

# Evolutionary Expansion, Gene Structure, and Expression of the Rice Wall-Associated Kinase Gene Family<sup>1[w]</sup>

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The wall-associated kinase (WAK) gene family, one of the receptor-like kinase (RLK) gene families in plants, plays important roles in cell expansion, pathogen resistance, and heavy-metal stress tolerance in Arabidopsis (*Arabidopsis thaliana*). Through a reiterative database search and manual reannotation, we identified 125 OsWAK gene family members from rice (*Oryza sativa*) japonica cv Nipponbare; 37 (approximately 30%) OsWAKs were corrected/reannotated from earlier automated annotations. Of the 125 OsWAKs, 67 are receptor-like kinases, 28 receptor-like cytoplasmic kinases, 13 receptor-like proteins, 12 short genes, and five pseudogenes. The two-intron gene structure of the Arabidopsis WAK/WAK-Likes is generally conserved in OsWAKs; however, extra/missed introns were observed in some OsWAKs either in extracellular regions or in protein kinase domains. In addition to the 38 OsWAKs with full-length cDNA sequences and the 11 with rice expressed sequence tag sequences, gene expression analyses, using tiling-microarray analysis of the 20 OsWAKs on chromosome 10 and reverse transcription-PCR analysis for five OsWAKs, indicate that the majority of identified OsWAKs are likely expressed in rice. Phylogenetic analyses of OsWAKs, Arabidopsis WAK/WAK-Likes, and barley (*Hordeum vulgare*) HvWAKs show that the OsWAK gene family expanded in the rice genome due to lineage-specific expansion of the family in monocots. Localized gene duplications appear to be the primary genetic event in OsWAK gene family expansion and the 125 OsWAKs, present on all 12 chromosomes, are mostly clustered.

Efficient communication between the plant cell wall and the cytoplasm is important in plant development and in responding to biotic and abiotic stresses (Kohorn, 2000; Brownlee, 2002; Somerville et al., 2004). The wall-associated kinase (WAK) gene family, which belongs to the receptor-like kinase (RLK) superfamily in plants (Shiu and Bleecker, 2001, 2003), plays a critical role in this communication (He et al., 1996; Kohorn, 2001). WAK1, the first member of the WAK gene family identified in Arabidopsis (*Arabidopsis thaliana*), encodes a protein containing an intracellular Ser-Thr kinase domain and extracellular domains with similarities to vertebrate epidermal growth factor (EGF)-like domains (He et al., 1996). From the Arabidopsis genome (Arabidopsis Genome Initiative, 2000),

26 WAK and WAK-Like (WAKL) genes were identified and most of the 26 members are expressed in Arabidopsis (Verica and He, 2002; Verica et al., 2003).

Functional studies of the different WAK members in Arabidopsis demonstrated that they are involved in various functions in plants, including pathogen resistance (He et al., 1998), heavy-metal tolerance (Sivaguru et al., 2003), and plant development (Lally et al., 2001; Wagner and Kohorn, 2001). Biochemical studies demonstrated that WAK proteins are covalently bound to pectin in the cell wall (Wagner and Kohorn, 2001). They can also form an approximately 500-kD protein complex via interactions with a Gly-rich extracellular protein, AtGRP-3 (Park et al., 2001), and can interact with a cytoplasmic, type 2C kinase-associated protein phosphatase (Anderson et al., 2001). Based on these attributes, it was suggested that WAKs serve as physical links between the extracellular matrix and the cytoplasm and also as a signaling component between the cell wall and the cytoplasm (He et al., 1996; Kohorn, 2001). Recently, a study of seven (group II) WAKL members in Arabidopsis showed that they have tissue-specific and developmentally regulated expression patterns and are functionally similar to WAKs (Verica et al., 2003).

To further understand the functions and evolution of the WAK gene family in plants, we analyzed the WAK gene family in rice (*Oryza sativa*). Rice is one of the most important cereals and also considered a

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model for other cereal species, including maize (*Zea mays*), wheat (*Triticum aestivum*), barley (*Hordeum vulgare*), and sorghum (*Sorghum bicolor*). Rice genomic sequences are now available from both subspecies *indica* (Yu et al., 2002) and *japonica* (Feng et al., 2002; Goff et al., 2002; Sasaki et al., 2002; Rice Chromosome 10 Sequencing Consortium, 2003; International Rice Genome Sequencing Project, 2005). Previous analyses of the RLK superfamily from the *indica* genome sequence indicated that the WAK/WAKL gene family was one of the few RLK subfamilies that expanded in rice compared to Arabidopsis (Shiu et al., 2004).

Here we present a detailed analysis of the rice WAK/WAKL (*OsWAK*) gene family from *japonica*. First, through public database searches, we retrieved all genes annotated as putative *OsWAKs*, followed by reiterative database searches from which we obtained additional putative *OsWAK* gene family members. Manual reannotation was performed to correct or reannotate the misannotated putative *OsWAK* genes, such as split genes, fused genes, short genes, and pseudogenes. We determined expression characteristics for certain *OsWAK* members and, based on domain composition, we classified *OsWAKs* into five groups and performed comparative phylogenetic analyses of WAKs in Arabidopsis, rice, and barley to understand the possible mechanisms of gene family expansion.

## RESULTS

### Identification and Classification of *OsWAKs* from Genome Sequences of Rice Subsp. *japonica* cv Nipponbare

Four analytical steps were used to identify and classify *OsWAKs* from *japonica* subsp. cv Nipponbare. First, all genes annotated as putative *OsWAKs* were retrieved from three public databases: (1) The Institute for Genomic Research (TIGR) Rice Genome Annotation Database (Osa1, release 1 and 2; Yuan et al., 2005); (2) Rice Protein Database in GRAMENE (Ware et al., 2002); and (3) GenBank in the National Center for Biotechnology Information (<http://www.ncbi.nlm.nih.gov>). There were different numbers of putative *OsWAKs* annotated in the different databases. Identical sequences from the same or overlapping contigs, but annotated with different putative *OsWAK* names in the various databases, were identified and removed. The second step, aimed at a more complete search for putative *OsWAKs* in rice, was performed using BLASTp and tBLASTn ( $E < 0.1$ ) searches of Osa1 and the *japonica* cv Nipponbare genome sequences in GenBank. These analyses utilized 10 uniquely different putative *OsWAK* protein sequences (2768.t00013, 2541.t00004, 8323.t00010, 4577.t00017, 9637.t02571, 4967.t00013, 3135.t00009, gi|19881772, 9638.t00862, and 6563.t00002), which contained both the extracellular EGF-like domains (EGF-2, PS01186/PROSITE and/or EGF-Ca<sup>2+</sup>, PS01187/PROSITE) and an intracellular protein kinase domain.

Most new genes obtained from this reiterative database search were originally annotated either as kinase domain-containing protein genes or receptor-like protein (RLP) genes.

In the third step, each putative *OsWAK* sequence was manually assessed for the EGF-like domains, kinase domain, or its sequence similarity to other putative *OsWAKs*. In our analyses, in order for a gene to be defined as a putative *OsWAK*, the gene had to fit into one of the five *OsWAK* types (see Table I). Based on these five criteria, a total of 129 putative *OsWAKs* were initially identified from *japonica* cv Nipponbare.

As the last step, manual reannotation was performed to correct or reannotate the misannotated putative *OsWAKs*, as described below. This analysis included 10 putative split *OsWAKs* merged into five *OsWAK-RLKs*, four fused putative *OsWAKs* split into five different *OsWAKs*, and six non-*OsWAKs*, leaving a final total of 125 *OsWAKs*; the numbers of each type of *OsWAK* are shown in Table I. For the 125 *OsWAKs*, the detailed annotations (Supplemental Tables I and II), genomic and predicted coding/full-length (FL)-cDNA sequences (Supplemental Data 1), and predicted protein sequences (Supplemental Data 2) are presented.

### Correction and Reannotation of Misannotated Putative *OsWAKs*

Automated annotations of rice genome sequences of *japonica* cv Nipponbare in TIGR and GenBank are useful as initial resources for gene annotation, but individual genes and gene families are generally not verified and corrected. Several types of annotation errors were found in the automated annotation of the Arabidopsis genome, including intron numbers/positions, merged/split genes, and missed short genes;

**Table I.** Classification of 125 *OsWAKs* identified from rice genome of *japonica* cv Nipponbare

Gene Type	Domain Content	No. in Category
<i>OsWAK-RLK</i>	Both extracellular EGF-like domains and a cytoplasmic protein kinase domain	67
<i>OsWAK-RLCK</i>	Only cytoplasmic protein kinase domain with its amino acid identity $\geq 40\%$ to a kinase domain of an <i>OsWAK-RLK</i> member	28
<i>OsWAK-RLP</i>	Only extracellular EGF-like domains	13
<i>OsWAK</i> short gene	No domain, protein sequence <300 amino acids but its amino acid identity $\geq 40\%$ to a longer <i>OsWAK</i> protein	12
<i>OsWAK</i> pseudogene	Stop codons or frame shifts in coding region	5

approximately 35% of the initially annotated genes from *Arabidopsis* were corrected from FL-cDNA sequences (Haas et al., 2003). We therefore performed a similar assessment of the initially identified, 129 putative *OsWAKs* derived from automated annotation using the rice FL-cDNA sequences in the Knowledge-Based *Oryza* Molecular Biological Encyclopedia database (Kikuchi et al., 2003; <http://cdna01.dna.affrc.go.jp/cDNA>). For 31 putative *OsWAKs*, one corresponding FL-cDNA sequence was obtained; for eight other *OsWAKs*, more than one corresponding FL-cDNA sequence was obtained (Supplemental Table III). Of the 39 putative *OsWAKs* that matched with FL-cDNAs, only 20 had predicted coding sequences that perfectly or nearly perfectly matched the corresponding FL-cDNA sequences (identity  $\geq 99.5\%$ ). For the other 19, the FL-cDNA sequences were either shorter (seven) or longer (12) than the predicted coding sequences. All of the FL-cDNA sequences, except one, have both 5' untranslated region (UTR) and 3' UTR, indicating they are most likely intact cDNA sequences. Therefore, these 19 putative *OsWAKs* were corrected using their corresponding FL-cDNA sequences, although it is possible that some or all of the shorter FL-cDNAs could derive from truncated sequences. Reannotation of 14 out of the 19, along with 13 additional putative *OsWAKs*, is described below.

#### *Split/Fused OsWAK Genes*

From the FL-cDNA sequence analysis described above, two putative *OsWAK-RLPs*, *OsWAK30* and *OsWAK31*, matched to the same FL-cDNA sequence (AK058435). By examining matched regions, the two *OsWAK-RLPs* were found to match complementally to the FL-cDNA sequence (*OsWAK30* on the 5' side and *OsWAK31* on the 3' side). Also, both putative genes were located within an approximately 12-kb genomic region on chromosome 4 and no extra genes were predicted in the region between the two putative *OsWAKs* (Fig. 1). Therefore, these two putative, split *OsWAK-RLPs* were merged into a single *OsWAK-RLP*, *OsWAK30/31*. Based on similar analyses, eight more putative, split *OsWAKs* (four putative *OsWAK-RLPs* plus four putative *OsWAK-receptor-like cytoplasmic kinases [RLCKs]*) were merged into four *OsWAK-RLKs* (*OsWAK7/8*, *OsWAK57/58*, *OsWAK96/97*, *OsWAK109/110*). Merging was based on the fact that each pair matched complementally to the same *OsWAK-RLK* and different pairs matched to different *OsWAK-RLKs* (Fig. 1). They were split in the initial annotations at an intron between the two EGF-like domains within the extracellular region.

Fourteen other putative, split *OsWAKs* (seven putative *OsWAK-RLPs* and seven putative *OsWAK-RLCKs*) also matched complementally to various *OsWAK-RLKs* (data not shown). However, for each of these pairs, one to eight different genes interrupt the sequence (Fig. 2). These split *OsWAKs* appear to have already become independent genes; for example, the

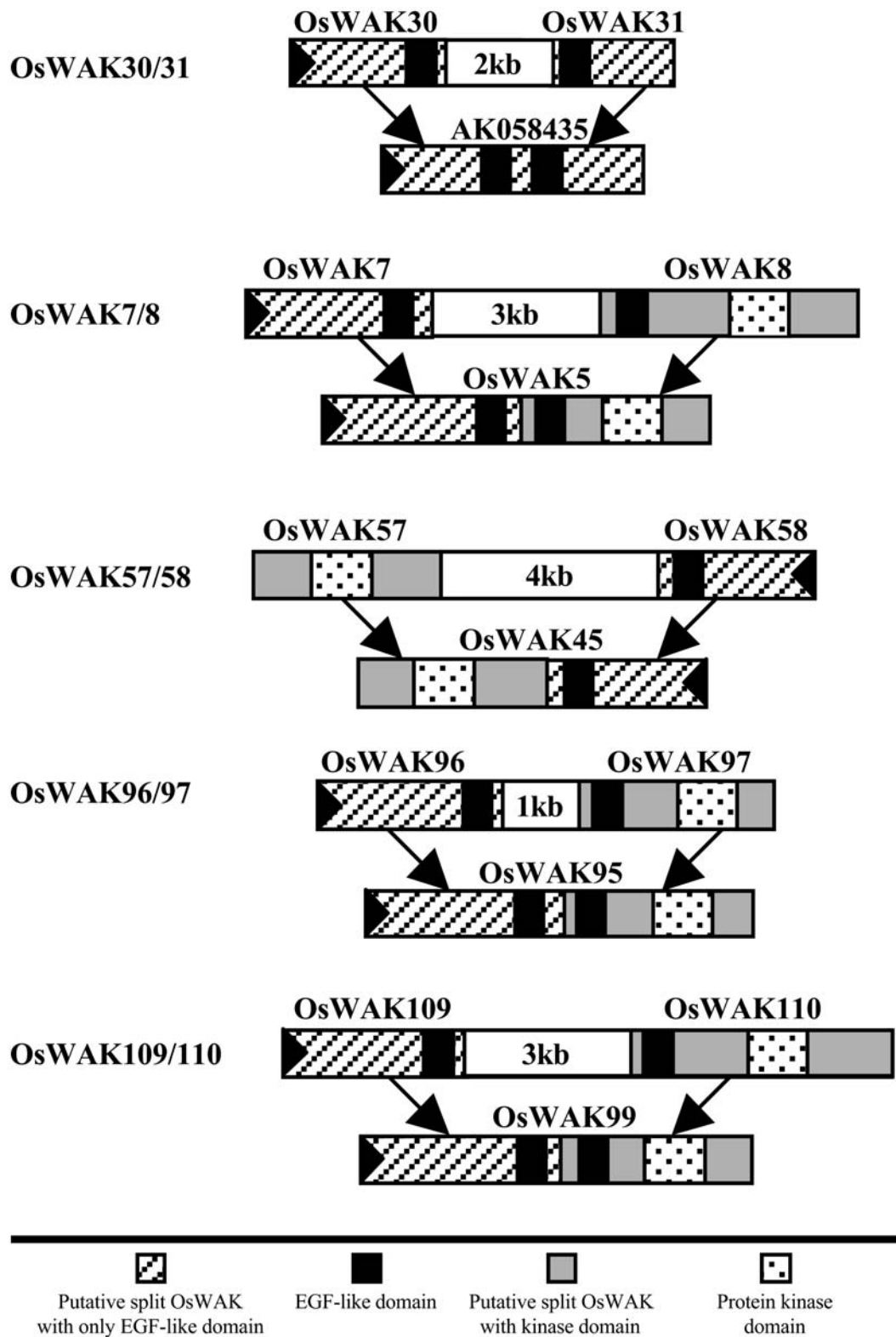
pair, *OsWAK84* and *OsWAK85*, has two different FL-cDNA sequences. Therefore, these 14 putative, split *OsWAKs* remain annotated as independent *OsWAKs*.

Three putative *OsWAKs* (*OsWAK5/2768.t00008*, *OsWAK73/9636.t03851*, *OsWAK89/9637.t03234*) from *Osa1* in TIGR were found to be three fused genes. Putative *OsWAK5* and *OsWAK73* are fused with non-*OsWAKs*, and the third has two fused *OsWAKs* (Fig. 3). Therefore, the three fused genes were split, obtaining four *OsWAKs* (*OsWAK5*, *OsWAK73*, *OsWAK89a*, *OsWAK89b*). *OsWAK89a* and *OsWAK89b* have 57% identity in overall protein sequence and  $>93\%$  identity in their kinase domains; both contain EGF-like domains in their extracellular regions, suggesting they are independent *OsWAK-RLKs* that likely derived from tandem duplication. Interestingly, putative *OsWAK69* is fused to a Pack-Mu-like element (MULE; Jiang et al., 2004) on its 5' side, and an approximately 2-kb space is found between the Pack-MULE and the kinase domain. The sequence of a FL-cDNA (AK064911) matched only to the Pack-MULE, not to the kinase domain (Fig. 4). Accordingly, the fusion is believed to be an annotation artifact and the kinase domain of the original *OsWAK69* was split out as an *OsWAK-RLCK*.

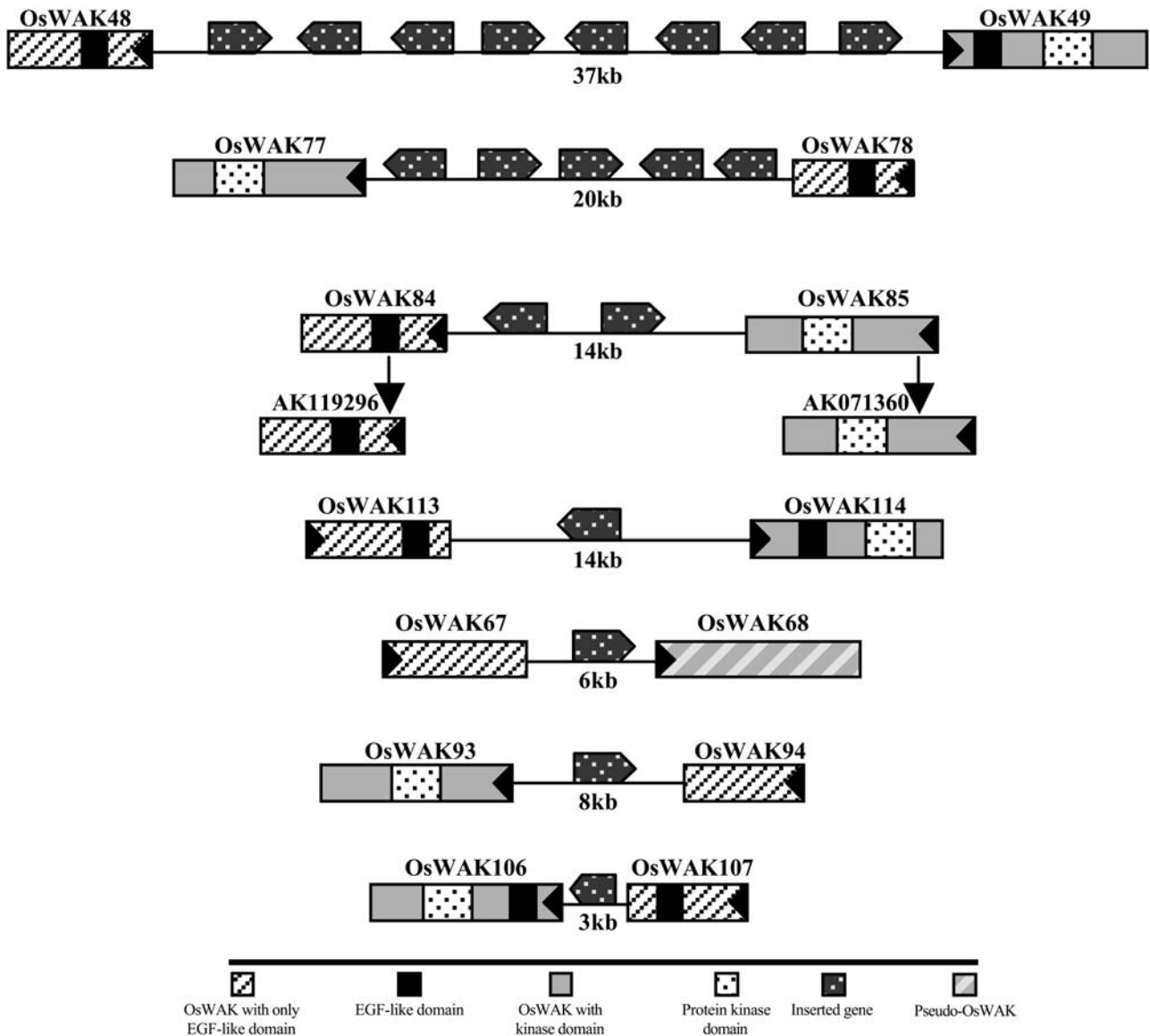
#### *OsWAK Short Genes*

Five putative *OsWAKs* (*OsWAK17*, *OsWAK19*, *OsWAK54*, *OsWAK62*, *OsWAK67*) were initially identified as *OsWAK* short genes. This was due to the fact that their predicted protein sequences were  $<300$  amino acids and did not encode an EGF-like or kinase domain, but they had  $>40\%$  amino acid identity to the longer *OsWAKs*. From the FL-cDNA sequence analyses described above, seven more putative *OsWAKs* (*OsWAK18*, *OsWAK23*, *OsWAK35*, *OsWAK37*, *OsWAK52*, *OsWAK101*, *OsWAK127*) were reannotated as *OsWAK* short genes. This was also due to the fact that the deduced protein sequences from their longest open reading frames (ORFs) are  $<300$  amino acids and do not encode an EGF-like or kinase domain. Of the seven *OsWAK* short genes, four (*OsWAK18*, *OsWAK52*, *OsWAK101*, *OsWAK127*) had antisense transcripts and the other three (*OsWAK23*, *OsWAK35*, *OsWAK37*) had sense transcripts. In order to further understand how the *OsWAK* short genes might have arisen in the rice genome, genomic sequences adjacent to the seven *OsWAK* short genes were analyzed. These analyses found various transposable elements (TEs) were inserted at either the 5' or 3' ends of the FL-cDNAs of five of the *OsWAK* short genes (Table II).

Further analyses indicated that the inserted TEs likely did not interrupt the *OsWAKs*; however, they either provided novel splicing sites or initialized antisense transcriptions, which resulted in short ORFs for the *OsWAKs*. For example, the 3' end of the *OsWAK23* FL-cDNA is located inside a long interspersed nuclear element (LINE) that appears to provide a novel splicing acceptor site. As a result, both the EGF-like and kinase domains were spliced out and thus do not present in



**Figure 1.** Five *OsWAK*s merged from five pairs of putative split *OsWAK*s. Each pair of putative split *OsWAK*s is shown to be split at an intron in the original annotation; size of the intron is indicated in kilobase pairs. Each pair complementarily matched either to a FL-cDNA or an *OsWAK* gene family member that is displayed below each pair; the corresponding matched regions are indicated by black arrows; large black triangles in boxes indicate direction of transcription.



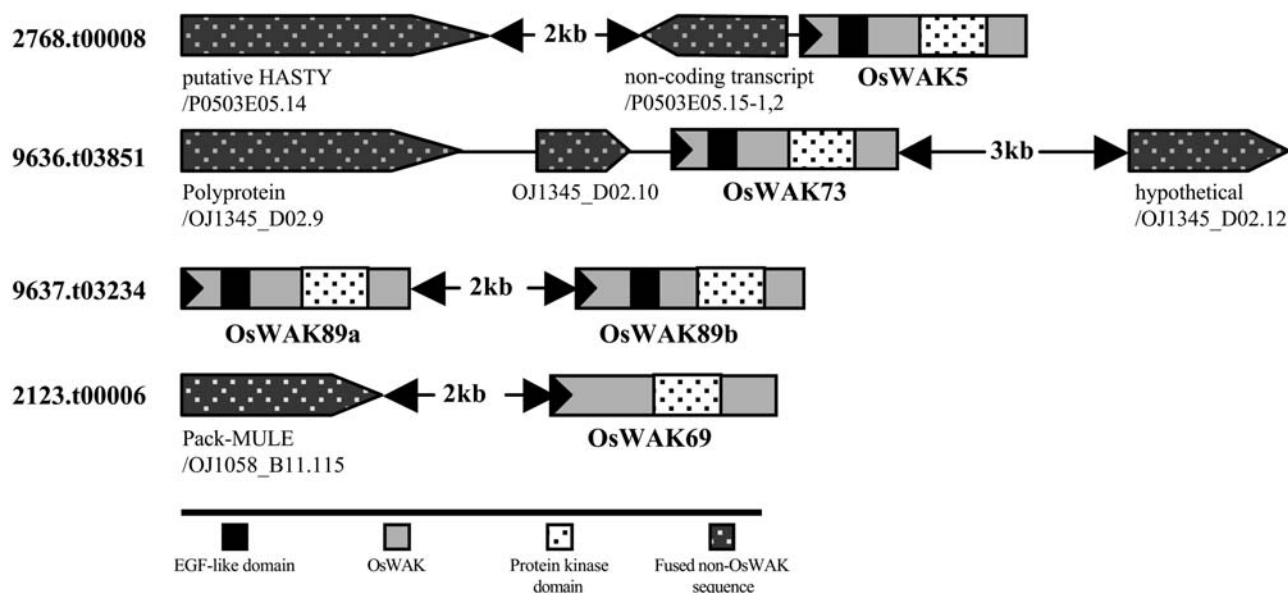
**Figure 2.** Genomic organization of 14 split *OsWAK*s. Each pair of *OsWAK*s is interrupted by one to eight different genes as indicated. The pair, *OsWAK84* and *OsWAK85*, matched two different FL-cDNA sequences, as indicated by vertical arrows. Large black triangles in boxes indicate direction of transcription of *OsWAK*s; distance between each pair of *OsWAK*s is indicated in kilobase pairs.

the FL-cDNA sequence of *OsWAK23* (Fig. 4). *OsWAK35* has one sense and three antisense transcripts, and one of the antisense transcripts (AK100568) appears to be initialized from a solo long-terminal repeat (LTR; Fig. 4).

#### *OsWAK Pseudogenes*

Twelve putative *OsWAK*s were initially annotated as pseudogenes in GenBank because of the apparent lack of the 5' exon, when compared to other *OsWAK-RLK*s, or the presence of stop codons or frame shifts in their coding regions (Supplemental Table II). However, four (*OsWAK21*, *OsWAK76*, *OsWAK82*, *OsWAK84*) had the

corresponding FL-cDNA sequences available (Fig. 5), indicating they were transcribed in rice. Since there are stop codons in the coding regions, it is possible that they are nonfunctional. However, the ORFs derived from the FL-cDNAs contained either a kinase domain (*OsWAK21*, *OsWAK76*, *OsWAK82*) or the EGF-like domains (*OsWAK84*), suggesting they could also be functional in rice. Even though further analysis is needed to verify their status, we have reannotated them as *OsWAK*s. Three other *OsWAK*s (*OsWAK3*, *OsWAK6*, *OsWAK80*) were initially annotated as pseudogenes because the 5' exon was missing; however, truncation does not always lead to loss of function (see "Discussion"). As a result, *OsWAK3* is reannotated as



**Figure 3.** Five *OsWAKs* split from four fused putative *OsWAKs*. The five *OsWAKs* are illustrated with boxes; large black triangles in boxes indicate direction of transcription. Fused non-*OsWAK* sequences are indicated (see key). Annotations with the GenBank accession number are shown below each non-*OsWAK* sequence; distances between fused genes are indicated in kilobase pairs, except in cases where the distance is less than 1 kb.

an *OsWAK-RLCK* because it contains a kinase domain. *OsWAK6* (524 amino acids) is shorter than other *OsWAK-RLKs* (>700 amino acids); however, it encodes an EGF-like domain and a kinase domain and was reannotated as an *OsWAK-RLK* member. For *OsWAK80*, the missing 5' exon was recovered by extending the 5' side from the bacterial artificial chromosome (BAC) sequence. These three putative pseudogenes, plus the four putative pseudogenes described above, were reannotated as true *OsWAKs*, leaving five annotated as *OsWAK* pseudogenes (*OsWAK66*, *OsWAK68*, *OsWAK83*, *OsWAK86*, *OsWAK88*).

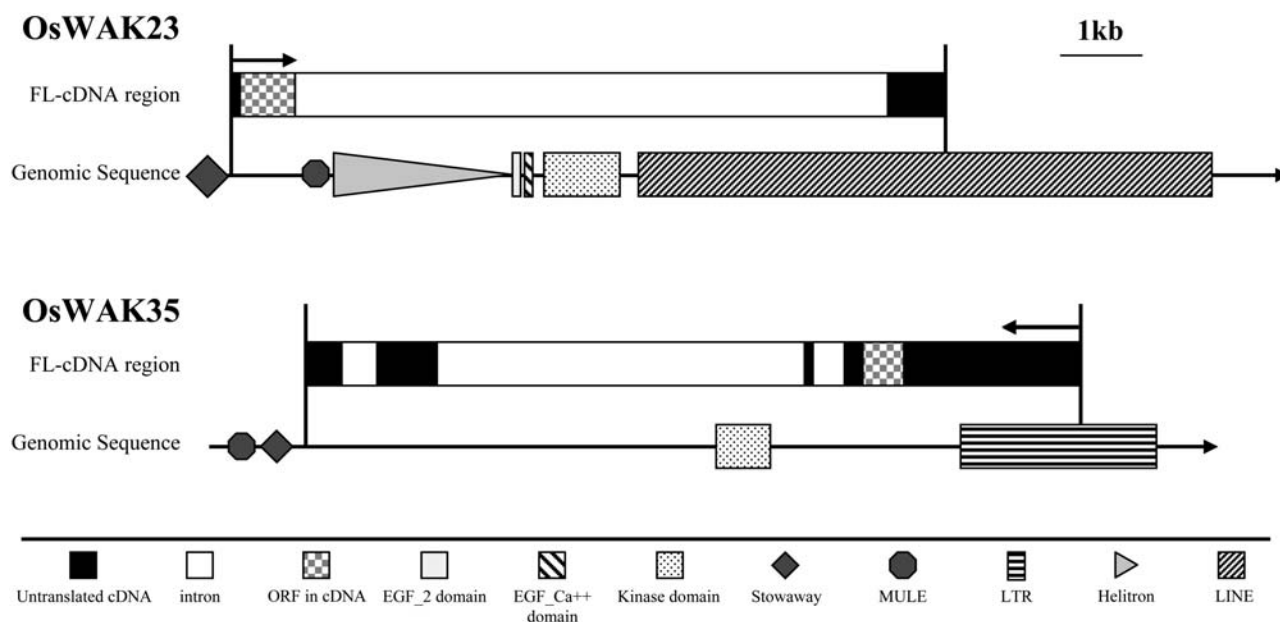
### Intron Number and Position of *OsWAKs*

Arabidopsis RLK-type *WAK/WAKLs* have a two-intron/three-exon gene structure pattern within their coding regions (Verica and He, 2002). Both introns are located within the extracellular regions; the first intron is between the two EGF-like domains and the second is between an EGF-like domain and the kinase domain. Based on predicted coding sequences or FL-cDNA sequences, intron numbers in the coding regions for

each of the 125 *OsWAKs* were determined (Supplemental Table II). For the 38 *OsWAKs* having a FL-cDNA sequence, intron numbers and positions, along with domain positions, are shown in Figure 5. One to three introns were observed in the 5' UTR regions in 14 of the 38 *OsWAKs*, and one intron was found in the 3' UTR regions in three *OsWAKs*. Of the 10 *OsWAK-RLKs* (Fig. 5A), five (*OsWAK11*, *OsWAK25*, *OsWAK46*, *OsWAK50*, *OsWAK92*) have the two-intron/three-exon pattern as observed in Arabidopsis *WAK/WAKLs*. However, three others (*OsWAK32*, *OsWAK87*, *OsWAK91*) have only one intron, likely due to a missing EGF-like domain in their extracellular regions. Another two (*OsWAK28*, *OsWAK29*) have three introns; both acquired an extra EGF-like domain in their extracellular regions. Of the 12 *OsWAK-RLCKs* (Fig. 5B), nine do not contain introns and three have one to two extra introns in their kinase domains. Of the four *OsWAK-RLPs* (Fig. 5C), three have the conserved two-intron pattern in their presumed extracellular regions, and one has an extra intron because of an additional EGF domain. Two of the seven *OsWAK* short genes have one to two introns in their coding regions (Fig. 5D).

**Table II.** *TEs* inserted within or adjacent to five *OsWAK* short genes

Name	Accession No. of FL-cDNA	Type of TE	Insertion Position of TE
<i>OsWAK18</i>	AK103831	Tourist	3' end of FL-cDNA in the element
<i>OsWAK23</i>	AK108554	LINE	3' end of FL-cDNA in the element
<i>OsWAK35c</i>	AK100568	SoloLTR	5' end of FL-cDNA in the element
<i>OsWAK35c</i>	AK100568	Stowaway	38 bp from 3' end of FL-cDNA
<i>OsWAK52</i>	AK111909	MULE	73 bp from 3' end of FL-cDNA
<i>OsWAK101</i>	AK111067	MULE	15 bp from 5' end of FL-cDNA



**Figure 4.** Two examples of *OsWAK* short genes resulting from TE insertions. MULE and Helitron elements inserted within *OsWAK23* and LINE at the 3' end of *OsWAK23* provide a novel splicing site, splicing out the EGF-like domain and kinase domain in the FL-cDNA of *OsWAK23*. The LTR element at the 3' end of *OsWAK35* provides a transcription start site producing an antisense transcript of *OsWAK35*, which has only a short ORF.

Various numbers of introns were observed in the different splicing forms of the five *OsWAKs* that have more than one FL-cDNA (Fig. 5E).

#### Domain Composition and Organization of *OsWAKs*

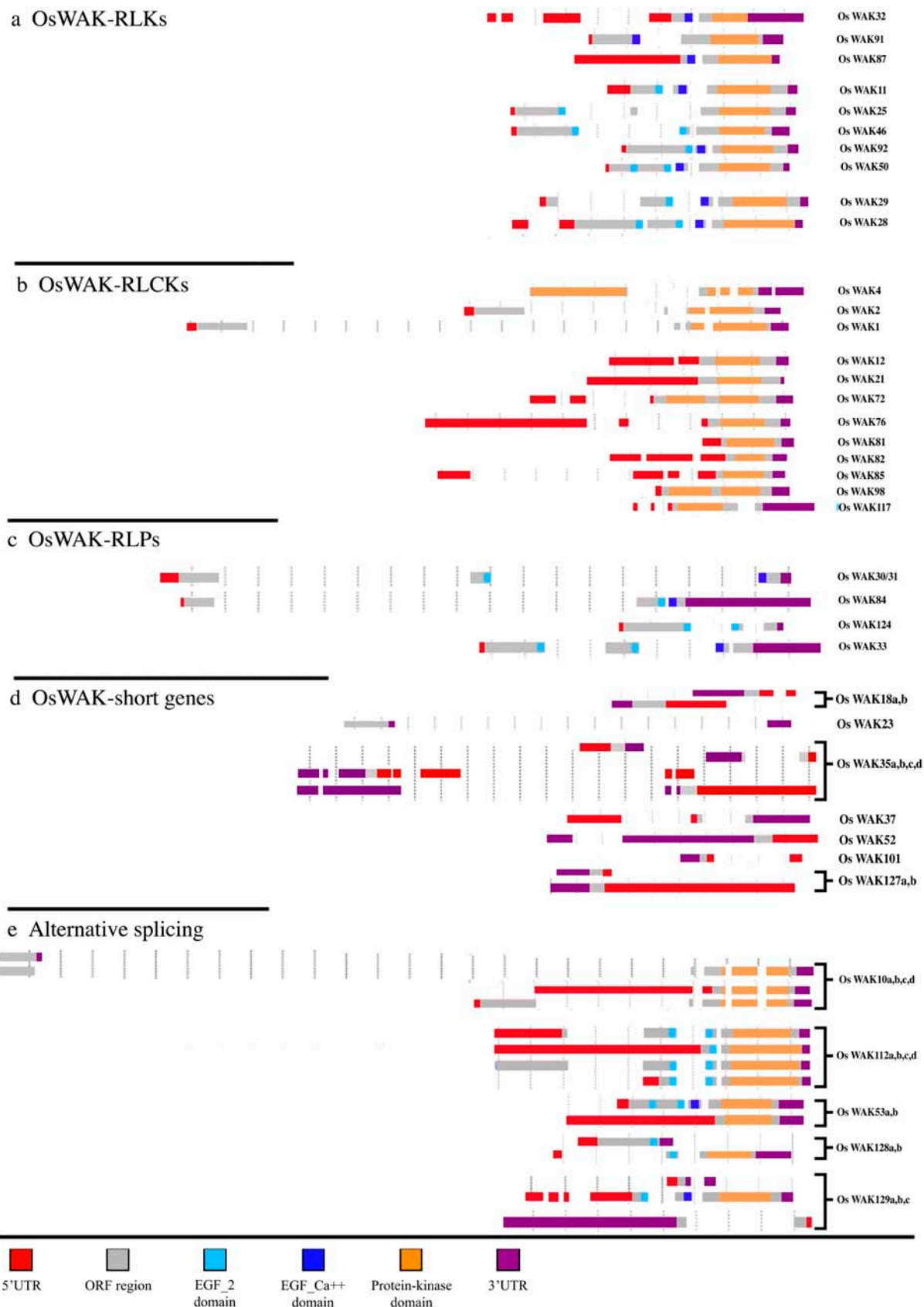
Excluding the five *OsWAK* pseudogenes and the 38 *OsWAKs* with FL-cDNAs (Fig. 5), domain composition and organization were analyzed for the other 82 *OsWAKs*, using their predicted protein sequences to search the Simple Modular Architecture Research Tool (SMART) database (<http://smart.embl-heidelberg.de>) for EGF-like domains, the kinase domain, and the transmembrane domain. Of the 82 *OsWAKs*, 53 *OsWAK*-RLKs contain both extracellular EGF-like domains (EGF-2 and/or EGF-Ca<sup>2+</sup>) and a cytoplasmic protein kinase domain; 15 *OsWAK*-RLCKs contain only one to two kinase domains; nine *OsWAK*-RLPs contain only the extracellular EGF-like domains; and five *OsWAK* short genes contain no domain (Fig. 6).

#### Expression Analysis of *OsWAKs*

Thirty-eight *OsWAKs* have corresponding FL-cDNA sequences (Fig. 5), indicating that they are expressed in rice. Rice EST database searches in Sputnik (<http://mips.gsf.de/proj/sputnik/oryza>) were also performed for each of the 125 *OsWAKs*. Eleven *OsWAKs* that previously had no FL-cDNA matched to one to five different EST sequences (details in Supplemental Table III), indicating they are also expressed. The matched EST sequences were derived from various rice tissues and also from libraries made from plants following

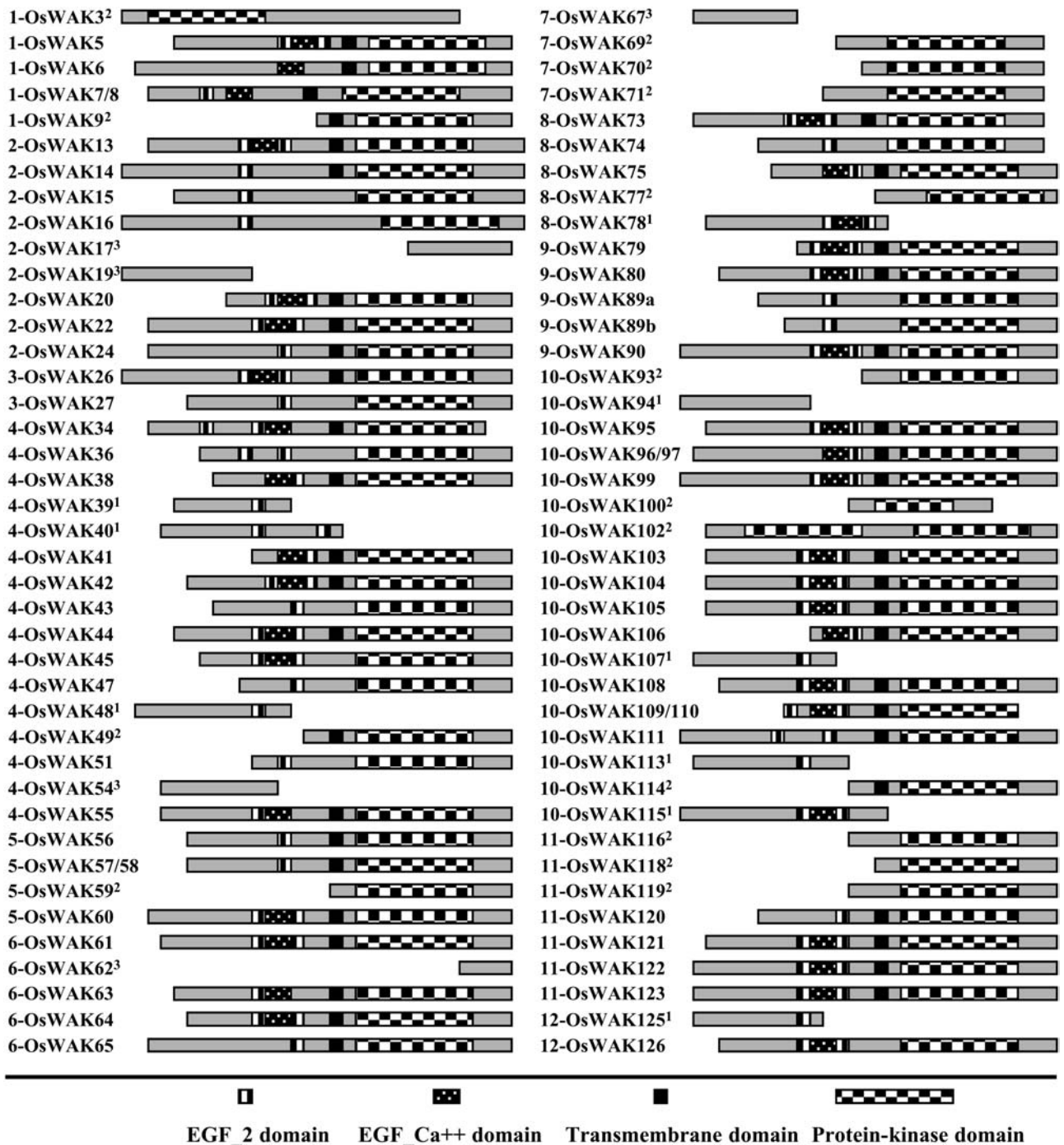
biotic and abiotic stress treatments. Of the 38 *OsWAKs* with FL-cDNAs, four (*OsWAK18*, *OsWAK52*, *OsWAK101*, *OsWAK127*) appear to be expressed as antisense genes because of the antisense orientation of their FL-cDNA sequences (Fig. 5D). Six *OsWAKs* (*OsWAK10*, *OsWAK53*, *OsWAK112*, *OsWAK118*, *OsWAK18*, *OsWAK127*) appear to be alternatively spliced, since they have more than one FL-cDNA sequence; and two (*OsWAK35*, *OsWAK129*) are transcribed in both directions (Fig. 5E).

In addition, five different *OsWAKs* were chosen for expression analysis using reverse transcription (RT)-PCR, *OsWAK7*, *OsWAK50*, *OsWAK125*, plus two short genes, *OsWAK17* (48% amino acid identity to the C terminus of *OsWAK53*) and *OsWAK62* (92% amino acid identity to the extracellular region of *OsWAK63*). DNA samples from leaf tissues were used in PCR analyses; three RNA samples from the root tip, root base, and shoot tissues were used in RT-PCR analyses. *OsWAK50*, which has a FL-cDNA sequence available, had RT-PCR products of an expected size (1,084 bp) from all three tissues; because two introns were present, its PCR product was larger (1,308 bp). This result confirmed *OsWAK50* is an expressed gene and showed there was no DNA contamination in RNA samples used for RT-PCR reactions. The same DNA and RNA samples were used for the other four *OsWAKs*. *OsWAK7* and *OsWAK125* that are without FL-cDNA sequences also had RT-PCR products from one or more rice tissues; because they had no intron, their RT-PCR products (*OsWAK7*, 758 bp; *OsWAK125*, 800 bp) were the same sizes as their PCR products (Fig. 7). The short gene, *OsWAK17*, which had an RT-PCR



**Figure 5.** Gene structure and domain composition of 38 *OsWAKs* having FL-cDNA sequences. Intron number and position are shown along with domains for each *OsWAK*; *OsWAKs* having alternative splicing forms are indicated with brackets; distance between each vertical bar is 500 bp.



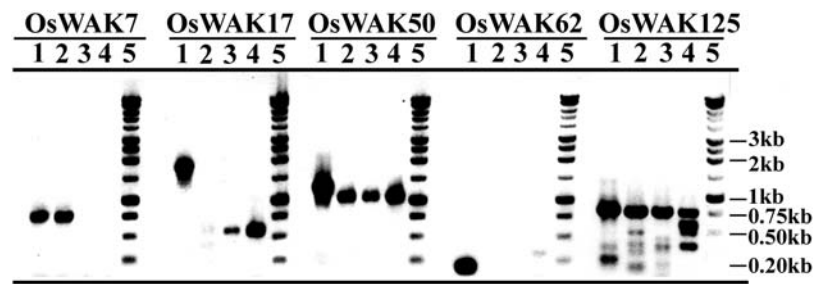


**Figure 6.** Diagrams of domain composition and organization of 82 OsWAKs without FL-cDNA sequences. Number to the left of each OsWAK is chromosome number on which OsWAK is encoded. Superscripts indicate gene type: 1, OsWAK-RLP; 2, OsWAK-RCLK; and 3, OsWAK short genes.

product from the root base and shoot, indicating the short gene, resembling the C-terminal region of an RLK, is also expressed. Because of the presence of two introns, the PCR product of *OsWAK17* was much larger (2,178 bp) than its RT-PCR product (731 bp). Only a PCR product (207 bp) was obtained from *OsWAK62*, indicating it may not be expressed or may be expressed in other tissues.

For functional studies of all OsWAKs, further experimental confirmation of gene expression patterns is needed. To perform this type of analysis, rice microarrays containing all 125 OsWAKs would be ideal; however, this type of custom microarray is not yet available. We therefore took advantage of available rice tiling-path microarrays representing the entire rice chromosome 10 that have been successfully used to

**Figure 7.** RT-PCR gene expression analyses of five *OsWAK* genes in rice. One DNA sample isolated from leaf tissue (lane 1) and three RNA samples isolated from root tip (lane 2), root base (lane 3), and shoot (lane 4) were used for all *OsWAK*s. Lane 5 contains *M<sub>r</sub>* markers for all *OsWAK*s, as indicated.



detect gene transcription activity (Li et al., 2005). Two independent sets of 36-mer probes, with 10-nucleotide intervals, tiled throughout chromosome 10 of *japonica* and *indica*, were designed for the microarrays; rice subsp. *japonica* cv Nipponbare and rice subsp. *indica* cv 93-11, respectively, were used for the experiments. The mRNA samples used for the microarray hybridization were from four normally growing tissues: 7-d-old seedling shoot and root, panicle (heading and filling stages), and suspension culture cells. Determination of the signal-to-noise ratio and expression level of a given gene model was previously described (Li et al., 2005). Based on the results of that analysis, we examined transcription activity for the 20 *OsWAK*s on chromosome 10. Fifteen out of the 20 *OsWAK*s were shown to be expressed in *japonica* tissues, and 10 in both *japonica* and *indica* (Table III). Surprisingly, of the three *OsWAK*s (*OsWAK98*, *OsWAK101*, *OsWAK112*) already having FL-cDNA sequences available, only *OsWAK101* was expressed based on the microarray experiment, suggesting the other two might be expressed under specific stress or developmental conditions.

#### Phylogenetic Analyses of Rice *OsWAK*s, Arabidopsis WAK/WAKLs, and Barley HvWAKs

Intracellular protein kinase domains are typically the conserved regions of RLKs and are used in phylogenetic analyses (Shiu and Bleecker, 2001). Ninety-five *OsWAK*s (67 *OsWAK*-RLKs plus 28 *OsWAK*-RCLKs) containing one or two kinase domains were initially analyzed phylogenetically together with the 21 Arabidopsis WAKs/WAKLs (Fig. 8).

This analysis revealed that most *OsWAK*s and Arabidopsis WAK/WAKLs cluster in species-specific distinct clades, except for four *OsWAK*s (*OsWAK1*, *OsWAK2*, *OsWAK10*, *OsWAK25*) and four WAKL-IV members (*WAK14*, *WAK15*, *WAK20*, *WAK21*) that cluster in the same clade. This result indicates that most *OsWAK*s and Arabidopsis WAK/WAKLs expanded in a species-specific manner; only a few members likely originated from the common ancestral genes that existed before divergence of monocots and dicots.

To further investigate whether expansion of *OsWAK*s is rice specific or due to lineage-specific expansion of this family in monocots, we identified 10 barley WAKs (*HvWAK*s) from the barley EST database (HarVEST; <http://harvest.ucr.edu>). Because only partial predicted protein sequences (of either extracellular regions or

kinase domains) of *HvWAK*s were obtained (Supplemental Data 3), we used predicted, FL protein sequences of 43 *OsWAK*-RLKs containing both extracellular regions and kinase domains in the phylogenetic analyses. Ten *HvWAK*s cluster with individual *OsWAK*s, rather than diverging in a group unique to barley (Fig. 9), suggesting that *OsWAK* expansion in rice likely resulted from lineage-specific expansion of the family in monocots.

#### Localized Duplications Resulted in Expansion of *OsWAK*s

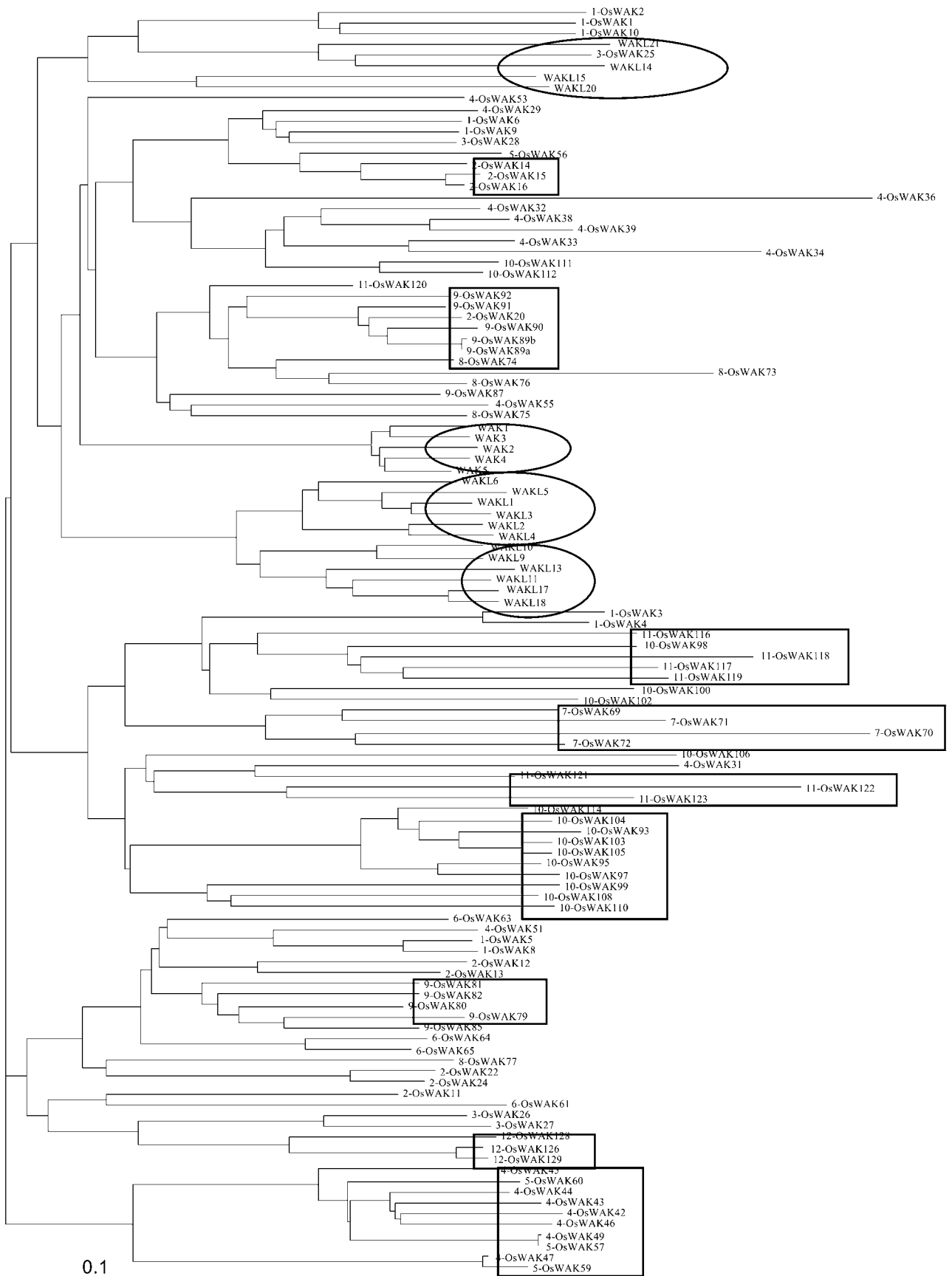
Examination of individual *OsWAK*s in different subclades of the rice-specific clades (Fig. 8) revealed

**Table III.** Tiling-microarray analysis of transcriptional activity in *japonica* and *indica* rice of 20 *OsWAK*s from chromosome 10

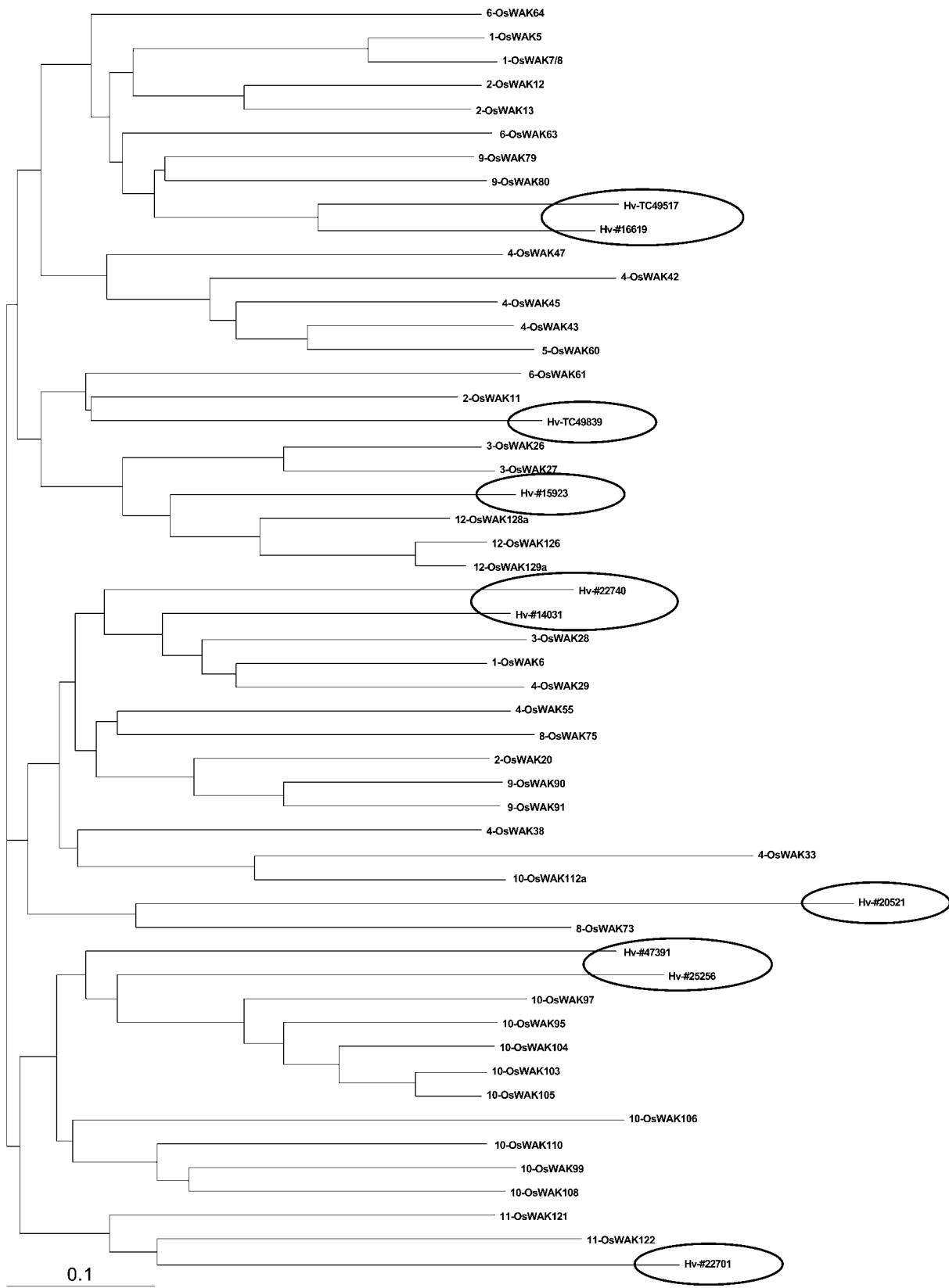
Two independent sets of tiling microarrays were used for *japonica* cv Nipponbare and *indica* cv 93-11, respectively. Analysis was performed using mRNA extracted from four tissues: 7-d-old seedling shoots and roots, panicles (heading and filling stages), and suspension culture cells from *japonica* cv Nipponbare and *indica* cv 93-11.

OsWAK Gene	Strand <sup>a</sup>	Expression	
		<i>japonica</i>	<i>indica</i>
<i>OsWAK93</i>	–	p <sup>b</sup>	P
<i>OsWAK94<sup>c</sup></i>	–	P	P
<i>OsWAK95</i>	–	P	P
<i>OsWAK97</i>	+	P	
<b><i>OsWAK98<sup>d,e</sup></i></b>	–		
<i>OsWAK99</i>	–	P	
<i>OsWAK100<sup>d</sup></i>	+		
<b><i>OsWAK101<sup>e,f</sup></i></b>	+	P	P
<i>OsWAK102<sup>d</sup></i>	+	P	P
<i>OsWAK103</i>	+	P	
<i>OsWAK104</i>	+	P	
<i>OsWAK105</i>	–	P	P
<i>OsWAK106</i>	–	P	P
<i>OsWAK107<sup>c</sup></i>	–	P	
<i>OsWAK108</i>	–	P	P
<i>OsWAK110</i>	+	P	P
<i>OsWAK111</i>	–		
<b><i>OsWAK112<sup>e</sup></i></b>	–		
<i>OsWAK113<sup>c</sup></i>	+	P	P
<i>OsWAK114</i>	+		

<sup>a</sup>Strand, Gene model orientation relative to the sequence; +, from the same orientation; –, from the complement. <sup>b</sup>P, Positively expressed. <sup>c</sup>*OsWAK*-RLP. <sup>d</sup>*OsWAK*-RLCK. <sup>e</sup>Genes in bold have FL-cDNA sequences. <sup>f</sup>Short genes.



**Figure 8.** Phylogenetic analysis using conserved protein kinase domain regions of 95 *Oryza sativa* WAKs and 21 *Arabidopsis thaliana* WAKs/WAKLs. Distinct species-specific clades of *Arabidopsis* WAK/WAKLs are circled. Rectangles highlight several *Oryza sativa* WAK gene groups closely located on the same chromosomes and clustered in the same subclades.



**Figure 9.** Phylogenetic analysis using predicted FL protein sequences of 43 OsWAK-RLK members and 10 partial protein sequences of HvWAKs. Circled HvWAK members aligned with individual OsWAK members.

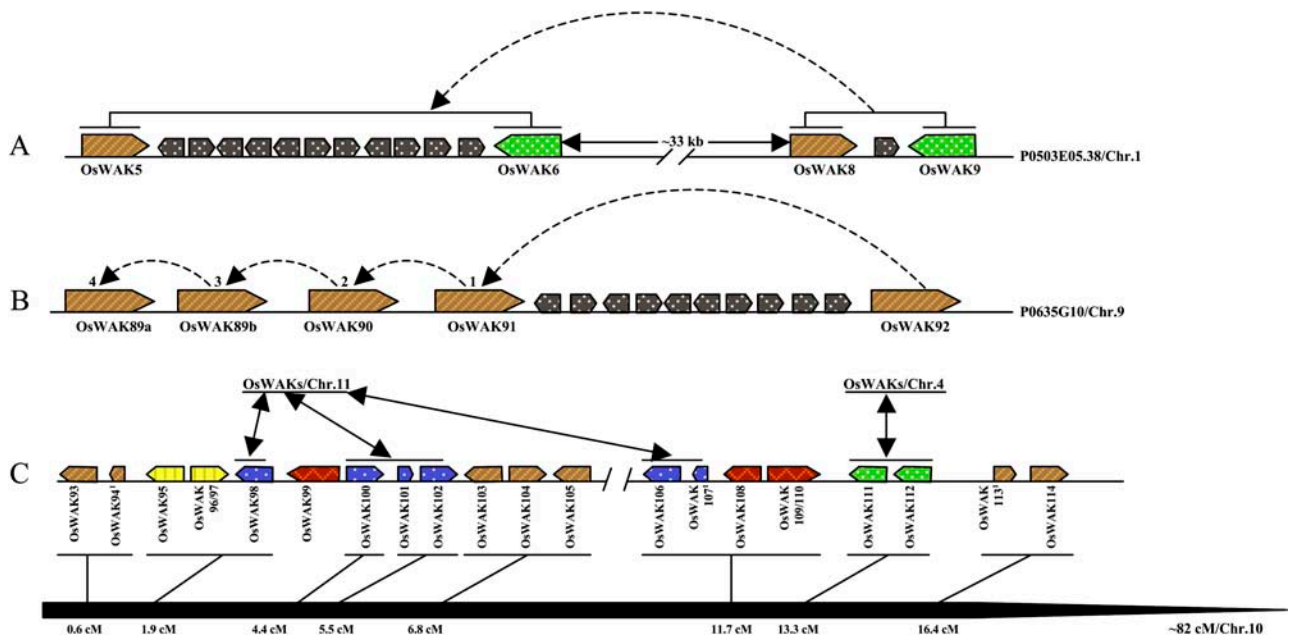
that many small *OsWAK* groups (two to seven genes) located close together on the same chromosomes are clustered in the same subclades, i.e., *OsWAKs* 14 to 16 from chromosome 2; *OsWAKs* 42 to 47 and 49 from chromosome 4; *OsWAKs* 70 to 72 from chromosome 7; *OsWAKs* 79 to 82, 85, and 89 to 92 from chromosome 9; *OsWAKs* 116 to 119 and 121 to 123 from chromosome 11; and *OsWAKs* 126, 128, and 129 from chromosome 12. The high sequence similarity of these physically, closely located *OsWAKs* indicates that likely they arose from localized gene duplications. For example, based on sequence similarity and physical location, the four *OsWAKs* (*OsWAK5*, *OsWAK6*, *OsWAK8*, *OsWAK9*) on chromosome 1 likely derived from duplication of a gene pair and, after duplication, 11 genes inserted between *OsWAK5* and *OsWAK6* (Fig. 10A). The five *OsWAKs* (*OsWAK89a*, *OsWAK89b*, *OsWAK90*, *OsWAK91*, *OsWAK92*) on chromosome 9 likely derived from four gene duplications and, after the duplication, 10 genes inserted between *OsWAK91* and *OsWAK92* (Fig. 10B). The 20 *OsWAKs* on chromosome 10 are physically located as two clusters within 16 cM at the end of the short arm. Twelve likely derived from several localized gene duplications; six were duplicated to or from the *OsWAKs* on chromosome 11, and two were duplicated to or from the *OsWAKs* on chromosome 4 (Fig. 10C).

### Ratios of Nonsynonymous versus Synonymous Substitution Rates between *OsWAK* EGF- $\text{Ca}^{2+}$ Domain Regions

The EGF- $\text{Ca}^{2+}$  domains, the typical extracellular domains in Arabidopsis WAK/WAKLs (Verica and He, 2002), are more conserved than the other extracellular regions and were shown to be under purifying selection (Verica et al., 2003). The EGF- $\text{Ca}^{2+}$  domain regions of *OsWAKs*, which were identified by searching for EGF- $\text{Ca}^{2+}$  domains ([DEQN]-x-[DEQN](2)-C-x(3,14)-C-x(3,7)-C-x-[DN]-x(4)-[FY]-x-C) using the SMART database (<http://smart.embl-heidelberg.de>), are also more conserved than other extracellular regions (data not shown). To determine whether *OsWAK* EGF- $\text{Ca}^{2+}$  domains are also under purifying selection, ratios of nonsynonymous versus synonymous substitution rates among EGF- $\text{Ca}^{2+}$  domain regions from 15 randomly selected *OsWAKs* were analyzed using an improved analytical method (Yang and Nielsen, 2000). Analysis of results shows that most ratios were significantly less than 1 ( $P < 0.05$ ; Supplemental Table IV), indicating *OsWAK* EGF- $\text{Ca}^{2+}$  domain regions are also under purifying selection.

### Physical Locations of *OsWAKs* on Chromosomes

Since all rice clones used for genomic sequencing in the International Rice Genome Sequencing Project can



**Figure 10.** Examples of localized gene duplications resulting in *OsWAK* gene clusters in rice chromosomes. Genes of the same color indicate duplications, and throughout the figure all symbols like those between *OsWAK5* and *OsWAK6* in A indicate inserted genes. A, Pair of *OsWAKs* duplicated on chromosome 1 with 11 genes inserted between *OsWAK5* and *OsWAK6*. BAC clone designation is at the right. B, Four localized duplications, resulting in a five-*OsWAK* gene cluster on chromosome 9; 10 genes were inserted between *OsWAK91* and *OsWAK92*. BAC clone designation is at the right. C, Twenty *OsWAKs* clustered within approximately 16 cM at the end of the short arm of chromosome 10, arose from several different gene duplication events (different colors represent different gene duplications). Six *OsWAKs* (depicted in blue) were duplicated to or from *OsWAKs* on chromosome 11; two *OsWAKs* (in green) duplicated to or from *OsWAKs* on chromosome 4. The remaining 12 *OsWAKs* derived from four gene duplications, which are highly similar to each other, as indicated by their being clustered in the same subclade (Fig. 8). Due to space limitation, genes between *OsWAKs* were not indicated in C.

be physically anchored on the 12 rice chromosomes, as are the pseudomolecules assembled in TIGR (<http://www.tigr.org/tdb/e2k1/osa1/pseudomolecules/info.shtml>), the 125 *OsWAKs* encoded on individual clones can be correspondingly mapped on *japonica* chromosomes (Fig. 11). This analysis showed that *OsWAKs* are clustered on chromosomes in most cases; however, in a few cases only a single *OsWAK* is present at a given chromosomal location, e.g., *OsWAK10* on chromosome 1. The majority of *OsWAKs* (56.8%; 71 of 125) are located on chromosomes 2, 4, 9, and 10.

## DISCUSSION

### Identification of *OsWAK* Gene Family Members

Through a reiterative database search and manual reannotation, we identified 125 *OsWAK* gene family members from rice *japonica* cv Nipponbare. Thirty-seven (approximately 30%) of the identified *OsWAKs* were corrected/reannotated from their earlier automated annotations, such as five merged genes from 10 split genes, five split genes from four fused genes, seven corrected as short genes, and six reannotated from putative pseudogenes. A few of the reannotations were based on sequence similarities to the other *OsWAK* family members; a few other reannotations/corrections were found by sequence extension of the previously annotated genes to recover the missing exons from the 5' or 3' ends.

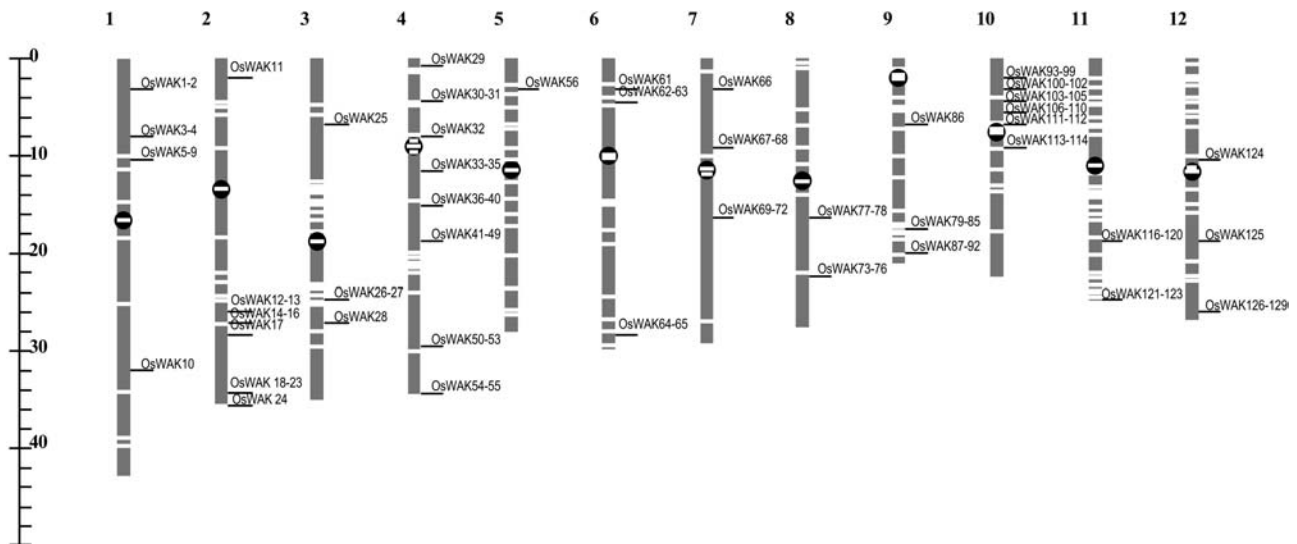
Most reannotations, however, were based on the corresponding FL-cDNA sequences available in rice (Kikuchi et al., 2003). This also occurred in the analysis of the Arabidopsis genome, where approximately 35% of automated annotations were reannotated using FL-

cDNA sequences (Haas et al., 2003). Misannotations from the current automated annotation programs could occur for several reasons, including insufficient numbers of experimentally confirmed genes being available to train the gene-prediction programs, unexpectedly large introns, presence of TE insertions that interrupt normal gene sequences and transcription, and possible sequencing errors (Haas et al., 2003; Wang et al., 2003; Castelli et al., 2004). In addition, few of the annotation errors in rice were due to incomplete sequencing information; of the 63 BACs that encode the 125 *OsWAKs*, six BACs were not yet completed at the time the annotation was generated (see Supplemental Table II, column D). None of the five *OsWAK* pseudogenes came from incomplete BACs; however, the merged *OsWAK57/58* was from an incomplete BAC (OJ1480\_H01). Also, two *OsWAKs* (*OsWAK10*, *OsWAK50*) were truncated because they were located at the end of the BAC sequences; the full length of these two *OsWAKs* was recovered from their FL-cDNA sequences. Therefore, further experimental validations of the reannotated *OsWAKs* are warranted.

### Classification of the *OsWAK* Gene Family

Of the 125 *OsWAK* gene family members, 67 are *OsWAK-RLKs* containing both extracellular EGF-like domains and an intracellular kinase domain; 28 are *OsWAK-RLCKs* containing only the kinase domain; 13 are *OsWAK-RLPs* with only the extracellular EGF-like domains; 12 are *OsWAK* short genes; and five are pseudogenes. Functions of these various *OsWAKs* are yet to be determined.

The unique character of *OsWAK-RLPs* and *OsWAK-RLKs* is the EGF-like domain at the N-terminal



**Figure 11.** Physical locations of 125 *OsWAKs* on the 12 *japonica* rice chromosomes, based on TIGR rice pseudomolecules (Osa1, release 2). Horizontal white lines within each chromosome indicate sequence gaps remaining; horizontal black lines indicate locations of *OsWAKs* on each chromosome. Three *OsWAKs* (*OsWAK57/58*, *OsWAK59*, *OsWAK60*) remain to be located on the map because, to date, the contig (OJ1480\_H01) encoding these three *OsWAKs* has not been mapped to the chromosome 5 pseudomolecule in TIGR.

extracellular region. The function of EGF-like domains in OsWAKs and Arabidopsis WAK/WAKLs is yet to be determined. Analysis of other EGF-containing proteins from plants and animals suggests several possible roles for the EGF-like domains in the OsWAK-RLKs and OsWAK-RLPs. In the plant vacuolar sorting protein, BP-80, EGF domains were shown to alter the structural conformation of the ligand-binding domains, thereby increasing their affinity for ligand binding (Cao et al., 2000). EGF-like domains in certain proteins from animals directly participate in protein-protein interactions (Appella et al., 1987; Rebay et al., 1991; Kuroda and Tanizawa, 1999; Stenberg et al., 1999). For example, in the fruit fly (*Drosophila melanogaster*), the transmembrane protein, Notch, which contains 36 tandem EGF-like repeats, interacts with other EGF-containing transmembrane proteins, including Delta and Serrate. Two of the Notch EGF repeats were shown to be both necessary and sufficient for mediating these interactions (Rebay et al., 1991). The formation of both homodimeric and heterodimeric receptor complexes has been proposed for a large number of receptor kinases (Heldin, 1995), including members of the EGF family of receptor kinases in animals (Yarden and Schlessinger, 1987; Sliwkowski et al., 1994) and the CLV1 receptor complex in plants (Clark et al., 1997; Trotochaud et al., 1999). It remains to be seen whether the EGF-like domains in OsWAK-RLKs and OsWAK-RLPs are involved in protein-protein interactions.

In plants, several different RLK members were characterized and found to function in a diverse array of signaling processes, including phytohormone responses (Chang et al., 1993 [ETR1]; Li and Chory, 1997 [BRI1]; Clark et al., 1998 [CTR1]), reproduction (Stein et al., 1991 [SRK]; Mu et al., 1994 [PRK1]), developmental regulation (Becraft et al., 1996 [CRINKLY4]; Clark et al., 1997 [CLV1]; Yokoyama et al., 1998 [ERECTA]; Jinn et al., 2000 [HAESA]), and plant disease resistance responses (Song et al., 1995 [Xa21]). These RLKs are divided into different subclasses based on sequence relationships between their predicted extracellular regions (Shiu and Bleecker, 2001).

Several RLP members in plants were also shown to have important functions. For example, CLV2 functions in shoot meristem development in Arabidopsis (Kayes and Clark, 1998; Jeong et al., 1999); Fasciated Ear 2, an ortholog of CLV2, in the regulation of ear inflorescence meristem proliferation in maize (Taguchi-Shiobara et al., 2001); Too Many Mouths (TMM) in stomatal patterning in Arabidopsis (Nadeau and Sack, 2002); Cf-9 in disease resistance in tomato (Jones et al., 1994); and Xa21D in pathogen resistance in rice (Wang et al., 1998). Arabidopsis WAKL7 is also an RLP gene that is wound inducible (Verica et al., 2003). A few plant RLCKs were found to be functional in disease resistance signaling processes, including Pto in pathogen resistance (Martin et al., 1993) and EDR1 in negative regulation of defense responses (Frye et al., 2001).

Especially interesting are recent studies showing that plant RLK, RLP, and RLCK members could function together in the same signaling pathways. For example, in the self-incompatibility signaling pathway in Brassica, the S-receptor kinase (an RLK member), the S-locus glycoprotein (an RLP member resembling the extracellular part of the S-receptor kinase), and the M-locus protein kinase (an RLCK member) function as a single complex in signaling (Cui et al., 2000; Takasaki et al., 2000; Murase et al., 2004). Another example is the complex interaction of TMM (Leu-rich repeat [LRR]-RLP) and three ERECTA family members (LRR-RLKs) in stomatal patterning and differentiation (Shpak et al., 2005); TMM negatively regulates specific ERECTA family members at critical steps in stomatal differentiation. The mechanism for regulation of the ERECTA family members by TMM is still unknown. It was suggested that TMM forms a receptor heterodimer with ERECTA family RLKs, preventing signaling; the lack of a signal transducer domain in TMM supports the idea of this inhibitory function. Alternatively, TMM may have the same ligands as the ERECTA family, hence representing the ERECTA family signaling pathway. These examples suggest that different OsWAKs could also function by heterodimerization in multimeric assemblies or alone in transduction cascades.

Twelve OsWAK family members are short genes with less than 300 amino acids and no known domain. This type of short gene was also identified in the Arabidopsis FL-cDNA sequence analyses (Haas et al., 2003). In some cases, the short genes appear to be functional in Arabidopsis, e.g. the CLE gene family members (Hobe et al., 2003; Sharma et al., 2003). A possible clue into the genesis of OsWAK short genes comes from the observation that recognizable TEs are found within or near the 5' or 3' end of five of the short genes (*OsWAK18*, *OsWAK23*, *OsWAK35*, *OsWAK52*, *OsWAK101*). They either provide a novel splicing site or initialize antisense transcription, which resulted in the short ORFs. Further studies of the function, if any, of the OsWAK short genes, especially of the seven OsWAK short genes with corresponding FL-cDNA sequences, should provide interesting insights into the role of OsWAK short genes in rice.

### Structure and Expression of OsWAKs

The two-intron gene structure pattern of Arabidopsis WAK/WAKLs (Verica and He, 2002) was generally conserved in OsWAKs. However, in OsWAKs with an extra/missing EGF-like domain, an extra/missing intron was observed in their extracellular regions. Extra introns were also observed in the kinase domains of a few OsWAK-RLCKs. Intron variation in OsWAKs demonstrates that OsWAK gene structure is not as conserved as it is in the Arabidopsis WAK/WAKLs and is a reflection of its expansion in rice.

Antisense expression, bidirectional transcription, and alternative splicing were observed in a few of

the 38 *OsWAKs* with FL-cDNA sequences. This type of antisense and bidirectional transcription was also observed from FL-cDNA sequence analysis of many other rice genes (Osato et al., 2003). However, biological implications of the different directions of transcription and the nature of the resulting products remain to be determined. Expression analyses based on the tiling-microarray experiment of the 20 *OsWAKs* on chromosome 10 and RT-PCR for five additional *OsWAKs* suggest that the majority of *OsWAKs* are expressed in rice.

### Evolutionary Expansion of *OsWAKs*

Compared to the 26 *Arabidopsis WAKs/WAKLs* (Verica and He, 2002), the *japonica* rice genome has nearly a 5 times greater number of *OsWAKs*. Previous comparative analyses between the two plant species of other gene families, e.g. the *CONSTANS*-like gene family (Griffiths et al., 2003), the *Dof* family (Lijavetzky et al., 2003), and the *LRR* extensin family (Baumberger et al., 2003), did not reveal the magnitude of difference in family sizes that is seen with the *OsWAKs*. Therefore, the evolutionary expansion of the *OsWAK* gene family does not appear to be due simply to the larger genome size of rice.

Phylogenetic analyses of *OsWAKs* and *Arabidopsis WAK/WAKLs* show that most *OsWAKs* and *Arabidopsis WAK/WAKLs* are clustered in distinct species-specific clades, suggesting species-specific expansion in both plants. Further phylogenetic analyses, comparing *OsWAKs* with barley *HvWAKs*, indicate that *OsWAK* expansion was mainly due to its lineage-specific expansion in monocot species. This type of divergence between monocot (rice) and dicot (*Arabidopsis*) species was also observed for a large gene family involved in pathogen resistance, the nucleotide-binding site (NBS)-*LRR* gene family (Bai et al., 2002). In rice, greater than 600 members of the *OsNBS-LRR* gene family were identified, 3 to 4 times greater than the number in *Arabidopsis*. Also, most *OsNBS-LRRs* do not encode *Toll* and mammalian interleukin-1 receptor domains; however, most *Arabidopsis NBS-LRRs* do contain an interleukin-1 receptor domain. This lineage-specific divergence of domains and expansion of certain gene families involved in pathogen resistance in both monocot and dicot species likely occurs to enable reaction of plants to pathogens unique to each species.

It has been suggested that the expansion of *Arabidopsis WAK/WAKLs* is due to both local tandem duplications and large-scale genomic duplications (Verica and He, 2002). However, it appears from our analyses in rice that expansion of *OsWAKs* is probably due in large part to localized gene duplications, since many small *OsWAK* groups, located in close proximity on the same chromosome, have high sequence similarity. Localized duplication was also suggested as the main reason for *WAK* gene family expansion in *indica* rice (Shiu et al., 2004). A recent study that looked at the

history of rice genome duplications showed that massive individual gene duplications are ongoing in the rice genome, providing a robust source of raw material for the genesis of new genes and gene functions (Yu et al., 2005). The results of our study provide the necessary genomic information for further in-depth study of the functions of *OsWAKs* and of their evolutionary expansion in the rice genome.

## MATERIALS AND METHODS

### Sequence Retrieval and Analysis

The DNA and predicted protein sequences, annotated as putative *OsWAKs* from the genome sequence of rice (*Oryza sativa japonica* cv Nipponbare, were retrieved from three public databases: (1) TIGR Rice Genome Annotation Database (Osa1; <http://www.tigr.org/tdb/e2k1/osa1>; Yuan et al., 2005); (2) Rice Protein Database in Gramene (<http://www.gramene.org>; Ware et al., 2002); and (3) GenBank in the National Center for Biotechnology Information (<http://www.ncbi.nlm.nih.gov>). The same sequences from the same or overlapping contigs, but annotated by different putative *OsWAK* names in the various databases, were checked using the multiple sequence alignment tool of ClustalW from the European Bioinformatics Institute (<http://www.ebi.ac.uk/clustalw/index.html>). Genes from rice varieties other than *japonica* cv Nipponbare were manually removed. BLAST search tools, BLASTp and tBLASTn (Altschul et al., 1997), were used to identify additional *OsWAK* sequences from Osa1 and GenBank, using 10 putative *OsWAK* protein sequences as queries.

### Domain Searches of *OsWAK* Protein Sequences

The SMART database (<http://smart.embl-heidelberg.de>) was used to search for EGF-like domains, protein kinase domains, and transmembrane domains. BioEdit software (<http://www.mbio.ncsu.edu/BioEdit/bioedit.html>) was utilized as a tool for diagramming domain positions for *OsWAKs*.

### RT-PCR Analysis of *OsWAKs*

Total DNA and RNA, used in PCR and RT-PCR respectively, were isolated from 2-week-old seedlings of rice *japonica* cv Nipponbare. DNA extraction was performed on young leaf tissues (Cone, 1989); shoot and root tissues were used for total RNA isolation with TRIzol reagent (Gibco-BRL, Life Technologies), according to the manufacturer's instructions. RNA purification, PCR, and RT-PCR were conducted as described (Meng et al., 2003). PCR or RT-PCR products were purified using QIAquick PCR purification kit (Qiagen) and sequenced (Elim Biopharmaceuticals). Gene-specific RT-PCR primers were designed for each *OsWAK* as follows: *OsWAK7* (F-AGTGTGCACTAGTC-ATGCTGCA, R-CTATGGCATGCATATGAAGTCATG); *OsWAK17* (F-GTTG-ATTGGTTCCTCTTGATGCAG, R-GAAGAGTGGAGAGTGGAGGATGA); *OsWAK50* (F-CAACTCAAGCTTAACGTCAACTC, R-GAGTCACTGGTGG-TGAATATC); *OsWAK62* (F-ACTCATGGACATTATAGGTCATC, R-GAATGTG-CACCATCTCTCC); and *OsWAK125* (F-AACCTCACCTGCAGCAGC-AAC, R-TCATGGCATCCACCAGCAACG).

### Multiple Sequence Alignment and Phylogenetic Tree Analysis

Predicted protein sequences of *OsWAKs* were retrieved from public databases as described in "Results." *Arabidopsis (Arabidopsis thaliana)* *WAK/WAKL* protein sequences were obtained from GenBank as described in Verica and He (2002). Barley (*Hordeum vulgare*) *HvWAKs* were retrieved from the barley EST database, HarVEST (<http://harvest.ucr.edu>), and their predicted protein sequences were translated by frame matching to known *WAK* protein sequences. Multiple sequence alignment analysis was performed using ClustalW, with default parameters set as in the European Bioinformatics Institute (<http://www.ebi.ac.uk/clustalw/index.html>) and BioEdit. The phylogenetic tree was produced using TreeView software (<http://taxonomy.zoology.gla.ac.uk/rod/treeview.html>).



## Physical Mapping of OsWAKs on Rice Chromosomes

All sequenced contigs from *japonica* cv Nipponbare were physically constructed as pseudomolecules (release 2) in TIGR (<http://www.tigr.org/tdb/e2k1/osa1/pseudomolecules/info.shtml>), representing the 12 rice chromosomes. OsWAKs were then mapped on individual chromosomes, based on corresponding contigs on rice chromosomes.

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