The Rice High-Affinity Potassium Transporter1;1 Is Involved in Salt Tolerance and Regulated by an MYB-Type Transcription Factor^{1[OPEN]}

Rong Wang, Wen Jing, Longyun Xiao, Yakang Jin, Like Shen, and Wenhua Zhang*

State Key Laboratory of Crop Genetics and Germplasm Enhancement, College of Life Sciences, Nanjing Agricultural University, Nanjing 210095, People's Republic of China

ORCID IDs: 0000-0002-9407-6895 (L.X.); 0000-0003-3122-6152 (W.Z.).

Sodium transporters play key roles in plant tolerance to salt stress. Here, we report that a member of the High-Affinity K⁺ Transporter (HKT) family, OsHKT1;1, in rice (*Oryza sativa* 'Nipponbare') plays an important role in reducing Na⁺ accumulation in shoots to cope with salt stress. The *oshkt1;1* mutant plants displayed hypersensitivity to salt stress. They contained less Na⁺ in the phloem sap and accumulated more Na⁺ in the shoots compared with the wild type. *OsHKT1;1* was expressed mainly in the phloem of leaf blades and up-regulated in response to salt stress. Using a yeast one-hybrid approach, a novel MYB coiled-coil type transcription factor, OsMYBc, was found to bind to the *OsHKT1;1* promoter. In vivo chromatin immunoprecipitation and in vitro electrophoresis mobility shift assays demonstrated that *OsMYBc* binds to AAANATNC(C/T) fragments within the *OsHKT1;1* promoter. Mutation of the *OsMYBc*-binding nucleotides resulted in a decrease in promoter activity of *OsHKT1;1*. Knockout of *OsMYBc* resulted in a reduction in NaCl-induced expression of *OsHKT1;1* and salt sensitivity. Taken together, these results suggest that OsHKT1;1 has a role in controlling Na⁺ concentration and preventing sodium toxicity in leaf blades and is regulated by the *OsMYBc* transcription factor.

Soil salinity is an abiotic stress that negatively affects plant growth and development, thus posing a serious threat to crop productivity (Munns et al., 2012). The adverse effects of high concentrations of salt on plants include osmotic stress, ionic toxicity, and nutritional imbalance (Munns and Tester, 2008). Sodium is taken up by the plant root system and transported to shoots via the transpiration stream (Tester and Davenport, 2003; Deinlein et al., 2014).

The mechanisms of influx of Na⁺ into the root system are not understood. It is thought that Na⁺ influx into root cells is in part via the voltage-independent, nonselective cation channels, such as the cyclic nucleotidegated channels (Apse and Blumwald, 2007; Ward et al., 2009; Jin et al., 2015). For glycophytes, the mechanisms of salt tolerance include the ability to limit Na⁺ accumulation on the shoot, exclude Na⁺ from the cytoplasm of cells, and sequester Na⁺ into the vacuoles (Hasegawa, 2013). Intracellular Na⁺ is exported out of the cell by the Salt Overly Sensitive1 plasma membrane Na⁺/H⁺ antiporter (Shi et al., 2000) or sequestered into the vacuole via the tonoplast Na^+/H^+ antiporter1 (Apse et al., 1999). At the tissue level, regulation of Na^+ loading into the root xylem is essential for limiting Na^+ accumulation in the shoot. Members of the high-affinity potassium transporter (HKT) family of transport proteins, encoded by AtHKT1;1 from Arabidopsis (Arabidopsis thaliana) and OsHKT1;5 from rice (Oryza sativa), reduce the transport of Na⁺ to shoots and positively regulate salt tolerance (Uozumi et al., 2000; Ren et al., 2005). Similar mechanisms have been reported in wheat (Triticum aestivum) for the *TmHKT1;4* and *TmHKT1;5* genes (Huang et al., 2006; Munns et al., 2012; Byrt et al., 2014).

Plant HKTs are allocated to two subfamilies (Platten et al., 2006). Subfamily 1 exists in monocotyledonous and dicotyledonous species, comprising Na⁺-selective transporters. Subfamily 2 is present in monocotyledonous species and comprises transporters permeable to both Na⁺ and K⁺ (Hauser and Horie, 2010). Rice contains seven to nine HKT transporters, depending on the variety (Platten et al., 2006; Hauser and Horie, 2010). Functional analyses in yeast (*Saccharomyces cerevisiae*) and *Xenopus laevis* oocytes reveal striking diversity. Subfamily 1 members OsHKT1;1, OsHKT1;3, and OsHKT1;5

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^{*} Address correspondence to whzhang@njau.edu.cn.

The author responsible for distribution of materials integral to the findings presented in this article in accordance with the policy described in the Instructions for Authors (www.plantphysiol.org) is: Wenhua Zhang (whzhang@njau.edu.cn).

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are permeable to Na⁺ only (Garciadeblás et al., 2003; Ren et al., 2005; Jabnoune et al., 2009). OsHKT2;1, which belongs to subfamily 2, displays diverse permeation modes, Na⁺-K⁺ symport, Na⁺ uniport, or inhibited states, depending on external Na⁺ and K⁺ concentrations (Horie et al., 2001, 2007; Garciadeblás et al., 2003; Jabnoune et al., 2009). Heterologous expression of OsHKT2;4 in *X. laevis* oocytes was reported to give rise to Ca²⁺ and Mg²⁺ membrane transport activity (Lan et al., 2010; Horie et al., 2011). As such, HKT proteins were predicted to participate in Ca²⁺ signaling in plant cells (Lan et al., 2010). However, the work of Sassi et al. (2012) suggests that OsHKT2;4 is a new functional HKT member, endowed with high K⁺ permeability and a particularly low Na⁺ permeability. Relatively little is known about HKT transporter

functions in planta. Reverse genetics approaches in Arabidopsis and analysis of quantitative trait loci for salt tolerance in rice have highlighted the roles, in planta, of the HKT transporters AtHKT1;1 and OsHKT1;5. These HKT transporters positively regulate salt tolerance by retrieving Na⁺ from the ascending xylem sap, thus limiting Na⁺ levels in the shoots (Uozumi et al., 2000; Ren et al., 2005; Møller et al., 2009). Crossing of the *TmHKT1;5-A* (*Na*⁺ *exclusion2*) gene locus from the wheat relative Triticum monococcum into a commercial durum wheat (Triticum durum) significantly reduces leaf Na⁺ content and increases durum wheat grain yield by 25% when grown in saline soil (Munns et al., 2012). These results suggest that similar xylem Na⁺-unloading mechanisms are essential for salt tolerance in monocotyledonous and dicotyledonous species.

By analyzing transposon-insertion rice mutants, OsHKT2;1 has been identified as a central transporter for nutritional Na⁺ uptake into K⁺-starved rice roots. However, the OsHKT2;1-mediated Na⁺ influx does not cause Na⁺ toxicity, as its transcription is down-regulated in the presence of salt stress (Horie et al., 2007). In barley (*Hordeum vulgare*), overexpression of *HvHKT2;1* enhances Na⁺ uptake in shoots and increases plant growth in the presence of 50 to 100 mM NaCl (Mian et al., 2011).

The expression of HKT genes is sensitive to K⁺ starvation and osmotic or salt stress (Wang et al., 1998; Ren et al., 2005; Sunarpi et al., 2005; Horie et al., 2007). Regulatory mechanisms of AtHKT1;1 expression have been identified recently. They include hormone regulation, transcription regulation, and DNA methylation (Baek et al., 2011; Shkolnik-Inbar et al., 2013). AtHKT1;1 is repressed by plant hormone cytokinin treatment but shows significantly elevated expression in the cytokinin Arabidopsis response regulator double mutant arr1-3 arr12-1. These data suggest that cytokinin, acting through the transcription factors ARR1 and ARR12, negatively regulate the expression of *AtHKT1;1* in roots and Na⁺ accumulation in shoots (Mason et al., 2010). ABSCISIC ACID-INSENSITIVE4 (ABI4) is also a transcription factor involved in the abscisic acid response. Recently, Shkolnik-Inbar et al. (2013) found higher *AtHKT1;1* expression (in xylem parenchyma cells) in *abi4* mutant plants and lower levels of expression in *ABI4*-overexpressing plants, suggesting that ABI4 represses *AtHKT1;1* expression. The increased salt tolerance of *abi4* mutants is believed to result from increased AtHKT1;1 activity, which leads to increased Na⁺ unloading from the xylem vessels and reduced shoot accumulation of Na⁺ (Shkolnik-Inbar et al., 2013).

In this study, we used genetic tools to understand the contribution of *OsHKT1;1* to salt tolerance. In addition, a novel transcription factor that binds to the *OsHKT1;1* promoter and regulates both its expression and salt tolerance was identified.

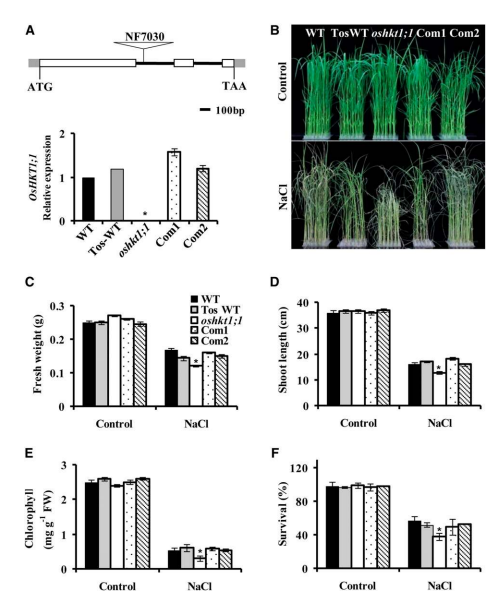
RESULTS

oshkt1;1 Mutants Exhibit Sensitivity to Salt Stress

The expression pattern of *OsHKT1;1* and its activity in X. laevis oocytes have been investigated (Jabnoune et al., 2009). However, the in vivo function of OsHKT1;1 in rice is unknown. A retrotransposon (Tos17) insertion mutant of OsHKT1;1 (line NF7030) was identified in the *Tos*17 Insertion Mutant Database (https://tos.nias.affrc. go.jp/index.html.en; Miyao et al., 2003). The Tos17 retrotransposon insertion resided within the first intron region of OsHKT1;1 (Fig. 1). We also isolated a related OsHKT1;1 wild-type control plant, named TosWT (Horie et al., 2007). Southern-blot analysis indicated that TosWT had the same *Tos17* insertions as *oshkt1;1* except for an insertion in the OsHKT1;1 gene (Supplemental Fig. S1A). Quantitative reverse transcription (qRT)-PCR analysis showed that OsHKT1;1 expression was abolished in oshkt1;1 but not affected in TosWT (Fig. 1A, bottom). The seedlings were hydroponically grown in the presence of 100 mM NaCl for 7 d; the *oshkt1;1* mutant showed sensitivity to salt stress, displaying reduced growth, decreased fresh weight and length of shoots, and lower chlorophyll content as compared with the wild type and TosWT (Fig. 1, B–E). The salttreated seedlings were recovered for an additional 7 d, and more seedlings of *oshkt1;1* died than those of the wild type and TosWT (Fig. 1F). TosWT showed a similar growth pattern in the presence or absence of salt stress (Fig. 1, B–F), suggesting that other Tos17 insertions did not obviously affect the growth and salt tolerance. In the following physiological experiments, we used the wild-type control only.

To further confirm the functions of *OsHKT1;1*, we transformed a 4.3-kb fragment to the *oshkt1;1* mutant that contained the 2.2-kb region with exons and introns of *OsHKT1;1* and a 2.1-kb upstream promoter sequence of the gene. The transformation was confirmed by qRT-PCR (Fig. 1A). Southern-blot analysis revealed that each complementary line had only one copy of *OsHKT1;1* in its genomic sequence (Supplemental Fig. S1B). The complementation of *OsHKT1;1* recovered the salt tolerance of the *oshkt1;1* mutant (Fig. 1, B–F). Taken together, these results suggest that *OsHKT1;1* confers salt tolerance to rice.

Figure 1. Knockout of OsHKT1;1 results in salt sensitivity. A, Isolation of homozygous Tos17 insertion mutants of the OsHKT1;1 gene. Top, schematic diagram of the oshkt1;1 mutant. The white boxes stand for exons, the lines stand for introns, and the triangle indicates the Tos17 insertion. Bottom, qRT-PCR analysis of OsHKT1;1 expression in wild-type (WT), TosWT, oshkt1;1 mutant, and OsHKT1;1-COM (Com1 and Com2) plants. Total RNA was extracted from the whole plant. The genes OsUbiquitin5 (OsUBQ5) and 18S Ribosomal RNA (18S rRNA) were used as internal controls. B to E, Phenotypes of wild-type, TosWT, oshkt1;1 mutant, and OsHKT1;1-COM plants under salt stress. Twenty-one-day-old hydroponically grown seedlings were treated with 100 mm NaCl for 7 d. B, Representative photographs of plants. C to E, Fresh weight (C), shoot length (D), and total chlorophyll content (E). The data represent means \pm se (n = 40-50for each line from three replicates). FW, Fresh weight. F, Survival rate of seedlings. Plants were grown under 100 mM NaCl for 7 d and recovered in culture solution without NaCl for an additional 7 d. The data represent means \pm sE from three biological repeats, each consisting of 50 seedlings of each line. Asterisks indicate statistically significant differences compared with the wild type: *, *P* < 0.05.



The *oshkt1;1* Mutation Results in More Accumulation of Sodium in Shoots

The regulation of Na⁺ content in shoots is essential for salt tolerance, and OsHKT1;1 is permeable to Na⁺ according to analyses in X. laevis oocytes (Jabnoune et al., 2009). We asked whether OsHKT1;1 is involved in the control of Na⁺ accumulation in the shoots. We next determined shoot Na⁺ content in the wild type and oshkt1;1 under saline conditions. As shown in Figure 2A, the *oshkt1;1* mutant accumulated more Na⁺ in the shoot than the wild type. In contrast, introduction of the native OsHKT1;1 gene into oshkt1;1 plants led to complementation of the Na⁺ content of *oshkt1;1* shoots (Fig. 2A). These results suggested that OsHKT1;1 is essential for Na⁺ accumulation in the shoot. We further determined the Na⁺ accumulation in leaf blade and sheath of complete leaves 1 and 2 separately. The oshkt1;1 mutant accumulated more Na⁺ in leaf blades

1 and 2 as compared with the wild type. In the sheaths of both leaves 1 and 2, however, the *oshkt1;1* mutant and the wild type stored similar Na⁺ contents (Supplemental Fig. S2; Supplemental Table S1). Therefore, knockout of *OsHKT1;1* resulted in more Na⁺ accumulation in leaves largely ascribed to the Na⁺ in leaf blades.

To investigate the mechanisms leading to Na⁺ accumulation in the shoot, we measured Na⁺ concentration in sap from the phloem and xylem after rice plants were treated with NaCl for 2 d. For Na⁺ in phloem sap, Gln was used as an internal standard to correct the errors caused by variation in volume of the phloem sap, and the Na⁺ amount was represented by a relative level as Na⁺-Gln ratio (Berthomieu et al., 2003; Ren et al., 2005). A reduction in the Na⁺ content of phloem sap was observed in the *oshkt1;1* mutant compared with the wild type. Transformation of *oshkt1;1* with the native *OsHKT1;1* gene complemented the Na⁺ reduction in phloem sap (Fig. 2B; Supplemental

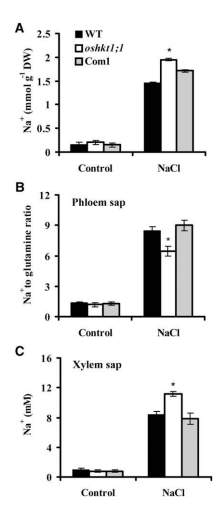


Figure 2. Na⁺ content in rice plants. A, Na⁺ content in shoots. Hydroponically grown seedlings were treated with 100 mM NaCl for 7 d, and the shoots were harvested for Na⁺ content assay. Error bars represent se (n = 5). DW, Dry weight. B, The *oshkt1;1* mutant plants show a decreased Na⁺ content in the phloem sap. Seedlings were hydroponically grown in culture solution for 21 d, then for 2 d in culture solution supplemented with 25 mM NaCl, before phloem sap was collected. One sample contained phloem sap from four seedlings. Gln content was used as an internal standard. Error bars represent se (n = 4). C, The *oshkt1;1* mutant plants display an increased Na⁺ concentration in the xylem sap. Seedlings. Error bars represent se (n = 4). Asterisks indicate statistically significant differences compared with the wild type (WT): *, P < 0.05.

Table S2). Disruption of the *OsHKT1;1* gene caused an increase in the Na⁺ concentration of xylem sap, which was decreased to the wild-type level by complementation of the native *OsHKT1;1* gene to the *oshkt1;1* mutant (Fig. 2C). These results suggested that OsHKT1;1 is probably involved in the control of Na⁺ concentrations in the phloem and xylem sap.

OsHKT1;1 Expression Pattern in Accordance with Its Function

To further explore whether the effects of OsHKT1;1 rely on its localization in plants, we investigated the spatial expression of the OsHKT1;1 gene by analyzing the transgenic rice lines containing the GUS reporter gene under the control of the OsHKT1;1 promoter. The results showed that the GUS activity was active in the vascular tissues of both leaves and roots (Fig. 3A, a and d). For the root, GUS activity was very weak in the tip region (Fig. 3A, d). Transverse cross sections of the leaf revealed that the majority of GUS signals were detected in the stele, mainly associated with the phloem and some with the xylem parenchyma cells (Fig. 3A, c). This result is identical to a previous observation by Jabnoune et al. (2009) using in situ hybridization methods. In the roots, OsHKT1;1 gene expression was identified mainly in the stele, including the cells surrounding protoxylem and phloem cells. Weak OsHKT1;1 expression was also found in the sclerenchyma (Fig. 3A, e and f). We next transiently coexpressed OsHKT1;1-GFP with the plasma membrane marker CBL1n-mCherry (red fluorescent protein; Held et al., 2011) in onion (*Allium cepa*) epidermal cells. GFP and mCherry signal showed an obvious overlap (Supplemental Fig. S3A; Supplemental Methods S1). OsHKT1;1-GFP was then coexpressed with CBL1nmCherry in Arabidopsis protoplasts. Most of the GFP signal overlapped with mCherry (Supplemental Fig. S3B). Only a small fluorescent signal was also observed inside the cells (Supplemental Fig. S3B), which may reflect endoplasmic reticulum localization, where the protein (e.g. OsHKT2;4) was assembled (Horie et al., 2011). These results suggested that the OsHKT1;1-GFP fusion protein was associated with the plasma membrane.

We next performed qRT-PCR to determine *OsHKT1;1* expression in different tissues and under saline conditions. The results showed that *OsHKT1;1* expression in shoots was 11-fold greater than that in roots (Fig. 3B). *OsHKT1;1* expression was increased approximately 3- to 5-fold by salt treatment in shoots but not in roots (Fig. 3, C and D).

In Vivo Analysis of the OsHKT1;1 Promoter by Agroinfiltrated Nicotiana benthamiana Leaves

To investigate the regulation of the expression of *OsHKT1;1* under salt stress, we applied a transient expression system using agroinfiltration of *N. benthamiana*. The goal was to identify the enhancer element in the promoter. The transient expression system of *N. benthamiana* and Arabidopsis is suitable for analyzing cis-element responses to stress stimuli, which include salt stress, pathogen infection, and heat shock (Yang et al., 2000; Tsuda et al., 2012). We constructed four 5' terminal deletion mutants of the *OsHKT1;1* promoter fused to the GUS reporter (Fig. 4A).

The *CaMV35S* promoter was used as a negative control, and the promoter of the NaCl up-regulated gene *Responsive to Desiccation 29A* (*RD29A*) was used as a positive control (Narusaka et al., 2003). All constructs were expressed uniformly in *N. benthamiana*

leaves. After expression for 60 h, the *N. benthamiana* leaves were cut into two parts. The left half was dipped into 100 mM NaCl solution and the right half was dipped into blank (water) solution for 2 h. Promoter activity was then determined by GUS staining (Fig. 4B) or measuring GUS activity (Fig. 4C). The leaves expressing ProCaMV35S:GUS showed strong GUS activity, with no differences between the NaCl- and water-treated parts. In contrast, the GUS activity of the *RD29A* and *OsHKT1;1* promoters in NaCl-treated leaves was greater than that in water-treated leaves. Although slightly lower compared with other *OsHKT1;1* promoters, Pro*OsHKT1;1*-629 retained the majority of the *OsHKT1;1* promoter activity and inducibility (Fig. 4C).

Characterization of the MYB-Like Transcription Factor

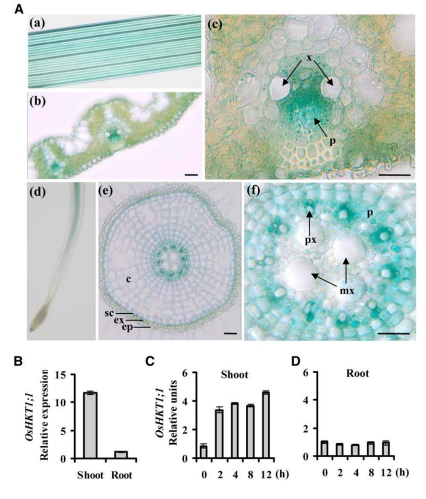
To identify the putative transcription factors regulating *OsHKT1;1* expression in response to salt treatment, we constructed a rice complementary DNA (cDNA) library and applied a yeast one-hybrid (Y1H) approach to search for novel proteins associated with the *OsHKT1;1* promoter. A 617-bp *OsHKT1;1* promoter

Figure 3. Expression pattern of OsHKT1;1 in tissues. A, Expression pattern of OsHKT1;1 promoter-GUS in transgenic plants. a, Leaves. b and c, Cross sections of leaves. GUS activity was detected mainly in vascular tissues of leaves (b). The positions of the phloem (p) and xylem (x) region are indicated by arrows (c). d, Expression of OsHKT1;1 in roots. e and f, Cross sections of roots. GUS activity was detected mainly in stele of root, including phloem and xylem parenchyma cells. c, Cortex; ep, epidermis; ex, exodermis; mx, metaxylem; p, phloem; px, protoxylem; sc, sclerenchyma; x, xylem. Bars = 50 μ m. B, Expression pattern of *OsHKT1;1* analyzed by qRT-PCR. Error bars represent se (n = 3). C and D, Time-course expression analysis of OsHKT1;1 in response to NaCl (100 mm) treatment. OsHKT1;1 expression in plants without NaCl was used as a reference of the basal expression level. The genes UBQ5 and 18S rRNA were used as internal controls. Error bars represent se (n = 3).

upstream of the ATG start codon, plus a 17-bp region downstream of the ATG, were used to screen the rice cDNA library. Positive interactions are listed in Supplemental Table S3. Of the 26 positive clones analyzed, seven originated from the same gene (LOC_Os09g12770), corresponding to two independent cDNAs of different sizes. LOC_Os09g12770 encodes a MYB-like transcription factor.

To confirm the interaction between *OsHKT1;1* and the MYB-like transcription factor LOC_Os09g12770, we performed the Y1H assay a second time, using a full-length coding sequence of the *MYB*-like gene. We used the 629-bp promoter fragment of *OsHKT1;1* as bait. The GoldH1Y yeast stain containing the bait was transferred with the MYB-like-pGADT7AD and grew normally on the selective medium containing aureobasidin A (AbA). The negative control (empty pGADT7AD vector) did not grow (Fig. 5A). These results suggest that the MYB-like transcription factor may interact with the promoter of *OsHKT1,1*.

The MYB-like transcription factor contains an open reading frame of 289 amino acid residues, with a calculated molecular mass of 31 kD. Structural analysis using the Rice Genomic Annotation Program showed that this protein contains an MYB-related DNA-binding



domain (residues 25–76) and a coiled-coil dimerization domain from residues 112 to 133, belonging to the MYB-CC family of transcription factors (Fig. 5B, top). We named this MYB-like protein as OsMYBc. Phylogenetic tree analysis showed the MYB-CC family members in plants (Fig. 5B, bottom). Some members have been identified, such as PHOSPHATE STARVATION RESPONSE (PHR) transcription factors from Arabidopsis, rice, maize (*Zea mays*), and wheat (Rubio et al., 2001; Zhou et al., 2008; Wang et al., 2013), and ALTERED PHLOEM DEVELOPMENT1 (APL1) from Arabidopsis (Bonke et al., 2003).

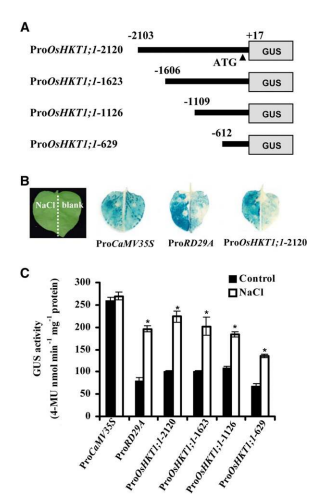


Figure 4. Salt stress induces *OsHKT1;1* promoter activity. A, The schematic diagram shows the constructs of 5' terminal deletion mutants of the *OsHKT1;1* promoter that are linked with the reporter gene *GUS*. B, GUS image of *N. benthamiana* leaves. The *N. benthamiana* leaves were infiltrated with agrobacterial stains containing constructs with different promoters indicated under the images at right. These leaves were separated into left and right parts as indicated in the image at left. The left part of each leaf was treated with 100 mM NaCl, and the right part was immersed in blank solution for 2 h. The treated leaves were stained with 5-bromo-4-chloro-3-indolyl- β -glucuronic acid and cleared with ethanol. Representative images are shown. C, GUS activity assay of the *N. benthamiana* leaves indicated in B. Asterisks indicate that the mean value is significantly different from that of the control: *, P < 0.05. Error bars represent st (n = 3). 4-MU, 4-Methylumbelliferone.

To examine OsMYBc localization, we expressed the OsMYBc-GFP fusion protein in onion epidermal cells. The OsMYBc-GFP protein accumulated mainly in the nucleus (Fig. 5C). To determine whether *OsMYBc* transcript levels were affected by salt stress, wild-type plants were treated with 100 mM NaCl and the expression patterns of *OsMYBc* were measured by qRT-PCR. *OsMYBc* transcription was induced by salt stress in both shoots and roots. With the prolongation of exposure to salt treatments, *OsMYBc* transcription decreased gradually (Fig. 5, D and E).

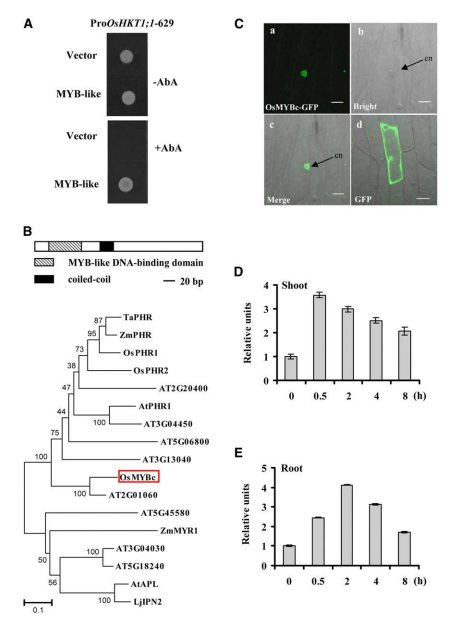
OsMYBc Protein Binds to a Specific Sequence in the Promoter of OsHKT1;1

The interaction of the OsMYBc protein with the promoter fragment of OsHKT1;1 (-612 to +17) suggests the presence of a cis-acting element in this region. We performed an electrophoretic mobility shift assay (EMSA) to identify the OsMYBc-binding element. The promoter fragment (-612 to +17) was partitioned into three regions, identified as F1 to F3 (Fig. 6A, top), and labeled with digoxigenin. An initial screen showed that the OsMYBc-GST fusion protein bound to the OsHKT1;1 promoter regions at -412 to -184 (F2) and -183 to +17 (F3) but not the region at -612 to -413(F1; Fig. 6A, bottom). The binding was specific, since it could be competed off with unlabeled probe (Fig. 6A, bottom). The F3 region was further divided into five 40-bp segments, a, b, c, d, and e, and banding was detected in probe c (Fig. 6B, lane 3).

We then compared the nucleotide sequences between segment c and the F2 region of the OsHKT1;1 promoter to identify OsMYBc-binding nucleotides. A 10-bp conserved segment, AAATATGC(C/T)A, was found (Fig. 6C). We then performed site mutation(s) for the 10 bp in fragment c and used a number of mutated fragments of segment c as probes to further characterize the OsMYBc-binding sites. When AAA (probe M1), TAT (probe M2), GC (probe M3), and CA (probe M4) were mutated, OsMYBc binding was prevented (Fig. 6E, lanes 2-5). The results indicate that at least four nucleotides, between positions -76 and -68, are essential for OsMYBc binding. We then performed single nucleotide mutation for the 10 nucleotides in fragment c. Mutation of the first two A nucleotides (probes M5 and M6) obviously reduced OsMYBc binding. The mutation of M7, M9, M10, M12, and M13 resulted in inhibiting the binding to OsMYBc, suggesting that the nucleotides A (M7), AT (M9 and M10), and CC (M12 and M13) are required for OsMYBc binding. By contrast, the presence of M8, M11, and M14 did not affect binding to OsMYBc (Fig. 6E). Therefore, the OsMYBc-binding region in fragment c is AAANATNCC (where N represents a changeable nucleotide). When the first C nucleotide of AAANATNCC was replaced by T, OsMYBc still bound to region F2 (Fig. 6, A and C). Taken together, the OsMYBc protein binds mainly to the region AAANATNC(C/T), the MYB-binding sequence,

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Figure 5. Characterization of MYBc-like protein. A, MYB-like protein binds the OsHKT1;1 promoter in yeast cells. The OsHKT1;1-629 promoter was linked to the Aureobasidin 1-C reporter gene that confers AbA resistance in yeast cells. The MYB-like gene isolated from the rice cDNA library was cloned and linked to the pGADT7AD vector as effector. After incubating plates on synthetic dropout-Leu plates with or without 250 ng mL^{-1} AbA for 3 d at 30°C, colonies were visualized. B, Description of OsMYBc. Top, schematic structures of OsMYBc. Bottom, the phylogenetic tree constructed using MEGA4.0 software. Sequences were found on the Phytozome (http:// www.phytozome.net/), The Arabidopsis Information Resource (http://www.arabidopsis.org/), and the Rice Genomic Annotation Project (http://rice. plantbiology.msu.edu/) databases. OsMYBc is boxed. At, Arabidopsis; Lj, Lotus japonicus; Os, rice; Ta, wheat; Zm, maize. C, Localization of OsMYBc protein in onion epidermal cells. Individual images show OsMYBc-GFP (a), bright field (b), merge (c), and GFP alone (d). Bars = 50 µm. cn, Cell nucleus. D and E, Salt stress regulates the expression of OsMYBc. Twenty-one-dayold plants were treated with 100 mM NaCl for 0, 0.5, 2, 4, and 8 h. gRT-PCR was performed by using the cDNA derived from shoots and roots separately. Plants cultured without NaCl were used as a reference of basal expression. The genes UBQ5 and 18S rRNA were used as internal controls. Error bars indicate se (n = 3).



between -275 to -267 and -76 to -68 of the *OsHKT1;1* promoter.

We next examined whether the cis-element also exists in other *HKT* genes. Searching the promoters (up to 2.5 kb from the start codon) of these genes led to the finding that the cis-element exists in the promoter region of four of seven *OsHKT* genes as well as two *HKT* genes from wheat and maize (Supplemental Table S4). For example, the MYB-binding sequence (AAATATTCC), present in the *OsHKT2*;1 gene, was bound by *OsMYBc* (Supplemental Fig. S4).

OsMYBc Binding to the OsHKT1;1 Promoter in Vivo

To confirm that OsMYBc binds directly to the OsHKT1;1 promoter in vivo, chromatin immunoprecipitation (ChIP)

assays were performed. The OsMYBc-Flag fusion protein was expressed in rice protoplasts and immunoprecipitated using an antibody recognizing the Flag tag. The genomic DNA fragments that coimmunoprecipitated with OsMYBc-Flag were analyzed by PCR. Fragment I (-422 to +13), which contains two cis-elements, was detected. However, the negative control fragment II (+226 to +665), which does not contain the cis-element, was not detected (Fig. 7, A and B).

Y1H, EMSA, and ChIP analyses demonstrated that OsMYBc binds to the *OsHKT1;1* promoter. To confirm the role of OsMYBc in the regulation of *OsHKT1;1* expression, we performed transient GUS assays in *N. benthamiana* as reported by Shim et al. (2013). OsMYBc, with the *CaMV35S* promoter, was used as the effector. *OsHKT1;1*-629 (P) with two cis-elements (AAATATGCC and AAATATGCT) or their mutated versions (mP)

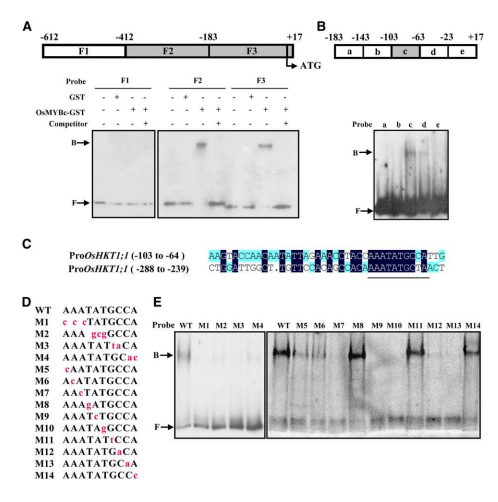


Figure 6. EMSA shows OsMYBc binding to specific *OsHKT1;1* promoter DNA fragments. A, Top, schematic diagram of the *OsHKT1;1* promoter fragment from -612 to +17 bp (translation start is +1) used in EMSA. F1 to F3 regions were prepared by PCR and labeled with digoxigenin. The region that did not bind to OsMYBc is shown as the white box, and those binding to OsMYBc are shown as gray boxes. Bottom, EMSA results. Glutathione *S*-transferase (GST)-OsMYBc fusion protein was expressed in *E. coli*. The labeled probes were also competed by excess unlabeled probes (lanes 4, 8, and 12). Arrows show OsMYBc-bound or free probe (B or F, respectively). B, Top, schematic diagram of five segments of the F3 fragment. Bottom, segment c shows binding activity with OsMYBc (lane 3). C, Alignment of the DNA sequence between segment c and a specific region from the F2 fragment. The highly related sequence is underlined. D and E, Identification of the OsMYBc-binding region in the *OsHKT1;1* promoter by base mutations M1 to M4, M7, M9, M10, M12, and M13 lost the function of binding with OsMYBc; mutations M5 and M6 had reduced binding; and mutations M8, M11, and M14 were not affected in binding. WT, The wild type.

fused to the GUS gene were used as reporters (Fig. 7, C and D). The reporter and effector plasmids were coinfiltrated into *N. benthamiana* leaves. The GUS reporter gene was activated by coexpressing *OsMYBc* with the wild-type promoter *OsHKT1;1-629* (P). However, the mutant reporter containing two cis-element mutations from AAATATGCC/T to AAAGCGGCC/T in *OsHKT1;1-629* (mP) was not activated (Fig. 7, D–F). These results demonstrate that *OsMYBc* can activate promoter transcription by interacting with the cis-element.

Expression of OsHKT1;1 and Salt Tolerance Are Reduced in osmybc Plants

We next investigated the genetic mechanism of OsMYBc regulation of OsHKT1;1 expression. A transfer

DNA (T-DNA) insertion mutant line, *osmybc* (K-00126), was isolated and confirmed by RNA expression (Fig. 8A, a and b). Southern-blot analysis indicated a one-copy insertion in the genomic DNA (Fig. 8A, c). Knockout of the *OsMYBc* gene did not affect the growth under normal conditions but resulted in salt sensitivity as compared with the wild type (cv Kitaake; Fig. 8, B–D). The *osmybc* mutant accumulated more Na⁺ in shoots than the wild type (Fig. 8, E and F).

We next investigated the genetic mechanism for *OsMYBc* regulation of *OsHKT1;1* expression. As *OsMYBc* increased the transcriptional activity of the *OsHKT1;1* promoter in the transient system (Fig. 7), the abolishment of *OsMYBc* in planta was expected to result in a low expression level of *OsHKT1;1*. Indeed, lower expression of *OsHKT1;1* was found in *osmybc* than in the wild type

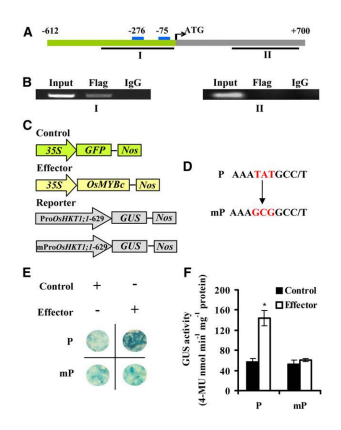


Figure 7. OsMYBc binds to the OsHKT1;1 promoter in vivo. A, Structure of the OsHKT1;1 gene. The green bar represents sequence upstream of the start codon, and the gray bar stands for coding regions of OsHKT1;1. The blue boxes indicate the positions of the two OsMYBc-binding elements. The lines below the binding elements and coding region indicate the fragments for ChIP-PCR in B. B, ChIP assay indicates that OsMYBc binds the OsHKT1;1 promoter in vivo. Fragmented chromatin DNA of rice protoplasts expressing the OsMYBc-Flag fusion protein was immunoprecipitated using anti-Flag antibody. Fragment I (-422 to +13) containing two cis-elements was amplified by PCR. Fragment II (+226 to +665) was used as a negative control. Input, Total input chromatin DNA; Flag, DNA precipitated using Flag antibody; IgG, DNA precipitated using mouse IgG. Each assay was repeated more than three times with independent biological materials. C and D, Schematic diagram of the effector and reporter used for transactivation studies. The plasmid 35S:OsMYBc was used as the effector, the plasmid ProOsHKT1;1-629:GUS (P) and its mutant version mProOsHKT1;1-629:GUS (mP) were used as the reporter, and 35S:GFP was used as an internal control. The sequences containing mutated nucleotides are shown in D. E, Transactivation activity was detected by GUS staining after reporter and effector plasmids were coinfiltrated into N. benthamiana. F, Quantitative analysis of the GUS activity indicated in E. Asterisks indicate that the mean value is significantly different from that of the control: *, P < 0.05. Error bars represent s_E (n = 3). 4-MU, 4-Methylumbelliferone.

in the presence or absence of salt treatment (Fig. 8G). Bioinformatic analysis indicated that OsMYBc also binds to at least three other *OsHKTs* and *HKTs* from wheat and maize, besides *OsHKT1;1* (Supplemental Table S4). We next determined the expression of *OsHKT1;5* and *OsHKT2;1* in the *osmybc* mutant background, which have been reported to be involved in Na⁺

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uptake and transport in rice (Ren et al., 2005; Horie et al., 2007). As shown in Figure 8H, NaCl-induced *OsHKT1;5* expression was pronouncedly reduced in the *osmybc* mutant, reminiscent of that of *OsHKT1;1* (Fig. 8G). The expression of *OsHKT2;1* was decreased in the *osmybc* mutant. Salt treatment (100 mM NaCl for 24 h) seriously inhibited *OsHKT2;1* expression, and its expression was lower in the *osmybc* mutant than in the wild type (Fig. 8I). These results suggest that OsMYBc regulates *OsHKT1;1* expression and probably the expression of other *OsHKTs*, playing an important role in salt tolerance.

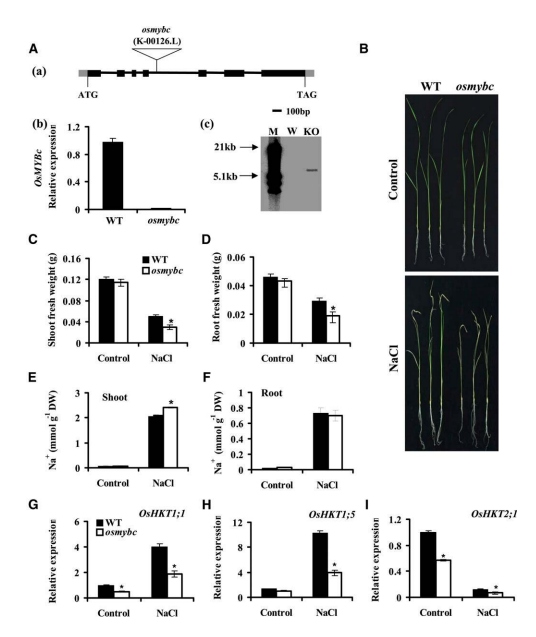
DISCUSSION

The main site of Na⁺ toxicity for many plants is the leaf, where photosynthesis and other metabolic processes occur (Munns and Tester, 2008). Some mechanisms of controlling Na⁺ accumulation in leaves have been identified and/or proposed. They include the control of Na⁺ uptake from soils, the reduction of the delivery of Na⁺ to the xylem, the storage of Na⁺ in the lower parts of the leaf (such as the sheath), and the recirculation of Na⁺ from shoots to roots (Davenport et al., 2005; Munns and Tester, 2008; Tian et al., 2010). In this article, we show the genetic evidence that OsHKT1;1, which is mainly in the phloem of leaf blades, is involved in limiting Na⁺ accumulation in leaves and salt tolerance in rice. Furthermore, this transporter is regulated by the *OsMYBc* transcription factor.

OsHKT1;1 Regulates Sodium Accumulation in Shoots

The phloem has been proposed to have a significant function in promoting $\bar{N}a^{+\bar{}}$ tolerance (Greenway and Munns, 1980), but only a few experimental reports have been published on the possible function of the phloem regulating Na⁺ concentration in leaves (Berthomieu et al., 2003; Tian et al., 2010). AtHKT1 was proposed to function as an Na⁺ transporter in the phloem (Berthomieu et al., 2003), but current studies indicated that it mainly works to remove Na⁺ from xylem sap (Sunarpi et al., 2005; Rus et al., 2006; Møller et al., 2009). Its orthologous gene in rice, OsHKT1;5, also mediates the retrieval of Na⁺ from the xylem (Ren et al., 2005). In situ hybridization experiments revealed that OsHKT1;1, OsHKT1;3, and OsHTK2;1 are expressed in the phloem (Jabnoune et al., 2009). OsHKT2;1 is involved in Na⁺ uptake as a mineral nutrient to enhance the growth of rice under K⁺ starvation conditions (Horie et al., 2007). In addition, knockout of the OsHKT2;1 gene did not affect growth under highsalt (140 mM NaCl) treatment compared with the wild type (data not shown). To our knowledge, there is no genetic evidence for OsHKT1;3's contribution to rice salt tolerance. To date, the significance of the recirculation of Na⁺ from shoots to roots has yet to be confirmed (Davenport et al., 2005; Munns and Tester, 2008).

The genetic analysis in this article revealed that the *oshkt1;1* mutant accumulated more Na⁺ in shoots,



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Figure 8. The *osmybc* mutant shows sensitivity to salt stress. A, Isolation of the T-DNA insertion *osmybc* mutant. a, Schematic diagram of the *osmybc* mutant. The boxes stand for exons, the lines stand for introns, and the triangle indicates the T-DNA insertion position. b, qRT-PCR analysis of *OsMYBc* expression in the wild type (WT) and *osmybc*. The cDNA was derived from whole plants. Error bars represent sE (n = 3). c, Southern-blot analysis of the T-DNA copy number in *osmybc* and wild-type plants. Genomic DNA was digested with restriction enzyme *Hin*dIII, and the *Hygromycin* gene was used as a probe. M, Marker; W, the wild type; KO, *osmybc* knockout mutant. B to D, Phenotypes under salt stress. Hydroponically grown seedlings were treated with 100 mM NaCl for 7 d, and *osmybc* showed more sensitivity to the salt stress than the wild type (cv Kitaake). B, Representative photographs of seedlings. C, Shoot fresh weight. D, Root fresh weight. The data represent means ± sE from three biological repeats, each consisting of 30 seedlings of each line. E and F, Na⁺ content in shoots (E) and roots (F). Error bars represent sE (n = 5). DW, Dry weight. G to I, Expression of *OsHKTs*. Plants were treated with or without 100 mM NaCl for 24 h. The genes *UBQ5* and *18S rRNA* were used as internal controls. Error bars represent sE (n = 3). Asterisks indicate that the mean value is significantly different from that of the wild type: *, P < 0.05.

especially in leaf blades, compared with the wild type (Fig. 1E; Supplemental Fig. S2). A failure in Na⁺ exclusion from shoots manifests its toxic effect and causes the premature death of older leaves (Munns and Tester, 2008; Fig. 1B). More accumulation of Na⁺ in shoots of

the *oshkt1;1* mutant could be owing to the disruption in the ability for Na⁺ exclusion from leaves, or in the control of Na⁺ transport from roots to leaves, or in both processes. Analysis of phloem sap revealed lower Na⁺ concentrations in the mutant than in the wild type (Fig. 2B), suggesting that OsHKT1;1 is potentially involved in the control of Na⁺ concentration in phloem sap. This speculation was partially supported by the observation that the bulk of *OsHKT1;1* expression was associated with the phloem of leaves (Jabnoune et al., 2009; Fig. 3A, c), although direct evidence for OsHKT1;1 functions in the phloem has yet to be presented.

OsHKT1;1 expression was also found in the vicinity of xylem tissues (Fig. 3A, c and f). After Na⁺ was transported to leaves, it was loaded to parenchyma cells and probably transferred to the phloem by OsHKT1;1, similar to the role of AtHKT1;1 in Arabidopsis (Sunarpi et al., 2005). The xylem-to-phloem transfer mechanism has been found for iron transport by the Oligopeptide transporter3 protein, which contributed to iron redistribution between the source and sink organs in Arabidopsis (Zhai et al., 2014). If the xylem-to-phloem transfer mechanism works for OsHKT1;1, it may function cooperatively between two types of cells. OsHKT1;1 was also expressed in xylem and phloem cells of roots, and its loss led to the increase of Na⁺ concentration in the xylem sap (Fig. 2C), suggesting that it may function in the control of Na⁺ transport to shoots.

The expression of OsHKT1;5 was also observed in roots and shoots. However, the induction of transcript levels of OsHKT1;5 by salt stress was found in the roots but not in shoots (Ren et al., 2005). This differs from that of OsHKT1;1, where salt-induced transcripts were observed in shoots but not in roots (Fig. 3). In wheat, an OsHKT1;5-like gene, TmHKT1;5-A (66% amino acid identity with OsHKT1;5), confers a root-specific constitutively active gene that is not induced by NaCl (Munns et al., 2012). Research using combined approaches has revealed that both OsHKT1;5 and TmHKT1;5-A are located on the plasma membrane of cells surrounding xylem vessels, withdrawing Na⁺ from the xylem and limiting the transport of Na⁺ to leaves (Ren et al., 2005; Munns et al., 2012). OsHKT1;4 was recently proposed to restrict leaf sheath-to-blade Na⁺ transfer in rice plants under salt stress (Cotsaftis et al., 2012). The capacity of the leaf sheath to sequester Na⁺ and thereby limit Na⁺ to the blade was also reported for salt tolerance in wheat (Davenport et al., 2005), although the candidate gene(s) is awaiting identification. oshkt1;1 showed a similar content of Na⁺ in leaf sheaths but a higher content in leaf blades, as compared with the wild type, suggesting its unique functions in the control of Na⁺ in shoots (Supplemental Fig. S2). Taken together, the above results suggest that different HKTs probably have distinct functions in plants. Further work is needed to decipher their mechanisms and cooperative functions in plants.

OsMYBc Regulates the Expression of *OsHKT1;1* to Affect Rice Salt Tolerance

Most membrane proteins are short-half-life proteins (Yen et al., 2008); therefore, they are expected to be affected largely by the rates of transcription and translation, coupled with their stability and function. The transcription factors that regulate the phosphate transporter and NH₄⁺ transporter genes were reported previously (Rubio et al., 2001; Zhou et al., 2008; Chiasson et al., 2014). For HKT regulation, a recent report indicates that the transcription factor ABI4 down-regulates the expression of *AtHKT1;1* in roots and affects salt tolerance (Shkolnik-Inbar et al., 2013).

We found in this study that transcriptional regulation of OsHKT1;1 affects the salt tolerance of rice. First, OsHKT1;1 expression is induced by salt stress (Fig. 3C), in accordance with the findings that HKT expression is regulated by salt, K⁺ starvation, or osmotic stress in other plants (Wang et al., 1998; Ren et al., 2005; Sunarpi et al., 2005; Horie et al., 2007; Shkolnik-Inbar et al., 2013). Second, OsHKT1;1 is regulated by a novel MYB coiled-coil transcription factor, OsMYBc. Genome-wide analysis led to the identification of 155 MYB genes in rice (Qu and Zhu, 2006). MYB-CC transcription factors constitute a family in plants (Rubio et al., 2001). These transcription factors have been reported to regulate transporter gene expression and vessel development. For instance, PHR transcription factors from Arabidopsis, rice, and maize regulate phosphate transporter gene expression (Rubio et al., 2001; Zhou et al., 2008; Wang et al., 2013). The APL1 gene, which encodes an MYB-CC transcription factor, controls phloem tissue identity in Arabidopsis (Bonke et al., 2003).

The isolated transcription factor *OsMYBc* in this study belongs to the MYB-CC family (Rubio et al., 2001; Fig. 5B). OsMYBc was up-regulated by salt (Fig. 5, D and E) and bound directly to the promoter of OsHKT1;1 (Figs. 6 and 7B). Scanning mutagenesis of the DNA target site in the OsHKT1;1 promoter for OsMYBc allowed us to identify the cis-element AAATATGCC/T. This element is similar to the previously reported PHR1binding consensus sequence GAATATGC at the upstream region of the PHR1-binding gene (AtIPS3; Rubio et al., 2001). The PHR1 homologous genes OsPHR1 and OsPHR2 have been identified in rice to function in the phosphate signaling pathway (Zhou et al., 2008), but its downstream target has not been found. In our study, we interpreted the interaction between OsMYBc and the OsHKT1;1 promoter using multiple approaches, showing that OsMYBc binding was essential for the promoter activity of OsHKT1;1 (Fig. 7, E and F). Knockout of the OsMYBc gene not only reduced the expression of OsHKT1;1 but also decreased salt tolerance (Fig. 8). These results support the importance of the OsMYBc regulation of OsHKT1;1 expression in salt tolerance. The MYB-binding element is also found in other OsHKTs and HKTs at least in wheat and maize (Supplemental Table S4), indicating that the identified cis-element is probably general among plant species.

In summary, we determined OsHKT1;1 function in the salt tolerance of rice using genetic and physiological approaches. Furthermore, the OsMYBc transcription factor was characterized to bind to the OsHKT1;1

MATERIALS AND METHODS

Plant Materials and Growth Conditions

Rice (*Oryza sativa japonica* 'Nipponbare') seeds were sterilized for 30 min with 2% (w/v) sodium hypochlorite solution and then washed three times with sterile distilled water. The seeds were soaked in water at room temperature for 3 d and then germinated for 1 d at 28°C. Seedlings were grown hydroponically in Yoshida's culture solution as described previously (Zhou et al., 2013). Plants were cultured in a growth chamber at 28°C/25°C (day/night) under a 14-hlight/10-h-dark photoperiod (approximately 500 μ mol m⁻² s⁻¹).

The *Tos17* insertion *oshkt1;1* mutant was obtained from the Rice *Tos17* Insertion Mutant Database (Hirochika, 1997; Yamazaki et al., 2001; Miyao et al., 2003). The T-DNA insertion mutant *osmybc* was obtained from the Crop Biotech Institute of Kyung Hee University. The homozygous lines were confirmed by a PCR-based method. Primers *OsHKT1;1-F*, *OsHKT1;1-R*, and *tail* 6 were used for the *oshkt1;1* mutant, and primers *OsMYBc-F*, *OsMYBc-R*, and the right bounder primer *2715RB* were used for the *osmybc* mutant. All primers are listed in Supplemental Table S5.

Nicotiana benthamiana seeds were germinated and grown in sterilized soil at $25^{\circ}C/20^{\circ}C$ (day/night) under a 16-h-light/8-h-dark photoperiod (approximately 100 μ mol m⁻² s⁻¹).

DNA Extraction and Southern-Blot Analysis

Genomic DNA was isolated and purified from 1 g of 21-d-old rice seedlings following the method described (Murray and Thompson, 1980). Southern-blot analyses were performed according to the method described Horbach et al. (2009). Ten micrograms of restriction enzyme-digested DNA was used for blotting. Specific primers *tos17ProbeF/tos17ProbeR* (Ding et al., 2007) and *HygProbeF/HygProbeR* were used to generate the digoxigenin-dUTP-labeled probe (Roche).

RNA Isolation and qRT-PCR

RNA was extracted using RNAiso Plus reagent (Takara) following the manufacturer's instructions. $\ddot{\mathrm{A}}$ total of 0.1 μg of RNA was used to synthesize first-strand cDNA using the PrimerScript RT Reagent Kit with gDNA Eraser (Takara). The primers OsHKT1;1-RT-F/OsHKT1;1-RT-R for OsHKT1;1, OsHKT1;5-RT-F/OsHKT1;5-RT-R for OsHKT1;5, OsHKT2;1-RT-F/OsHKT2;1-RT-R for OsHKT2;1, and OsMYBc-RT-F/OsMYBc-RT-R for OsMYBc were used for qRT-PCR. The quantified expression levels of the tested genes were normalized against the housekeeping genes, OsUBQ5 with primers OsUBQ5-F and OsUBQ5-R and 18S rRNA with primers 18S rRNA-F and 18S rRNA-R (Jain et al., 2006). qRT-PCR was performed using the SYBR Green master mix (Takara) on the ABI-7500 Fast Real-Time PCR System (Applied Biosystems). Conditions for quantitative analysis were as follows: 94°C for 2 min; 35 cycles of 94°C for 15 s, 60°C for 20 s, and 72°C for 30 s; and a final extension at 72°C for 10 min. Efficiency-adjusted gene expression was normalized with the geometric mean of the control primers using the following equation: Square root $(E_{\text{Ubq5}}^{Cq(\text{Ubq5})} \times E_{185}^{Cq(185)}) / E_{\text{Test}}^{Cq(\text{Test})}$ (Bartley et al., 2013), where E and Cq indicate the average reaction efficiency and cycle number at which the threshold fluorescence level is exceeded for the designated gene.

Vector Construction and Rice Plant Transformation

To complement the *oshkt1;1* mutant, a DNA fragment containing the *OsHKT1;1* promoter region 2,103 bp upstream of the initial codon and a 2,248bp region containing exons and introns was cloned from rice genomic DNA using primers *com-OsHKT1;1-F* and *com-OsHKT1;1-R* and linked into the modified vector pSuper1300. The resulting construct was introduced into the *Agrobacterium tumefaciens* strain EHA105 and transformed into an *oshkt1;1* mutant following the protocol reported by Nishimura et al. (2006).

The deletion derivatives of the OsHKT1;1 promoter were cloned from rice genomic DNA by PCR. The primers were Pro-OsHKT1;1-2120/Pro-OsHKT1;1-R,

Pro-OsHKT1;1-1623/Pro-OsHKT1;1-R, Pro-OsHKT1;1-1126/Pro-OsHKT1;1-R, and Pro-OsHKT1;1-629/Pro-OsHKT1;1-R for ProOsHKT1;1-2120, ProOsHKT1;1-1623, ProOsHKT1;1-629/Pro-OsHKT1;1-629, respectively. The promoter of RD29A was cloned from Arabidopsis (*Arabidopsis thaliana*) genomic DNA using the primers Pro-RD29A-F and Pro-RD29A-R. The products were transferred to a pCambia1301 vector containing a GUS gene.

The QuickChange site-directed mutagenesis kit (Stratagene) was used to generate the site-directed mutants. Using ProOsHKT1;1-629 as the template, the primers Point1-F/Point1-R and Point2-F/Point2-R were used for mutation.

To construct the GFP-OsMYBc, GST-OsMYBc, and Flag-OsMYBc fusion expression vectors, the full-length coding sequence of *OsMYBc* was cloned from rice cDNA using the primers *GFP-OsMYBc-F/GFP-OsMYBc-R*, *GST-OsMYBc-F/GST-OsMYBc-R*, and *Flag-OsMYBc-F/Flag-OsMYBc-R*, respectively. The PCR products were inserted into the vectors pSuper1300-GFP, pGEX-4T-1, and pFLAG, respectively.

Transient Expression in N. benthamiana Leaves

Transient expression was according to the method of Yang et al. (2000). Four-week-old *N. benthamiana* plants were used for infiltration. The constructs were individually transformed into the *A. tumefaciens* strain EHA105. The *A. tumefaciens* cells were infiltrated onto the abaxial surface of *N. benthamiana* leaves using 1-mL needleless syringes. After infiltration, *N. benthamiana* was grown in a greenhouse for 48 to 60 h.

GUS Staining and Activity Assay

Histochemical activity of GUS in transgenic plant materials and quantitative analysis of GUS activity in *N. benthamiana* leaves were detected according to the method of Jefferson et al. (1987).

Subcellular Localization of OsMYBc

The OsMYBc localization assay was performed as described by Campo et al. (2014). The GFP-OsMYBc fusion protein expression construct was transformed into onion (*Allium cepa*) epidermis cells using a gene gun (Bio-Rad). After bombardment, the onion layers were incubated in the dark for 20 h at 24°C. The cell layers were imaged with the LSM 780 Exciter confocal laser scanning microscope (Zeiss) with an excitation wavelength of 488 nm and a 505- to 530- nm band-pass emission filter.

Y1H Analysis

A Y1H library screen was performed using the Matchmaker Gold Yeast One-Hybrid Library Screening System kit and the Yeastmaker Transformation System 2 kit (Clontech), following the manufacturer's instructions. The OsHKT1;1 promoter region -612 to +17 (translation start is +1) was amplified by PCR using the primers Bait-F and Bait-R and inserted into the vector pAbA. The construct was linearized by BstBI digestion and transformed into a Y1HGold strain to generate a Y1H bait strain. Rice RNA, used to create a cDNA library, was extracted from the seedlings treated with 100 mm NaCl for 0, 1, 2, 4, 8, 24, and 48 h. A total of 5 µg of combined RNA at equal ratio was used to prepare the cDNA library. Screening of interaction clones was carried out via mating according to the manufacturer's instructions (Clontech). We screened approximately 1.75 million yeast (Saccharomyces cerevisiae) transformants and were able to isolate 26 potential positives that specifically interact with the bait protein. The plasmids of the 26 selected positives were extracted and amplified in Escherichia coli. All plasmids from E. coli were sequenced and BLASTed, and OsMYBc was selected for further study.

To confirm the interaction between the *OsHKT1;1* promoter (region -612 to +17) and OsMYBc, the full-length coding sequence of *OsMYBc* was cloned into the pGADT7AD vector using primers *pGADT7-OsMYBc-F* and *pGADT7-OsMYBc-R*. The *OsMYBc* construct or empty vector was transformed into the Y1H bait strain and selected on a synthetic dropout-Leu plate containing 250 ng mL⁻¹ AbA.

EMSA

EMSA was performed according to the method of Wang (2012). The GST-OsMYBc fusion protein was expressed in *E. coli* BL21 at 23° C for 8 h in the

presence of 0.5 mM isopropyl β -D-1-thiogalactopyranoside. GST-OSMYBc protein was purified using Glutathione Sepharose (Genscript Life Sciences) beads according to the manufacturer's instructions.

OsHKT1;1 promoter fragments F1 (-612 to -413), F2 (-412 to -183), and F3 (-182 to +17) were amplified by PCR using the primers *F1-L/F1-R*, *F2-L/F2-R*, and *F3-L/F3-R*. Five regions, each containing 40 bp in fragment F3, were synthesized by Shanghai Sangon Biotech. The digoxigenin gel-shift kit, second generation (Roche), was used to perform EMSA following the manufacturer's instructions. Two nanograms of digoxigenin-labeled DNA probes was incubated with 4 μ g of purified recombinant proteins (OSMYBc) in a total volume of 20 μ L. The reaction mixtures were incubated at room temperature for 15 min and loaded onto a 6% (w/v) native polyacrylamide gel. Electrophoresis was conducted at 80 V for 3 h in 0.5× TBE buffer (44.5 mM Tris, 44.5 mM boric acid, and 1 mM EDTA, pH 8) at 4°C. The gel was sandwiched and transferred to an N⁺ nylon membrane (Roche) in 0.5× TBE buffer at 400 mA for 30 min at 4°C. DNA labeled by chemiluminescence was exposed and detected using x-ray film.

ChIP

ChIP was performed as described by Lee et al. (2007) using the EpiQuik Plant ChIP Kit (Merck Millipore). In brief, the FLAG-OsMYBc fusion protein expression vector was transformed into rice protoplasts according to the protocols described previously (Ge et al., 2012). Protoplasts were fixed with 17% (v/v) formaldehyde, and chromatin was isolated and sheared by sonication to obtain fragments of sizes between 200 and 1,500 bp. Anti-Flag monoclonal antibodies bound to protein G-coated beads were used to immunoprecipitate the genomic DNA fragments. PCR was performed with immunoprecipitated genomic DNA using primers *ChIP-IF-JChIP-IR* and *ChIP-II-F/ChIP-II-R*. ChIP experiments were performed independently three times.

Physiological Analysis for Salt Tolerance

Three-week-old hydroponically grown seedlings were treated with 100 mM NaCl for 7 d. The growth phenotypes were recorded by taking photographs. Fresh weight, shoot length, and total chlorophyll content were determined. The survival rate of seedlings was analyzed according to the method of Zhang et al. (2009). Briefly, the seedlings after salt treatment were transferred to culture solution without NaCl for recovery. After the seedlings were grown for an additional 1 week, the plants with aerial parts completely yellow were scored as dead.

Determination of Na⁺ Content

Shoots and roots of rice seedlings were harvested separately, and Na⁺ content was determined according to the methods described by Ali et al. (2012) with minor modifications. Briefly, samples were dried at 80°C for 2 d. Each dry sample was digested in 5 mL of nitric acid at 90°C for 8 h, diluted to 25 mL with distilled water, and analyzed using an inductively coupled plasma-optical emission spectrometry instrument (Pekin Elmer).

Na⁺ concentration in the phloem sap was determined according to the method of Ren et al. (2005). Rice plants were grown hydroponically in culture solution for 21 d. The phloem sap was collected after NaCl treatment (15 mm) for 2 d. For each replicate, four plants were detached using a blade, and the wound sections were dipped in a solution of 20 mM K2EDTA, pH 7.5, for at least 1 min. The shoots were then dipped in 1 mL of 15 mM K2EDTA, pH 7.5, for 4 h in an illuminated growth room under a water-saturated atmosphere. The quantity of Na⁺ was measured by inductively coupled plasma-optical emission spectrometry as described previously. The volume of the collected phloem sieve is highly variable and cannot be measured directly because of technical reasons (Berthomieu et al., 2003; Ren et al., 2005), and Gln is usually used as an internal standard because it is abundant in the phloem sap and remains quite constant during the 24-h day/night cycle (Berthomieu et al., 2003). The Gln released in the EDTA solution was analyzed using an amino acid analyzer (L8900; Hitachi). The Na+ concentration in the phloem sap is expressed as Na⁺-Gln ratio.

We measured Na⁺ concentration in xylem sap according to the method of Horie et al. (2007) with minor modifications. Rice plants were grown as for collection of the phloem sap. The shoots (2 cm above the roots) were excised using a blade. The xylem sap was collected, except for the first drop, using a micropipette for 2 h after decapitation of the shoot. Total Na⁺ content in the sap was measured.

Chlorophyll Content Measurement

Estimation of the total chlorophyll content was performed according to Porra et al. (1989). Rice seedlings were incubated in 80% (w/v) acetone, and after vigorous shaking in the dark for 24 h at room temperature, and then centrifugation for 10 min at 10,000g, the supernatant was collected for chlorophyll determination.

Statistical Analysis

Statistical analysis was performed using SPSS 3.0, and differences were analyzed with one-way ANOVA followed by Tukey's multiple comparison test.

Sequence data from this article can be found in the Rice Annotation Project Database under the following accession numbers: *OsHKT1;1*, LOC_Os04g51820; *OsMYBc*, LOC_Os09g12770.

Supplemental Data

The following supplemental materials are available.

- Supplemental Figure S1. Southern-blot analysis for the plant materials used in this study.
- Supplemental Figure S2. Na⁺ content in the leaf sheaths and leaf blades.
- Supplemental Figure S3. OsHKT1;1 localization in the cell.
- Supplemental Figure S4. OsMYBc binds to the promoter of OsHKT2;1.
- **Supplemental Table S1.** Raw data of Na⁺ content in the leaf sheaths and leaf blades.
- Supplemental Table S2. Raw data of Na⁺ content in phloem sap.
- Supplemental Table S3. Positive interactions from the Y1H screen.
- **Supplemental Table S4.** *HKT* genes containing the MYB-CC cis-elementbinding region in their promoters.
- Supplemental Table S5. Primers used in this study.
- Supplemental Methods S1. Subcellular localization of OsHKT1;1.

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