5 spikes per F$_2$ plant were tested for PHS resistance using completely randomized design; the selected homozygous recombinant NILs were evaluated for PHS resistance using a randomized complete block design with two replications and five spikes per replication.

**Comparative fine-mapping**

$Qphs.pseru-3AS$ was initially mapped to a 2.0 cM region at the distal end of the short arm of wheat chromosome 3A (Supplemental Figure S1B). Comparing the QTL map to previously published 3A maps (Song et al., 2005) revealed four common markers for the QTL (Supplemental Figure S1C). Among them, $Xbarc12$ and $Xgwm369$ flanking the QTL were physically mapped in the deletion bin 3AS4-0.45-1.00 (Sourdille et al., 2004) (Supplemental Figure S1D), indicating the QTL was located in the bin of 3AS4-0.45-1.00. Previously, 224 wheat expressed sequence tags (ESTs) were physically mapped in the bin (Munkvold et al., 2004) (http://wheat.pw.usda.gov/cgi-bin/westsql/map_locus.cgi). We randomly selected 114 of these ESTs to develop sequence tag site (STS) markers or sequence directly to identify polymorphisms between the two parents and the two NILs (08F485 and 08F481) for fine-mapping of the QTL.

Comparative mapping allowed us to identify the syntenic region of $Qphs.pseru-3AS$ in rice and *Brachypodium* genomes by BLASTN search for polymorphic wheat EST sequences in the rice and *Brachypodium* genome databases (http://www.jcvi.org; http://www.phytozome.org) (The International Brachypodium Initiative, 2010; Fu et al., 2009). The annotated gene sequences in the syntenic region of rice and *Brachypodium* genomes were used as queries in BLASTN (Altschul et al., 1997) to search the wheat EST database (http://www.ncbi.nlm.nih.gov/ or http://www.plantgdb.org) or Chinese Spring 5x coverage of 454 reads (http://www.cerealsdb.uk.net/CerealsDB/Documents/DOC_CerealsDB.php) to identify additional unmapped wheat ESTs or DNA sequences that may be mapped in the $Qphs.pseru-3AS$ region to develop a fine
map. In the BLASTN searches, a significant match was declared when there was at least 70% nucleotide identity for at least half of the query sequences, but no fewer than 300 bases, and with an e-value lower than $e^{-20}$. For tBLASTX searches, significance was declared when there was at least 40% amino acid identity over at least half of the EST sequences, but no fewer than 200 amino acids at an e-value lower than $e^{-11}$. When several significant matches were found for a single predicted rice or *Brachypodium* gene sequence, the best match was selected (Kuraparthty et al., 2008).

**Development and analysis of STS and SNP markers**

STS primers were designed using Primer3 software (http://frodo.wi.mit.edu/cgi-bin/primer3/primer3_www.cgi) to target amplicons of 400–1200 bp. Polymerase chain reactions (PCR) of 25 μL mixture contained 100 ng of template DNA; 1 mM each of reverse and M13-tailed forward primers; 1 pmole fluorescence-labeled M13 primer; and 0.2 mM each of dNTP, 1X PCR buffer, 2.5 mM MgCl$_2$, and 1.5 units of *Taq* polymerase. PCR was performed in a DNA Engine Tetrad Peltier Thermal Cycler (Bio-Rad Lab, Hercules, CA) using a touch-town program (Liu et al., 2008).

Those ESTs that did not show STS polymorphism were re-sequenced to identify SNPs. The PCR products were sequenced using big dye-terminator chemistry after they were cleaned up by adding 2 U shrimp alkaline phosphatase and 0.4 U exonuclease I, then incubating at 37 °C for 1 hr and 75 °C for 15 min to eliminate enzyme activity. DNA sequence data were checked for sequencing errors using Sequencer software (Gene Codes Corporation, Ann Arbor, MI). Sequences were aligned using ClustlW2 (http://www.ebi.ac.uk/Tools/msa/clustalw2/), and SNPs between the parents and between the NILs were identified manually.

After an SNP was identified, the SNP primer was designed as 18–24 bases ending right before the SNP for genotyping the fine-mapping population. SNP was analyzed using SNaPshot kit (Applied Biosystems, Inc., Foster City, CA) and following the manufacturer’s manual. All primers were checked for possible extendable primer–dimmer formation using PerlPrimer v1.1.9. Fragments were scored using GeneMarker version 1.6 (SoftGenetics, LLC, State College, PA).

**Physical mapping and sequencing of BACs and TaPHS1 gene in wheat accessions**

Five flanking or co-segregating markers were used to screen six-dimensional BAC pools from the Chinese Spring 3AS chromosome-specific BAC library, as reported previously (Safar et al., 2007; Sehgal et al., 2012), to construct the physical map for *Qphs.pseru-3AS*. Each marker was PCR-amplified from the BAC pools, and the positive pools were de-convoluted using in-house scripts to map the markers to individual BACs. Positive BACs were traced to fingerprint assembly, and the contig covering the QTL region was identified. The minimum tiling path BACs spanning the QTL region were identified and sequenced using 454 GS FLX+
System (Roche Company, Branford, CT) as pools. The sequence was assembled by gsAssembler using default parameters. A sequence with 5951 bp was annotated with the ORF of the TaPHS1. To amplify this gene in Rio Blanco and NW97S186, a pair of TaPHS1 gene-specific primers (TaPHS1-GS) were developed using the Chinese Spring group 3 nulli-tetrasomic lines N3AT3B, N3AT3D, N3BT3A, N3BT3D, N3DT3A, and N3DT3B (Supplemental Figure S2A and Table S1). Seven pairs of PCR primers (TaPHS1-P, TaPHS1-17, TaPHS1-18, TaPHS1-19, TaPHS1-20, TaPHS1-21, and TaPHS1-SNPShot) were designed to sequence the genomic DNA of the TaPHS1 in the mapping parents and the 82 accessions for association analysis (Supplemental Table S1). The PCR products were purified with shrimp alkaline phosphatase and exonuclease I and then used for sequencing.

**Gene expression analysis**

RNA was extracted and purified using the RNeasy Plant Kit with on-column DNaseI treatment (Qiagen, Valencia, CA). Complementary DNA from reverse-transcription reaction using a SuperScriptII kit (Invitrogen, Grand Island, NY) was amplified by conventional reverse transcriptase-PCR (RT-PCR). Supplemental Table S1 lists all primer sequences used in the gene expression analysis.

**RNA interference-mediated knockdown of TaPHS1**

An RNAi-based gene-silencing construct was made using the Gateway system described by Miki and Shimamoto (2004). The insert in the silencing construct consisted of a 230 bp fragment from 3’UTR of TaPHS1. The insert DNA fragment was amplified from a cDNA pool synthesized from total RNA extracted from the embryos of Rio Blanco collected 17 days after anthesis using the primers listed in Supplementary Table S1. The fragment was sub-cloned into pENTR/D-TOPO using a directional TOPO cloning kit (Invitrogen, Carlsbad, CA). The final RNAi-based silencing construct was made by recombination from an LR clonase reaction using a Gateway LR clonase enzyme mix (Invitrogen, Grand Island, NY) between the entry vector carrying the TaPHS1 fragment and the pANDA-mini vector. The resulting TaPHS1 silencing construct, PALi7, was bombarded into the cultured immature embryo of two PHS-resistant cultivars, Bobwhite and Rio Blanco, as described by Altpeter et al. (1996).

Transgenic wheat plants were identified by PCR for the presence of the antisense TaPHS1 insert using the gus linker primer, Gus-F1, and the TaPHS1 specific reverse primer, RNAi-R. The sense fragment of TaPHS1 insert was confirmed using gus linker reverse primer Gus-R2 with RNAi-R (Supplemental Table S1). The transformants with both sense and antisense fragments of TaPHS1 gene in RT-PCR were selected, and progenies from three T0 plants (three, two, and three T1 plants lacking or showing weak TaPHS1 expression were evaluated for spike sprouting using non-transformed Bobwhite as control.
**Bioinformatics and statistical analysis**

Genomic sequences from the three BACs covering the *Qphs.pseru-3AS* region were annotated using FGENESH (http://linux1.softberry.com/berry.phtml). The promoter region of *TaPHS1* was analyzed by PlantCare to display the cis-acting regulatory elements (http://bioinformatics.psb.ugent.be/webtools/plantcare/html/). Amino acid sequence alignments were conducted using ClustalW (http://www.ebi.ac.uk/Tools/msa/clustalw2/).

The *TaPHS1* homologs were identified in rice, *Brachypodium*, maize, sorghum, *Arabidopsis*, and barley by BLAST at the Rice Annotation Project (RAP) Database (http://www.jcvi.org/), phytozome (http://www.phytozome.org) and *Arabidopsis* Information Resource (http://www.arabidopsis.org/index.jsp) using the entire deduced amino acid sequence of *TaPHS1*. All statistical analyses were conducted using SAS 9.0 for Windows (SAS Inc, Cary, NC).

Thirty-three unlinked SSR markers covering 21 wheat chromosomes were selected for structure analysis using the STRUCTURE 2.2 (Pritchard et al., 2000). SPAGeDi was used for Kinship analysis (Hardy, et al. 2002). Tassel was used for association analysis (Bradbury et al., 2007) using a threshold of $P < 0.001$ to claim significant association between SNPs and PHS resistance. The protein structures of different *TaPHS1* alleles were predicted using I-TASSER (Zhang et al., 2008).

**Results**

**Comparative fine-mapping of *Qphs.pseru-3AS***

We developed a set of near-isogenic lines (NILs) contrasting at the *Qphs.pseru-3AS* region using a heterogeneous inbred family method (Tuinstra et al., 1997). PHS resistance was evaluated in 1874 F$_2$ plants to identify recombinants in the QTL region. All homozygous NILs harboring the QTL showed a low sprouting rate (2.9%) similar to Rio Blanco (3.5%), whereas all the NILs without the QTL sprouted at a higher rate (94.5%), similar to susceptible parent NW97S186 (95.2%) (Supplemental Figure S1A). The heterozygotes showed a sprouting rate of 20.5%, much lower than the middle value (49.3%) between the two parents. These results confirmed that the partially dominant *Qphs.pseru-3AS* allele in Rio Blanco significantly enhanced PHS resistance. We selected two NILs, 08F485 (resistant) and 08F481 (susceptible), for further gene expression and marker analysis (Figure. 1A & B) and 56 recombinant NILs with a recombination in the QTL region for further fine-mapping (Supplemental Table S2).

Based on the published 3A linkage and physical maps, *Qphs.pseru-3AS* was further located to deletion bin 3AS4-0.45-1.00 at the distal end of the short arm of chromosome 3A (Supplemental Figure S1B, C, & D). A total of 114 expressed sequence tags (ESTs) in this bin were used to develop sequence tag site (STS) markers for fine-mapping. One STS (*XBE423484*) was polymorphic between the parents and between the NILs and mapped in the QTL region (Figure 1C and Supplemental Figure S2B). Comparative sequence analysis of the
EST BE423484 using BLASTN against rice and *Brachypodium* genomes identified a rice homolog, *LOC_Os01g02070*, at the distal end of rice chromosome 1, but the homolog was not found in *Brachypodium* (Figure 1C and Supplemental Table S3). Further analysis using two closely linked rice genes (*LOC_Os01g02060* and *LOC_Os01g02080*) to *LOC_Os01g02070* identified a syntenic region on *Brachypodium* chromosome 2. Among the 45 genes (from *LOC_Os01g01800* to *LOC_Os01g02510*) in the rice syntenic region and 38 genes (from *Bradi2g00780* to *Bradi2g01150*) in the *Brachypodium* syntenic region that were blasted in the wheat EST database, 13 wheat homologous ESTs were mapped to the *Qphs.pseru-3AS* region (Figure 1C, Supplemental Figure S2C & D and Supplemental Table S3). *Qphs.pseru-3AS* was delimited at an interval between STS markers XBE423484 and XAL815375. This interval corresponds to a 35-kb region between *LOC_Os01g02070* and *LOC_Os01g02130* in rice chromosome 1 and a 73-kb region between *Bradi2g00910* and *Bradi2g01030* in *Brachypodium* chromosome 2 (Figure 1D). Three wheat STS markers, XBE401794, XCD910417, and XCA654295, co-segregated with PHS resistance (Supplemental Table S2).

**Physical mapping and determination of the candidate gene for Qphs.pseru-3AS**

The two flanking and three co-segregating markers were then used to screen a Chinese Spring 3AS chromosome-arm specific BAC library. One 210kb contig (Ctg619) containing 22 BACs was found to harbor the QTL (Supplemental Figure S3). A high-confidence minimum tiling path of three BACs (3ASHA_0054M15, 3ASHA_0065J02, and 3ASHB_0159A06) was identified to cover the entire Ctg619. After the three BACs were sequenced, six genes were identified and considered as candidate genes for *Qphs.pseru-3AS* (Figure 1E and Supplemental Table S4).

The expression of the six candidate genes was studied during seed sprouting. Only the wheat gene *TaMFT*, a wheat homolog to the rice gene *LOC_Os01g02120*, was expressed in the PHS-resistant NIL and not in the susceptible NIL (Figure 2A). *MFT* is involved in seed germination of *Arabidopsis* and encodes a phosphatidylethanolamine-binding protein (PEPB) in the embryo (Xi et al., 2010). *TaMFT* was highly expressed only in the embryo of the resistant wheat genotypes during seed development 17 and 26 days after anthesis and between imbition and full seed germination during sprouting (Figure 2B & C). Treating imbibed seeds at low temperature (4°C) for 3 days terminated the gene expression in the resistant genotype (Figure 2A). Among the other five genes (Supplemental Table S4), four showed no differential expression between resistant and susceptible NILs, and the fifth, a transposon gene homologous to *LOC_Os01g02100* in rice, was not expressed during sprouting (data not shown). Therefore, *TaMFT* is the candidate gene for *Qphs.pseru-3AS*.

To validate the function of *TaMFT* for PHS resistance, an RNA interference (RNAi) construct, PALi7 (Figure 3A), was constructed and transformed into PHS-resistant wheat cultivars Rio Blanco and Bobwhite using the particle bombardment mediated method. Bobwhite shares an identical *TaMFT* DNA sequence with Rio Blanco; however, regeneration of transgenic plants was not successful for Rio Blanco. Of the 10
independent T0 transformants from Bobwhite, only three (672C, 838A, and 839A) were confirmed to have both the sense and the anti-sense strands of TaMFT (data not shown). T1 plants from these three transformation events showed significantly reduced endogenous RNA levels of TaMFT but significantly increased sprouting rates: from 2.7% in non-transgenic control to 52.9–66.6% in transgenic Bobwhite plants (Figure 3B, C, & D). These results confirm that TaMFT is the gene underlying the Qphs.pseru-3AS QTL and is responsible for PHS resistance. Because of the confirmed biological function of TaMFT in wheat pre-harvest sprouting resistance, rather than in regulating flower time of Arabidopsis thaliana, we designated it as TaPHS1.

**Structure of TaPHS1**

Genomic sequence of TaPHS1 (5165bp), including a 941 bp promoter sequence, was obtained from Rio Blanco using specific primers designed from the BAC sequences and the Chinese Spring nulli-tetrasomic lines (Supplemental Figure S2A and Table S1). Gene annotation predicted four exons and three introns in TaPHS1 encoding a peptide of 175 amino acids (Figure 4A & E). To verify the coding sequence annotation, primers were designed based on a wheat full-length cDNA, AK330655, from the Triticeae Full-Length CDS database and used to amplify the full-length cDNA from Rio Blanco (Mochida et al., 2009). The resulting sequence is identical to previous coding sequence annotations, and the deduced peptide sequence is homologous to MFT proteins from several species (Supplemental Figure S4).

**Mis-splicing in TaPHS1 causes PHS susceptibility**

Sequence comparison of TaPHS1 alleles between Rio Blanco and NW97S186 identified 31 SNPs or InDels in the promoter region and 32 SNPs in the gene. Two important mutations in Taphs1 of NW97S186 were identified. One was a GT-to-AT transition at the 5’ donor splice-site (position +646) of the intron 3, which extended the exon 3 into intron 3 (Figure 4A & D); the other was an A-to-T transversion at position +666 that generated a premature stop codon and resulted in a truncated protein in NW97S186 (106 amino acids) (Figure 4C & E). These two SNPs are most likely the candidate SNPs for wheat sprouting resistance. The primers designed from different exons of Rio Blanco and NW97S186 were used for RT-PCR. The primers designed from the identical sequence of exons 1 to 3 amplified one band in both Rio Blanco and NW97S186, whereas the primers designed from the fourth exon amplified one band only in Rio Blanco. In contrast, the primers designed from the truncated cDNA sequence of NW97S186 amplified one band only in NW97S186 (Figure 4A & B), which confirmed the extension of exon 3 and truncation of the transcript in NW97S186.

Analysis of the sequences in the promoter region of Rio Blanco identified six cis-acting abscisic acid (ABA) responsive elements (ABREs) that were reported to be involved in ABA responsiveness, two RY-elements involved in seed-specific regulation, one motif IIB, and an ABA responsive element (Baumlein et al.,
Two SNPs, both transversions, between Rio Blanco and NW97S186 were detected in two ABREs at positions -314 and -222.

To further determine the SNPs for PHS resistance in \textit{TaPHS1}, we sequenced both the promoter and gene-coding regions of the gene from 82 wheat cultivars and found 84 SNPs/InDels. Structure analysis using SSR markers from all 21 chromosomes stratified these cultivars into two groups (Supplemental Table S5). Significant associations were found between 20 sequence polymorphisms and sprouting rates ($P < 0.001$), with 8 in the promoter region and 12 in the gene (Figure 5A). The two SNPs at positions +646 and +666 that resulted in the mis-splicing site and the premature stop codon, respectively, were the two most significant SNPs ($P<8.27E-5$); the other 18 significant signals were most likely due to the linkage disequilibrium (LD) of the two SNPs (Supplemental Figure S6). The association between sprouting rate and the two SNPs in the two ABRE cis-acting elements at positions -314 and -222 was not significant (Figure 5A).

Haplotype analysis based on the SNPs (+646 and +666) and the SNP in the promoter region (-222) allowed the 82 cultivars to be divided into three groups: Rio Blanco type (Group I), Chinese Spring type (Group II), and NW97S196 type (Group III). Group I differed from Group II in the SNP at position -222; their sprouting rates were not significantly different, but Group I had a slightly lower sprouting rate than Group II (Figure 5B). The SNPs in Group III differed from Group I and II at both positions +646 and +666, where mis-splicing and the premature stop codon mutations occurred. Group III had significantly higher sprouting rates (79.5%, $P=2.98 \times 10^{-5}$) than groups I and II (Figure 5B), demonstrating that two SNPs at positions +646 and +666 were responsible for the change in PHS resistance. Although red wheat had a slightly lower sprouting rate than white wheat in both haplotypes (Supplemental Figure S7), the two SNPs at +646 and +666 were primarily responsible for the shift from PHS-resistant to susceptible. Thus, the two mutations that cause mis-splicing and the premature stop codon are the SNPs for PHS resistance. We designed two SNP markers, \textit{TaPHS1-SNP1} based on the mis-splicing SNP and \textit{TaPHS1-SNP2} based on the premature stop codon. Because the mis-splicing mutation at position +646 was required for the splicing change and both SNPs appeared together in all cultivars tested, we propose to use \textit{TaPHS1-SNP1} (Supplemental Figure S8 and Table S1) as the diagnostic marker for identifying the resistance allele of \textit{TaPHS1} in breeding.

\section*{Discussion}

In a recent microarray study, \textit{TaMFT} was identified as one of several genes that regulate seed dormancy; it is located in the region with a previously reported QTL for seed dormancy, \textit{QPhs.ocs-3A.1}, from a red wheat (Nakamura et al., 2011). However, whether the gene regulates overall PHS resistance was not determined in that study. In our study, we independently cloned the same \textit{TaMFT} gene using map-based cloning of a QTL (\textit{Qphs.pseru-3AS}) on chromosome 3AS for PHS resistance. Fine-mapping of the QTL was achieved by taking advantage of draft reference genome sequences of rice (http://www.jcvi.org) and the \textit{Brachypodium} and wheat
EST database (Munkvold et al., 2004; The International Brachypodium Initiative, 2010). By measuring the spike-sprouting rate, we directly evaluated overall PHS resistance in the fine-mapping population. We used comparative mapping, physical mapping, BAC sequencing, and candidate gene expression to pinpoint TaMFT as the only gene underlining \textit{Qphs.pseru-3AS}. The result of RNA interference-mediated knockdown of \textit{TaPHS1} confirmed that \textit{TaPHS1} positively regulates overall PHS resistance. Thus, the gene underlining the PHS resistance conditioned by \textit{Qphs.pseru-3AS} is the same gene that regulates seed dormancy as reported by Nakamura et al. (2011).

An SNP in the promoter region (at position -222) of \textit{TaMFT} was previously reported to be the key SNP responsible for the change in seed dormancy level (Nakamura et al., 2011). We observed a high frequency of sequence variation in both promoter and coding regions of \textit{TaPHS1} among wheat cultivars. Gene structure analysis of these SNPs revealed five SNPs (two in the promoter region and three in the gene) that could be potential candidates for the causal SNPs for changing PHS resistance to susceptibility. Two SNPs at positions -222 and -314 of the promoter region are ABREs binding sites; one SNP at position +386 in the second exon leads to one amino acid change; one SNP at the 5' donor splicing site at +646 results in extending exon 3 into intron 3; and one SNP at +666 in intron 3 creates a premature stop codon when the exon 3 is extended. The association between the two SNPs in the promoter region (-222 and -314) and PHS resistance, however, was not significant (Figure 5A). The SNP at position -222 of the promoter region might regulate differences in seed dormancy between PHS-resistant and moderately resistant cultivars at low temperatures (Nakamura et al., 2011). The discrepancy between the two studies is most likely due to different materials used in different studies and different traits evaluated under different conditions. In this study, we evaluated sprouting rate per spike for overall PHS resistance at normal wheat growing temperatures (~21-25 °C) instead of evaluating germination rate after threshing at 13 °C (Nakamura et al., 2011), and we used a population developed from two parents with the largest contrast in sprouting rates. The susceptible cultivar (Chinese Spring) used in the previous study (Nakamura et al., 2011) showed moderate PHS resistance in our study. These results suggested that the same gene conditioned both seed dormancy and PHS resistance on the 3AS chromosome, and other morphological factors didn’t play a significant role in PHS resistance expression in this study. This might be because all tested spikes were provided with sufficient moisture for sprouting in a moist chamber, and the difference in spike structures that attract water for seed germination in natural conditions did not contribute to PHS resistance; thus, PHS resistance was mainly contributed by long seed dormancy in this study. Genetic control of different levels of PHS resistance, however, may be different among different sources, so it is possible that quantitative difference in seed dormancy between highly PHS-resistant and moderately resistant genotypes is regulated by sequence variation in the promoter region of \textit{TaPHS1} (Nakamura et al., 2011), whereas qualitative difference in PHS resistance between resistant and susceptible genotypes is determined by the sequence variation in the coding region through functional change of the gene. The coding region of
TaPHS1 determining qualitative PHS resistance was validated in a panel of 82 wheat cultivars in this study. The difference in germination temperatures between the two studies may also contribute to difference in the gene regulation.

Mis-splicing in Vp1 transcripts in wheat has been shown to cause reduction in Vp1 protein activity, thus compromise the gene function (McKibbin et al., 2002). The mis-splicing at position +646 of TaPHS1 together with the mutation at +666 that forms a premature stop codon results in a nonfunctional transcript, which leads to a change from a PHS-resistant genotype to a susceptible genotype. When all SNPs from a panel of 82 wheat cultivars were analyzed, the highest association occurred between the two SNPs in intron 3 and PHS resistance (Figure 5A), which validated our results. These two key SNPs were not found in the previous study (Nakamura et al., 2011, Figure 5B). Gene expression analysis using the primers designed from different exons of both resistant and susceptible genotypes confirmed the extension of exon 3 and truncation of exon 4 in the TaPHS1 transcript of the susceptible genotypes. Thus, these newly identified SNPs are the key mutations for changing PHS resistance and together cause the loss in function of TaPHS1, changing wheat from PHS-resistant to susceptible. Based on these results, we designed two SNP markers, TaPHS1-SNP1 and TaPHS1-SNP2, at the two mutation sites for screening of TaPHS1 in breeding. The two mutations always come together in all wheat germplasm screened to date, so TaPHS1-SNP1 for the mutation at the mis-splicing site suffices for diagnosing the presence of TaPHS1 in wheat breeding.

Red wheat usually shows a higher level of PHS resistance than white wheat. Red color pericarp/testa has been used as a genetic marker for selecting PHS resistance (Gale and Lenton, 1987; Groos et al., 2002; Himi et al., 2002). Several other DNA markers for grain color genes have been developed recently (Himi et al., 2011). In this study, both parents of the mapping populations are white wheat, suggesting that major differences in PHS resistance are independent of grain color. In the association panel, both resistant white wheat and highly susceptible red wheat were identified, so this study indicated that TaPHS1 is much more important for PHS resistance than seed color genes are. However, seed color may modify PHS resistance that is regulated by TaPHS1. In either group containing the resistance or susceptibility allele of TaPHS1, the red wheat subgroup had a slightly lower sprouting rate than the white wheat subgroup, although the differences were not significant (Supplemental Figure S7). This suggested that TaPHS1 is the major gene determining PHS resistance (qualitative), although seed color genes may modify the level of PHS (quantitative) in either PHS-resistant or susceptible genotype groups as determined by TaPHS1. Thus, TaPHS1 is a highly valuable PHS resistance gene for breeding white wheat cultivars.

Global climate change with unstable temperatures and rainfall threatens wheat production. Developing PHS-resistant cultivars that can adapt to unpredictable environments is essential to reduce losses from grain sprouting in fields under wet weather conditions. The characterization and functional determination of TaPHS1 as the PHS resistance gene for QTL Qphs.pseru-3AS in white wheat significantly advances the understanding of
the mechanisms underlying PHS resistance in wheat, and identifying causal SNPs and developing the diagnostic markers for TaPHS1 and should facilitate effective manipulation of PHS resistance to improve grain yield and quality in wheat.

**Acknowledgements**

We thank Hyeonju Lee and Dehlia McAfee from the Kansas State University Plant Pathology Department for technical assistance in wheat transformation and Dr. Graybosch, USDA/ARS in Lincoln, NE, for providing original recombinant inbred line population. This project was partially supported by the National Research Initiative Competitive Grants CAP project 2011-68002-30029, and NRICG grants 2006-35604-17248 and 2008-35300-04588 from the National Institute of Food and Agriculture of the US Department of Agriculture. Mention of trade names or commercial products in this article is solely for the purpose of providing specific information and does not imply recommendation or endorsement by the US Department of Agriculture. USDA is an equal opportunity provider and employer. Contribution number 13-277-J from the Kansas Agricultural Experiment Station.

**Literature Cited**


Hardy, O.J., X. Vekemans, 2002 SPAGeDi: a versatile computer program to analyse spatial genetic structure at the individual or population levels. Mol Ecol Notes **2**: 618-620.


Kuraparthy, V., S. Sood S, B. S. Gill, 2008 Genomic targeting and mapping of tiller inhibition gene (*tin3*) of wheat using ESTs and synteny with rice. Funct Integr Genomics **8**: 33–42.


Figure 1. Map-based cloning of QTL (Qphs.pseru-3AS) for pre-harvest sprouting (PHS) resistance. (A) PHS phenotypes of different genotypes after one week in a moist chamber. The top picture shows wheat spikes of resistant (Rio Blanco) and susceptible parents (NW97S186), and resistant (08F485) and susceptible (08F481) near-isogenic lines (NILs). The bottom graph shows the difference in spike sprouting rates (%) with standard deviations for different genotypes. (B) Spike sprouting rates for both parents and NILs evaluated from harvesting date to 12 weeks after harvest. Wheat spikes were stored at room temperature for different weeks before moist treatment. (C) Comparative maps of the Qphs.pseru-3AS region across wheat chromosome 3A, Brachypodium chromosome 2, and rice chromosome 1. The green arrow indicates the location of TaPHS1. The blue line connects the other syntenic genes in Brachypodium, and the red line connects rice to wheat homologs in the Qphs.pseru-3AS region. The wheat chromosome 3A fine map was generated using a high resolution mapping population from the analysis of 1874 F2 plants using three SSR markers Xbarc12, Xbarc57, and Xbarc321; the other markers are added to the map using the high resolution mapping population. The numbers below the linkage map indicate the number of recombinants between the intervals of two markers on the two bars. (D) Graphical genotypes of the Qphs.pseru-3AS region in 10 near-isogenic recombinants with unique recombination among markers (left) and graphical phenotypes (10 plants for each line) showing their sprouting levels (right). Solid and open bars in the genotypic graph represent chromosomal segments from Rio Blanco.
and NW97S186. “R” represents sprouting resistant and “S” represents sprouting susceptible. (E) Six candidate genes, including *TaMFT* (first on the right), were identified after sequencing the three BACs covering the *Qphs.pseru-3AS* region.

**Figure 2.** Expression of all the genes in the *Qphs.pseru-3AS* region during spike sprouting determined the wheat MFT homolog (*TaMFT*) as *TaPHS1*. (A) *TaMFT* was amplified in the PHS resistant (R) NIL, but not in the susceptible (S) NIL, during sprouting. RNA was extracted from the embryos of seeds harvested from the spikes at 0 hour, 1 day, 3 days, and 7 days after imbibing. “All germinated” embryos were obtained by incubating the spikes at 4°C for 3 days after imbibing and then keeping them at room temperature for 1 day. (B) *TaPHS1* is expressed only in the embryo of the resistant (R) NIL after 10 days and before physical maturity during seed development, not in the susceptible (S) NIL. (C) *TaPHS1* expression occurs only in the embryo, not in other tissues at 15 days after anthesis. Actin was used as control. *TaPHS1*-CDS refers to the primer pair P1+P6 that were used in RT-PCR and designed from the full-length cDNA sequence of Rio Blanco (Supplemental Table S1).
Figure 3. Confirmation of TaPHS1’s function on PHS resistance by RNA interference (RNAi). (A) Structure of the RNAi construct, PALi7. The gray arrow shows that the fragment in 3’-UTR of Rio Blanco TaPHS1 was introduced into the construct as inverted repeats. (B) Representative spikes from PHS resistant cultivar Bobwhite and three Bobwhite RNAi T1 transgenic lines at 5 days in a moist chamber. (C) Reverse transcription PCR detected PALi7 only in the transgenic T1 lines, not in the non-transgenic control; in contrast, TaPHS1 strongly expressed in non-transgenic control but not in its RNAi lines. RNA was extracted from the embryo 1 day after the spikes were put into a moist chamber. (D) Spike sprouting rates of non-transgenic Bobwhite (Cont) and its T1 lines from the three transgenic events with standard deviations.
**Figure 4.** Gene and protein structures of *TaPHS1* in Rio Blanco (R) and NW97S186 (S). (A) Genomic sequence and the full-length cDNA sequences of *TaPHS1* from both parents. Boxes represent exons; the green box shows the extension of exon 3 caused by mis-splicing and truncation of exon 4 in NW97S186. Lines represent introns and 5' and 3' UTR; Arrows point to two critical mutation sites (mis-splicing site and the premature stop codon) in NW97S186. P1 to P6 were the primers used to confirm the truncated mRNA in NW97S186. (B) The two critical mutations in intron 3 of NW97S186 were validated by RT-PCR using four pairs of primers (see A). (C) The predicted disruption of the protein structure of the *TaPHS1* in NW97S186, not in Rio Blanco. (D) Partial sequences of *TaPHS1* show the two key mutations that form mis-splicing site and a premature stop codon in intron 3 of NW97S186. Arrows point to the key mutation sites, and * indicates SNPs between two parents. (E) Deduced amino acid sequences of *TaPHS1*. * points polymorphic amino acids between parents.
Figure 5. Association analysis to validate two causal SNPs in intron 3 of TaPHS1 for PHS resistance. (A) Twenty SNPs were significantly associated with PHS resistance at $P < 0.001$. The SNP positions are based on TaPHS1 sequence of Rio Blanco with the start codon as 0. Red arrows point to the two SNPs causing the mis-splicing and the premature stop codon in intron 3 and the one in the promoter region that was previously reported as the causal SNP (Nakamura et al. 2011). (B) Mean sprouting rates of three contrasting haplotype groups according to the three SNPs, where haplotype I represent 20 cultivars that shares the same haplotype with Rio Blanco; haplotype II represent 46 cultivars that shares the same haplotype with Chinese Spring that has one mutation at the promoter region (Nakamura et al. 2011), and haplotype III represent 16 cultivars that shares the same haplotype with NW97S186 that has all mutations at three sites. Error bar denotes standard deviation. Different letters, a and b, indicate significant difference at $P < 0.05$. 

<table>
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Supplemental data

The following materials are available in the online version of this article.

Supplemental Figure S1. Linkage and physical maps of Qphs.pseru-3AS. (A) Segregation of sprouting resistance in the near-isogenic recombinant population. Plants with Rio Blanco genotypes at the three marker loci, Xbarc321, Xbarc57, Xbarc12, were highly sprouting-resistant; the heterozygous plants were moderately resistant; plants with all three NW97S186 marker alleles were highly susceptible. Error bar denotes standard deviation, and a, b and c indicate significant difference at $P<0.01$. (B) The QTL map of Qphs.pseru-3AS was developed using two recombinant inbred populations derived from Rio Blanco/NW97S186 and Rio Blanco/NW97S078 (17). The green bar represents the QTL interval on the short arm of chromosome 3A (3AS). (C) The four SSR markers (Xbarc321, Xbarc57, Xbarc12, and Xgwm369) close to Qphs.pseru-3AS show similar order and genetic distance in another map (Song et al., 2005). (D) Physical map of 3A (Sourdille et al., 2004). Two of the four SSR markers (Xbarc12, Xgwm369) close to Qphs.pseru-3AS are located in the deletion bin 3AS4-0.45-1.00 at the distal end of 3AS.

Supplemental Figure S2. Markers used to determine the chromosome location of Qphs.pseru-3AS. (A) An agarose gel image shows TaPHS1 specific primers amplified only in Chinese Spring nulli-tetrasomic lines carrying 3A chromosome. No PCR product was amplified when the 3A chromosome was replaced by 3B (N3A-T3B) or 3D (N3A-T3D). (B) An agarose gel image shows an STS marker developed from the wheat EST BE423484 and detected polymorphism between the resistant genotypes (Rio Blanco and 08F485) and the susceptible genotypes (NW97S186 and 08F481). (C) An electrophorogram of polymorphic STS marker developed from the EST CA654295 analyzed using capillary electrophoresis in an ABI 3730 DNA analyzer. (D) An electrophorogram of an SNP marker developed by re-sequencing wheat EST CD910417 analyzed using SNPSht in an ABI 3730 DNA analyzer.

Supplemental Figure S3. BAC contig Ctg619 (bottom solid bar) spans the entire Qphs.pseru-3AS region. Top bar is the linkage map of Qphs.pseru-3AS with the Qphs.pseru-3AS region labeled by red oval. Solid bars between the linkage map and the contig bar are different BACs in the contig. Three BACs with red color that cover the entire QTL region were selected for sequencing to identify candidate genes for TaPHS1.

Supplemental Figure S4. Sequence alignment of the deduced amino acids of TaPHS1 homologs from maize (GRMZM2G021614, GRMZM2G152689, GRMZM2G059358), sorghum (Sb03g8270), Brachypodium (Bradi2g01020), barley (HvMFT), rice (Os01g02120, Os06g30370), Arabidopsis (At01g18100), and wheat (Qphs.pseru-3AS).

Supplemental Figure S5. Comparison of the promoter sequences between Rio Blanco and NW97S186. Identical sequences are labeled by *. ABRE CEs are indicated by blue or yellow boxes if two ABRE CEs
overlapped. RY repeats are indicated by green letters. Motif IIB is underlined. Two SNPs in the ABRE CEs at -222 and -314 positions are shown in red letters.

Supplemental Figure S6. Linkage disequilibrium (LD) between the two causal SNPs (+646 and +666) in intron 3 and other polymorphic sites of TaPHS1 identified in 82 wheat cultivars. The upper part is the gene structure of TaPHS1, and the lower part is the LD graph. From left to right, each grid indicates one SNP or InDel of the 84 SNPs or InDels in order from 5’-end of TaPHS1 across the 82 cultivars. Among the 84 SNPs, only 20 SNPs or InDels that were significantly associated with PHS resistance were labeled on the graph, including two causal SNPs (+646 and +666) in intron 3 and the 18 non-causal polymorphic sites (Figure 5A) associated with sprouting resistance due to their LD to the two causal SNPs.

Supplemental Figure S7. Two causal SNPs determine PHS resistance. White wheat had slightly higher spike sprouting rates than red wheat in both haplotypes of TaPHS1 based on the two causal SNPs loci (+646 and +666), but the difference was not significant (a and b indicate significant difference at P < 0.05) as the two causal SNPs (P=2.98 x 10^-5, Figure 5B).

Supplemental Figure S8. One diagnostic SNP marker for TaPHS1 (TaPHS1-SNP1) was developed based on the GT-to-AT mutation at the mis-splicing site (+646) in susceptible cultivars. The SNP was analyzed using SNaPshot kit (Life Technology, Grand Island, NY) via an ABI3730 DNA analyzer. The top four cultivars with blue (G) peaks are sprouting-resistant, and the bottom four cultivars with green (A) peaks are susceptible.

Supplemental Table S1 Primers used for sequencing, gene expression, and gene transformation
Supplemental Table S2 Genotypic data and sprouting resistance levels of the 56 recombinants analyzed with the markers in the QTL Qphs.pseru-3AS region
Supplemental Table S3 Thirteen genes that were mapped in the QTL Qphs.pseru-3AS region by comparative mapping
Supplemental Table S4 Genes identified by sequencing the three BACs spanning the Qphs.pseru-3AS region
Supplemental Table S5 Relationship among two structure groups, seed color, and causal SNPs for PHS resistance in TaPHS1.