

RESEARCH ARTICLE

Exploring *OshAK* Transporters for the Salt Responsiveness in Rice

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ABSTRACT

Environmental stresses are the major constraint in crop growth and productivity, particularly in rice. Improving rice against major abiotic stressors such as salinity is one of the major thrust areas. In this direction, the trait of sodium exclusion was the primary focus among the rice breeders to enhance salt tolerance. Along similar lines, increased uptake of potassium, the counter ion of Na⁺, through high-affinity potassium transporters (HAK) assumes significance in the context of maintaining a favorable Na/K ratio, a low Na/K is an indicator of salt tolerance. The *OshAK* family in rice includes 27 members and an insight into the evolution, structure, and expression through *in silico* approaches was attempted. Phylogenetic analysis exhibited four distinct groupings of the *OshAK* family, and functional motif analysis revealed a characteristic consensus sequence among the *OshAK* transporters. Relative gene expression analysis based on published microarray data put forth the differential regulation of *OshAK3 X2*, *OshAK8*, *OshAK14(X2)*, *OshAK15* and *OshAK26* between salt-tolerant landrace *Pokkali* and susceptible cultivar IR29 and their tissue-specific expression profiles predicted. This study narrowed down a set of putative *OshAKs* from the larger family for their role in enhancing cellular potassium under salt stress thus paving the way for crop improvement.

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INTRODUCTION

Rice, the major calorie crop for half of the world's population was classified as salt-sensitive (Sen *et al.*, 2020). Salinity is one of the major abiotic stress factors, severely affecting growth and productivity, particularly rice is highly susceptible to salt stress (Amaravel *et al.*, 2019; Akilan *et al.*, 2023). Salt stress was attributed to an excess of dissolved salts, primarily the chlorides and sulfates of sodium, magnesium, and calcium, in soil and irrigation water (Bernstein,1975; Wali *et al.*, 2021). Accordingly, the saline soil was characteristic of an electrical conductivity (EC) of above 4 dS/m at 25 °C, exchangeable sodium percent (ESP) of less than 15, and pH of below 8.5 (Allison and Richards, 1954). Besides, NO₃⁻ was also reported to contribute to soil salinity (Stavi *et al.*, 2021). Salt stress impacts crop growth via stuntedness, smaller leaves, reduced vegetative vigor, and ultimately poor yield (FAO, 2020).

Physiologically, excess sodium ions lead to damage to chloroplast, and cell membranes and affect various vital cellular processes (Hameed *et al.*, 2021). Towards improving rice crop in particular against salt stress, breeders have been continuously engaged in the introgression of several salt tolerant quantitative trait loci (QTLs) of which SALTOL harboring *OshKT1;5* transporter remains the mainstay (Ren *et al.*, 2005; Thomson *et al.*, 2010), this in turn had resulted in a fair success in improving rice crop for salt tolerance.

In contrast, enhancing cellular K⁺ content could be another promising and complementing approach toward the maintenance of Na/K homeostasis (Shabala and Cuin, 2008; Shabala and Pottosin, 2010). Potassium is the most abundant inorganic cation in

plants and has turned out to be an inevitable ion in plant growth and development (Talbot and Zeiger, 1996; Pandey and Mahiwal, 2020). Thus, potassium has evolved to be a functional component of several enzymes involved in different metabolic processes such as photosynthesis, carbon, nitrogen metabolism, protein synthesis, osmoregulation, transport of sugars, disease resistance, etc. (Hasanuzzaman *et al.*, 2018). The K⁺ uptake and its homeostasis were in turn regulated by several membrane-bound transporters and channels that facilitate the K⁺ uptake from the external environment and its transport within the plant (Maathuis and Sanders, 1994; Maathuis *et al.*, 1997). These transporters and channels were further classified into four multigene families (i.e.) high-affinity K/Na transporter (*HKT*), K⁺ uptake permease (*KT/KUP/HAK*), K⁺ exchange antiporters (*KEA*) and cation/H⁺ exchangers (*CHX* transporters) (Zhang *et al.*, 2012). Among these, the *KT/KUP/HAK* family is the largest of the K⁺ transporter family comprising 13 members in *Arabidopsis*, 24 in *Poplar*, and 27 members in rice. Baneulos *et al.* (2002) have identified 14 gene encodes for HAK transporters in rice viz., *OshAK2*, *OshAK3*, *OshAK5-15*, and *OshAK17*. Later, Gupta *et al.* (2008) identified twenty-six *OshAK* transporters and Amrutha *et al.* (2007) reported 27 *OshAK* transporters in rice. Further, the *OshAK* transporter family was known to enhance the potassium content even under K⁺ limiting conditions by enhancing their expression levels thereby improving the salinity tolerance in rice (Okada *et al.*, 2008). In this study, we have attempted to characterize the members of the *OshAK* family for their evolutionary relatedness, analysis of the functional motifs, and their differential expression status between salt-tolerant and susceptible rice genotypes.

MATERIALS AND METHODS

Construction of phylogenetic tree

The nucleotide sequences for all 27 *OshAK* transporters in rice were retrieved from the National Center for Biotechnology Information (NCBI) (<https://www.ncbi.nlm.nih.gov/>) and downloaded in FASTA format. Multiple sequence alignment was done using Clustal W2 (www.ebi.ac.uk/Tools/) and the phylogenetic tree was constructed using the MEGA-X software (<https://www.megasoftware.net/>).

Identification of functional motifs

The amino acid sequences for all 27 *OshAKs* were retrieved from NCBI and downloaded in FASTA format. Multiple Em for Motif Elicitation (MEME) suite was

used to identify the functional motifs (<https://meme-suite.org/meme/>).

In silico differential expression analysis of OshAK transporters

The publically available microarray dataset involving the roots of salt-tolerant rice landrace *Pokkali* and salt-susceptible cultivar IR29 performed as a seedling stage salt stress experiment by Cotsaftis *et al.* (2011) with the following identifier GSE14403 was downloaded from the NCBI GEO Omnibus database (<https://www.ncbi.nlm.nih.gov/geo/>). The expression values for the 27 *OshAKs* were extracted from the series matrix file both from the salt-tolerant landrace *Pokkali* and salt-susceptible cv. IR29. The CREP database facilitated the querying process (CREP-Query (ncpgr. cn).

Tissue-specific expression analysis of selected OshAKs

Selected differentially expressed *OshAK* genes were further analyzed to explore their tissue-specific expression profiles using the RNASeq dataset available in ExPath2.0 (expath.itps.ncku.edu.tw/expression/rice/search.php). Transcript abundance of selected *OshAKs* in the roots and shoots of one-week-old seedlings under 150 mM salt stress were analyzed for their tissue-specific expression.

RESULTS AND DISCUSSION

Towards understanding the evolutionary relatedness amongst the 27 *OshAK* transporters identified in rice, a phylogenetic tree was constructed based on the gene sequences using Mega X software. The 27 *OshAKs* were distinctly grouped into four clades viz., clade I, clade II, clade III, and clade IV wherein, the majority of the *OshAKs* were found clustered under clade IV. This is as per the earlier report by Gupta *et al.* (2008), wherein, the twenty-seven transporters were subgrouped into four clades. The clade I consisted of three members viz., *OshAK14*, *OshAK15* and *OshAK23*. The clade II included seven transporters viz., *OshAK5*, *OshAK18*, *OshAK12* and *OshAK11*. Similarly, clade III comprised seven members which were further classified into two clusters, cluster I consisted of *OshAK2* and *OshAK7* and cluster II included *OshAK8*, *OshAK9*, *OshAK10*, *OshAK24*, and *OshAK25*. The clade IV consisted of the remaining 13 *OshAKs* which were further classified into two clusters. The cluster I of clade IV consisted of only one member *OshAK13* and the cluster II included rest of them viz., *OshAK26*,

OsHAK6, *OsHAK3*, *OsHAK17*, *OsHAK4*, *OsHAK20*, *OsHAK19*, *OsHAK1*, *OsHAK22*, *OsHAK16*, *OsHAK21* and *OsHAK27* (Fig. 1). The chromosomal locations of the 27 *OsHAKs* were found distributed across eight of the 12 rice chromosomes viz., chr 1, chr 2, chr 3, chr 4, chr 6, chr 7, chr 8 and chr 9. No *OsHAK* genes were detected on chromosomes 5, 10, 11, and 12 (Yang *et al.*, 2009; Gupta *et al.*, 2008). The average length of the coding sequence (CDS) and polypeptide were 3070 bp and 770 amino acids, respectively (Table 1).

Further, the functional motifs present in the 27 *OsHAK* transporters were predicted using the MEME suite (Fig. 2). A total of 14 different functional motifs were predicted across the transporters.

The characteristic feature of these transporters was the presence of a consensus motif viz., 'GVVYGD LGTSPLY' (Rodriguez-Navarro, 2000). In our study, the 10th motif was found to harbor the above consensus amino acid sequence which appeared to be the signature of the *OsHAK* transporter family (Fig. 3). The consensus amino acid sequence across the 14 identified motifs showed a higher frequency of alanine and glycine.

Further, towards exploring the role of putative *OsHAK* transporter(s) involved in enhancing the K⁺ uptake under salt stress conditions in rice, *in silico*

differential expression analysis of the 27 *OsHAK* genes was performed based on the expression values extracted from the publically available microarray data set, NCBI (GSE14403; Cotsaftis *et al.*, 2011). Landrace *Pokkali* is a well-known rice genotype and was extensively subjected to molecular investigations towards uncovering the mechanisms underlying salt tolerance. On the other hand, the rice cultivar IR29 was known for its salt susceptibility. Differential expression analysis revealed the putative role of a few *OsHAKs* which were found to be differentially regulated under salt stress between the salt-tolerant landrace *Pokkali* and salt susceptible cv. IR29 at the seedling stage. As stated above, *OsHAK* transporters viz., *OsHAK3* splice variant (X2), *OsHAK8*, *OsHAK14* splice variant (X2), *OsHAK15* and *OsHAK26* were upregulated upon salt stress in *Pokkali*, and these might be considered to have a role during salt stress. Besides, *OsHAK6* and *OsHAK13* were found to show no variations in their expression levels under salt stress for *Pokkali* and cv. IR29. On the contrary, a few *OsHAK* transporters viz., *OsHAK1*, *OsHAK2*, *OsHAK10*, and *OsHAK18* showed a declining expression in both *Pokkali* and cv. IR29. On expected lines, a majority of the *OsHAK* transporters showed downregulation in salt susceptible rice cv. IR29 (Fig. 4).

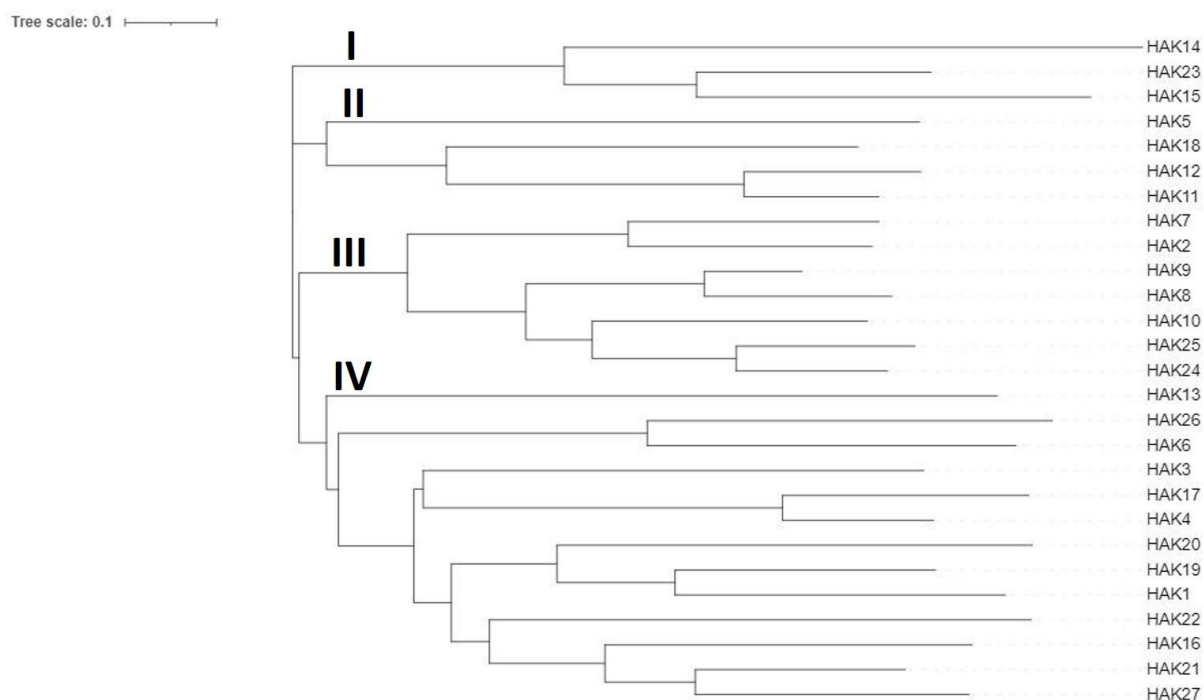


Fig. 1. Phylogenetic tree showing the evolutionary relationship among 27 *OsHAK* transporters in rice.

I, II, III, and IV denote clades of the tree

Table 1: List of OsHAK transporters in rice with gene name, locus ID, chromosome, length of coding sequence (CDS), length of the polypeptide

Sl. No	Gene	Gene Id.	Chromosome	CDS length (bp)	Length of polypeptide (aa)
1	Os HAK1 (X1)	LOC_Os04g32920	4	3092	792
2	Os HAK1 (X2)	LOC_Os04g32920	4	3174	801
3	OsHAK2	LOC_Os01g70940	1	3678	640
4	OsHAK3 (X1)	LOC_Os01g27170	1	3152	808
5	OsHAK3 (X2)	LOC_Os01g27170	1	3041	791
6	OsHAK4	LOC_Os08g36340	8	2584	697
7	OsHAK5	LOC_Os01g70490	1	2757	770
8	OsHAK6	LOC_Os01g70660	1	2411	748
9	OsHAK7	LOC_Os07g47350	7	2803	811
10	OsHAK8	LOC_Os03g21890	3	2577	793
11	OsHAK9	LOC_Os07g48130	7	3282	635
12	OsHAK10	LOC_Os06g42030	6	5864	843
13	OsHAK11	LOC_Os04g52390	4	3493	791
14	OsHAK12	LOC_Os08g10550	8	3404	582
15	OsHAK13	LOC_Os06g45940	6	2954	778
16	OsHAK14 (X1)	LOC_Os07g32530	7	3121	842
17	OsHAK14 (X2)	LOC_Os07g32530	7	3246	859
18	OsHAK15	LOC_Os04g52120	4	3138	867
19	OsHAK16	LOC_Os03g37840	3	2909	811
20	OsHAK17 (X1)	LOC_Os09g27580	9	2894	707
21	OsHAK17 (X2)	LOC_Os09g27580	9	2352	649
22	OsHAK18	LOC_Os09g38960	9	3408	793
23	OsHAK19	LOC_Os02g31910	2	2874	742
24	OsHAK20	LOC_Os02g31940	2	2795	747
25	OsHAK21	LOC_Os03g37930	3	2625	799
26	OsHAK22	LOC_Os07g01214	7	3255	808
27	OsHAK23	LOC_Os09g21000	9	3170	877
28	OsHAK24	LOC_Os06g15910	6	2319	772
29	OsHAK25	LOC_Os02g49760	2	3204	771
30	OsHAK26	LOC_Os08g39950	8	2739	739
31	OsHAK27	LOC_Os03g37830	3	2832	811

Note: X1 and X2 are the splice variants of the respective genes

In this investigation, the *OsHAK3* splice variant (X2) was one of the *OsHAK* family found to be upregulated in the salt-tolerant rice *Pokkali*. In support of this observation, the work of Zhang *et al.* (2020) aimed at functionally characterizing *OsHAK3* reported that the loss of function of *OsHAK3* resulted in a reduction in cellular K⁺ uptake leading to stunted plant growth, especially under K⁺ limiting conditions. Subsequently, in an attempt to identify the key players involved in

salinity tolerance during the seedling germination, Ju *et al.* (2022) explored the importance of *OsHAK3* along with *OsHAK5* as a promising candidate gene for improving salt stress tolerance in rice. *OsHAK3* expression was found to be primarily associated with roots and was found to play a crucial role in maintaining the cellular K homeostasis in turn leading to a favorable Na/K ratio.

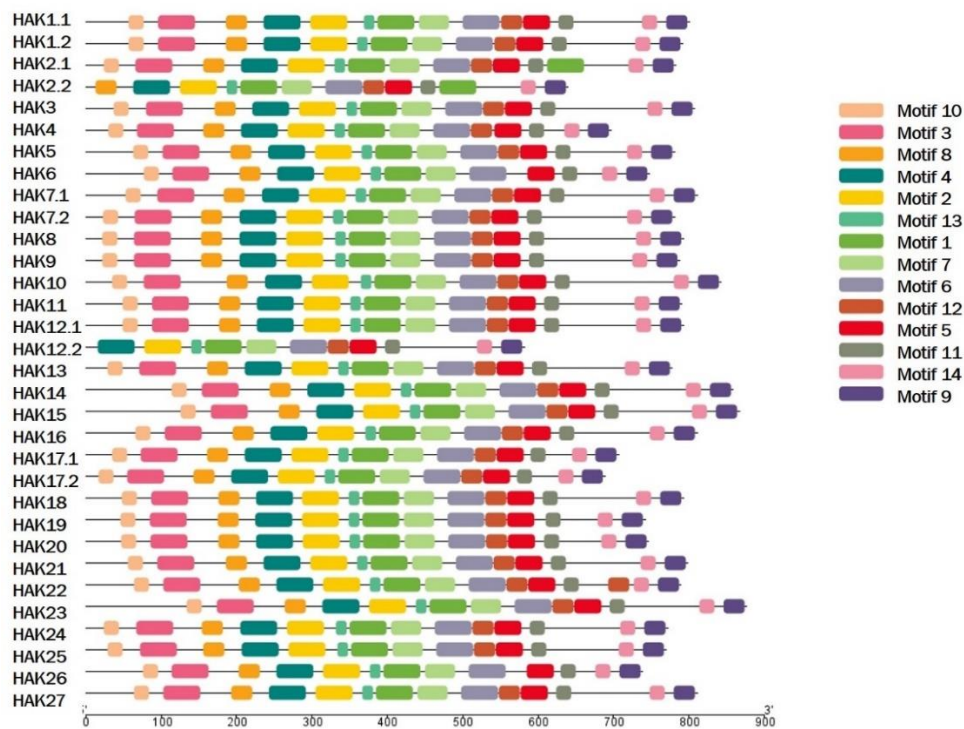


Fig. 2: Functional motif analysis among the OSHAK family using MEME Suite.

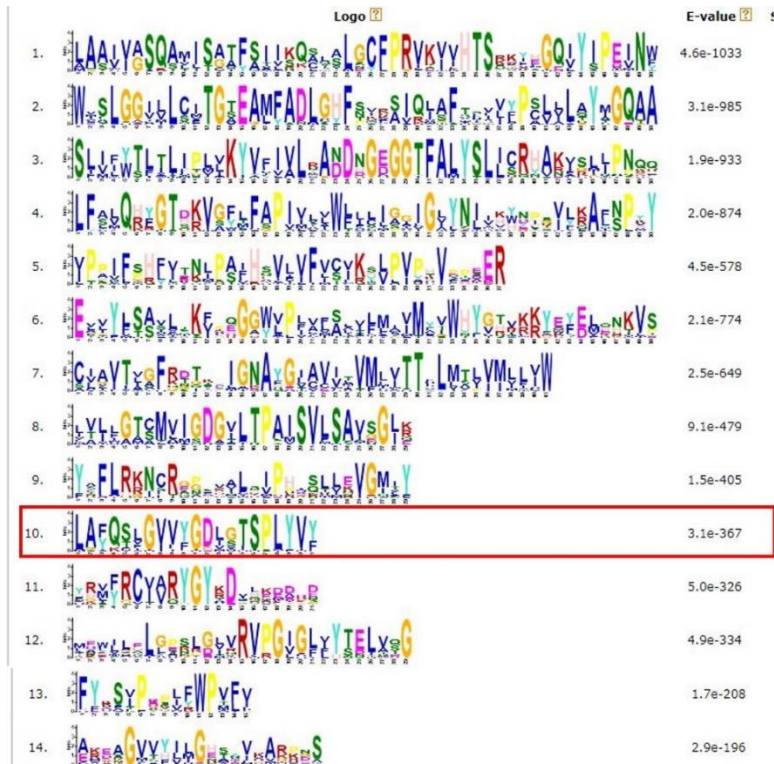


Fig.3: Consensus sequence observed among the OSHAK transporters in rice.

The red box on motif 10 reveals the consensus sequence for the OSHAK transporter family

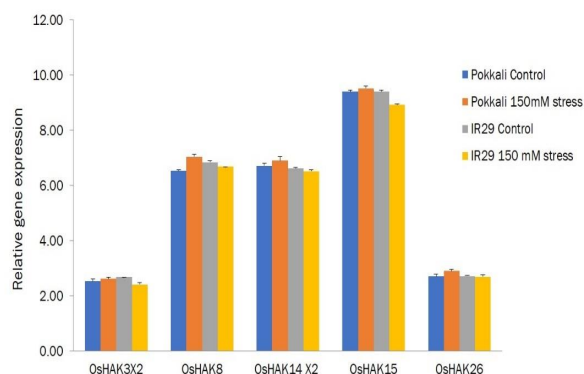


Fig. 4: Differentially expressed *OshAK* transporters between *Pokkali* and cv. *IR29* under salt stress

A study conducted by Wang *et al.* (2021) revealed the major play of *OshAK8* in potassium uptake as well as root-to-shoot K^+ transport within the plant. The expression level of *OshAK8* was significantly downregulated upon salt stress in *OshAK8* mutants and K^+ uptake was impaired. In another study by Gupta *et al.* (2008), *OshAK8* was found to be differentially expressed upon salt stress, especially in panicles implicating its role in panicle development and in turn crop productivity.

In this study, we observed that *OshAK14* and *OshAK15* were grouped under the clade I. A genome-wide association analysis conducted by Li *et al.* (2019) within *japonica* cultivars at the seedling stage revealed that *OshAK15* plays a key role in salinity tolerance in rice besides, demonstrating a significant homology with *OshAK14*. This probably reflects on the co-expression of *OshAK14* and *OshAK15* as observed from gene expression analysis. Earlier, Baneulos *et al.* (2002) also reported that *OshAK15* shared a close homology with *OshAK14*.

Further analysis of these differentially expressed genes was performed to explore the tissue-specific expression in rice seedlings under salt stress using EXPath 2.0. *OshAK14* was found to be very highly expressed in the leaf sheath and root upon salt stress followed by *OshAK15*. The other differentially expressed *OshAKs* viz., *OshAK3* and *OshAK8* showed only a moderate level of expression as compared to *OshAK14* and *OshAK15* whereas, *OshAK26* showed the least in both leaves and roots. Majorly, the differentially expressed set of transporters exhibited a low level of expression in the leaf sheath as compared to the leaf blade and roots during the vegetative stage (Fig. 5).

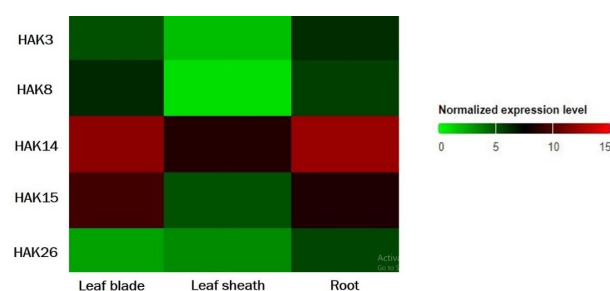


Fig 5: Tissue-specific expression levels of selected *OshAK* transporters in rice seedlings under salt stress

A study on pollen development and fertility by Li *et al.* (2022) implicated the role of golgi bound *OshAK26* to play a major role in anther development. Knockout of *OshAK26* in *Nipponbare* caused a reduction in the number, viability, and germination of pollen grains. As the *OshAK26* plays a major role in pollen development, they might be expressed in anthers and this could be the reason for the reduced level of expression of *OshAK26* within the vegetative tissues of the plant.

EXPath 2.0 analysis also paved the way for exploring the transcript abundance of *OshAK3(X2)*, *OshAK8*, *OshAK14(X2)*, *OshAK15* and *OshAK26* under salt stress (Fig. 6 and Fig. 7). Among the five differentially expressed set of genes, *OshAK8*, *OshAK14* and *OshAK15* were significantly upregulated in the roots as compared to *OshAK3* and *OshAK26*. In the case of shoots, *OshAK8* is found to be significantly upregulated compared to other HAK transporters such as *OshAK3*, *OshAK14*, *OshAK15*, and *OshAK26*.

Most of the differentially expressed set of transporters showed significant upregulation in roots as compared to shoots and therefore, these genes might play a role in enhancing the root K^+ uptake under salt stress. Thus, identification and characterization of differentially expressed genes involved in K^+ uptake as well as transport and their subsequent introgression into elite rice cultivar(s) help in improving the K/Na ratio thereby leading to salt tolerance in rice.

CONCLUSION

In silico analysis of the *OshAK* transporter gene family indicated the upregulation of a few selected transporters such as *OshAK3(X2)*, *OshAK8*, *OshAK14(X2)*, *OshAK15* and *OshAK26* under salt stress in salt-tolerant rice landrace *Pokkali* compared

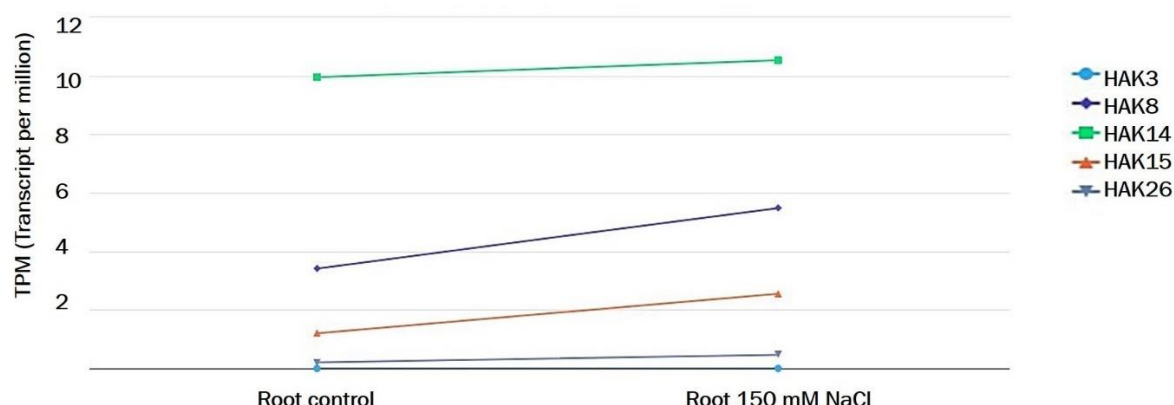


Fig.6: Transcript abundance levels of selected *OsHAK* transporters in roots of rice seedlings under salt stress

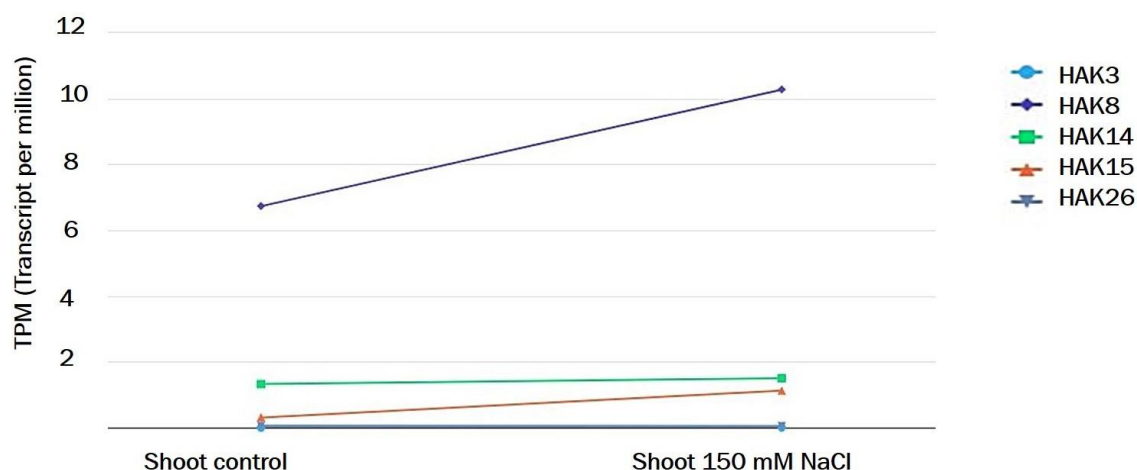


Fig.7: Transcript abundance levels of selected *OsHAK* transporters in shoots of rice seedlings under salt stress

to the salt susceptible cv. IR29. In addition, the tissue-specific expression of these differentially expressed transporters is suggestive of their expression confined to the leaf blade as well as roots. Further experimental validation of the differentially upregulated set of *OsHAK* transporters through real-time PCR may pave the way for exploiting their usefulness in translational research leading to rice improvement against salt stress.

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Conflict of interest

The authors disclose no conflict of interest.

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