Identification of an Apoplastic Protein Involved in the Initial Phase of Salt Stress Response in Rice Root by Two-Dimensional Electrophoresis1[C][W][OA]

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Salt stress is one of the most significant abiotic stresses and affects every aspect of plant physiology and metabolism. During salt stress, Na⁺ enters the cells and accumulates to a concentration that induces ionic and osmotic stress in plants. Plant cells respond and adapt to these adverse conditions through signaling networks (Lee et al., 2004). Understanding the signaling pathway of plant salt resistance is important for improving plant salt tolerance, especially for improving agricultural productivity in irrigated land.

To study the signaling network of plant salt adaptation, the most important thing is identification of the components involved. Much effort has been made to discover components or elements of signaling pathways involved in plant salt stress responses. Of all the molecular components and signaling pathways known so far, the best understood signaling pathway in salt ionic stress is the SOS pathway (Zhu, 2002). In addition to the SOS pathway, other signaling pathways and components have also been suggested to be involved in salt osmotic stress signaling. Several plant protein kinases have been found to be activated by osmotic stress (Liu et al., 2000; Zhu, 2002). Osmotic stress can increase the transcription levels of a number of protein kinase genes, including genes for a two-component His kinase, mitogen-activated protein kinase kinase, mitogen-activated protein kinase, mitogen-activated protein kinase kinase, and mitogen-activated protein kinase (Munnik et al., 1999; Mikolajczyk et al., 2000; Mizoguchi et al., 2000). Also, membrane phospholipids make up a dynamic system that generates a multitude of signaling molecules in addition to serving important structural roles during stress responses (Munnik et al., 2000; DeWald et al., 2001; Zhu, 2002). Recently, a new Ca²⁺-dependent membrane-binding protein (AnnAt1) involved in salt stress was identified by a proteomic approach (Lee et al., 2004). To date, most identified salt stress resistance-related signaling components have been localized to the cytosol or cell membrane. Although it is an important component of the plant cell, the plant apoplast has been ignored in studies of the salt stress response.

The apoplast is the portion of the plant cell outside the cell membrane. This region includes the cell walls and intercellular space of the plant (Dietz, 1997). The apoplast, which plays important roles in regulating plant physiological and developmental processes, is not only a barrier but also a linker between the environment and the protoplast; there are many organic and inorganic molecules, as well as enzymes, present in the plant apoplast. These molecules have...
An Apoplastic Protein Involved in Salt Stress Response

Table 1. MDH enzyme assay

Results shown present data collected from one independent experiment. Of all the MDH activities assayed in three independent experiments, the results were similar. One unit (U) was calculated using the change of 0.01 A\text{\textsubscript{410}} per minute. ND, Not detected.

<table>
<thead>
<tr>
<th>Extract</th>
<th>Protein Concentration (mg mL\textsuperscript{-1})</th>
<th>Enzyme Activities (U ml\textsuperscript{-1})</th>
<th>Relative Enzyme Activities (U mg\textsuperscript{-1})</th>
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<tr>
<td>Rice root total cell extracts</td>
<td>0.65</td>
<td>180 ± 28</td>
<td>277 ± 43</td>
</tr>
<tr>
<td>Diluted rice root total cell extracts</td>
<td>0.065</td>
<td>38 ± 7</td>
<td>585 ± 108</td>
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<td>Rice root apoplastic extracts</td>
<td>0.06</td>
<td>ND</td>
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The sequencing of many plant and animal genomes has revealed large numbers of putative receptor-like kinase genes. The potential ligands of many of these putative receptors have been predicted to be peptides. It is very likely that some of these receptor-like protein kinases may be involved in plant perception of, and response to, biotic and/or abiotic stress signals (Haffani et al., 2004). These data suggest that there may be a number of proteins in the plant apoplast that are involved in the plant stress response. The identification of these proteins during the initial phase of salt stress will be an important step toward understanding the role of these apoplastic proteins in the salt stress response.

The sequencing of many plant and animal genomes has revealed the fact that the regulation of a cell’s biological activities mostly occurs at the level of protein degradation, interactions, and posttranslational modification. Proteomics is now emerging as a powerful tool for studying these protein dynamics, especially in plant stress responses (Salekdeh et al., 2002; Lee et al., 2004). A number of studies have been performed using proteomic approaches to identify pathogen response proteins in rice and Arabidopsis (Kim et al., 2003; Ndima et al., 2003) and salt stress response proteins in the rice microsome (Lee et al., 2004), root (Yan et al., 2005), and leaf sheath (Abbasi and Komatsu, 2004). Dani et al. (2005) used a proteomic approach to analyze the proteome changes in the tobacco (Nicotiana tabacum) leaf apoplast during long-term (20 d) salt stress. Nevertheless, few studies have addressed the changes in the apoplastic proteome in response to salt stress, especially during the initial phase of salt stress. In this study, the changes in the rice root apoplastic proteins during the initial phase of salt stress were carefully investigated by two-dimensional electrophoresis (2-DE). Eight salt stress-regulated apoplastic proteins were identified, including a secreted protein with extracellular domain-like Cys-rich motifs (DUF26) that has been reported to be involved in O. sativa root meander curling (OsRMC; Jiang et al., 2007). In our work, the roles of OsRMC in the response to salt stress have been revealed. OsRMC was up-regulated at the transcriptional and translational levels during the initial phase of salt stress. Our findings showed that knocking down the expression level of OsRMC resulted in more resistance to salt stress in transgenic rice.

RESULTS

Apoplastic Extracts Are Free of Cytosolic Contamination

The vacuum infiltration method was used to obtain apoplastic fluid from the rice root. Because the procedures of vacuum infiltration and centrifugation could damage cells and contaminate apoplastic extracts with
cytosolic ingredients, we used malate dehydrogenase (MDH) activity assay and immunobloting against tubulin to evaluate the quality of apoplastic extracts. Our results show that MDH activity could not be detected in our apoplast extract (Table I). To eliminate the possibility that the difference in MDH activity between the total cell extracts sample and the apoplastic extracts sample was due to the protein concentration difference, the total cell extract was diluted to a concentration that was similar to that of the apoplastic extracts and analyzed again for MDH activity (Table I); the normalized MDH activity in total cell extracts was even higher than in the undiluted samples. In addition, apoplastic extracts from three independent extraction procedures and soluble cell total extracts were separated and probed with the anti-tubulin monoclonal antibody. With an apparent molecular mass of 49 kD, tubulin was detectable in whole cell extracts but not in any of our apoplastic extracts (Fig. 1). These results proved that the apoplast extract samples used in this study were free of cytosolic contamination.

Figure 2. Protein expression profiles of the rice root apoplast after treatment for 1 h (C1 and T1), 3 h (C3 and T3), and 6 h (C6 and T6) with 200 ms NaCl (T1, T2, and T3) or regular medium (C1, C2, and C3). Arrows indicate the MALDI-TOF MS-identified protein spots whose abundance increased or decreased consistently following NaCl treatments.

The Apoplastic Proteins Changed in Response to Salt Stress

Rice root apoplast proteins from the 1-, 3-, and 6-h salt treatments and from the controls were analyzed by 2-DE. Approximately 100 spots from each sample were visualized by Coomassie Brilliant Blue staining (Fig. 2). Three independent experiments were conducted to ensure that the protein abundance changes at each time point were reproducible and significant. Software quantification showed that, although the protein expression profiles between the salt-stressed and untreated samples were similar to each other, the abundance of some spots changed significantly with salt stress treatment over time (Figs. 2 and 3). We found 10 spots that showed at least a 1.5-fold increase or decrease in abundance (P < 0.05) in response to salt stress (Figs. 2 and 3). Based on their expression profiles, these proteins were classified into three groups. The first group contains spots 1, 2, 5, 6, 7, 8, and 10, which showed a steady increase in abundance over the...
course of the treatment (Fig. 3). The second group contains spot 9 only, which showed a steady decrease in abundance across all treatment time points (Fig. 3). The third group contains spots 3 and 4. The abundance of the spots in this group increased at 3 h and decreased at 6 h of salt treatment (Fig. 3). The different expression patterns of the spots might imply different roles for these apoplastic proteins in plant salt stress responses.

Identification of Salt Stress-Responsive Proteins by Mass Spectrometry

All 10 salt stress-responsive spots were subjected to in-gel digestion and analyzed by matrix-assisted laser-desorption ionization time of flight mass spectrometry (MALDI-TOF MS). The obtained peptide mass fingerprints were used to search the National Center for Biotechnology Information database using Mascot (http://www.matrixscience.com). The 10 identified proteins spots are listed in Supplemental Table S1. The experimentally calculated molecular mass and pI values of identified protein spots were consistent with the theoretical values, except for spots 1 and 9 (Supplemental Table S1).

To confirm that the proteins we identified were indeed apoplastic proteins, all identified protein sequences were searched using the TargetP program (www.cbs.dtu.dk/services/TargetP) for their predicted subcellular localization (Emanuelsson et al., 2000). As we can see from Supplemental Table S1, eight of 10 spots were predicted to be typical secretory proteins with signal peptide sequences.

OsRMC Is an Apoplastic Protein

Both spots 1 and 6 were identified as protein OsRMC (gi|19387274). To confirm the identities of spots 1 and 6, MS/MS was used for further analysis (Supplemental Figs. S1 and S2). The results showed that spots 1 and 6 represent the same protein, and the pI/molecular mass difference between the two spots may be due to posttranslational modification or dimerization. This protein has been further analyzed using bioinformatics tools. OsRMC belongs to the secreted type of DUF26 protein (Chen, 2001), with two extracellular domain-like Cys-rich motifs (DUF26). A sequence including 23 residues in the N terminus was predicted to be the signal peptide (http://www.cbs.dtu.dk/services/SignalP/; Nielsen et al., 1997; Moller et al., 2001; Supplemental Fig. S3).

As OsRMC was obtained from an apoplastic sample and bioinformatics analysis also indicated that OsRMC is a putative secreted protein, we sought experimental evidence for its apoplastic localization. We used Citrine reporter gene, which encodes a new yellow fluorescent protein (YFP) that has lower pH sensitivity (Griesbeck et al., 2001), to determine the subcellular localization of OsRMC in vivo. The recombinant constructs of the 35S::OsRMC-Citrine fusion gene were transiently expressed in onion (Allium cepa) epidermal cells by particle bombardment. Citrine expressed from the empty vector pAVA321 was used as a negative control. As shown in Figure 4, in the onion epidermal cell, OsRMC-Citrine was detected in the cell wall region after plasmolysis with 0.9 M mannitol, whereas for the empty vector control, the Citrine was not detected in the extracellular region but only throughout the intracellular region. This result confirms that OsRMC is an apoplastic protein.

Figure 3. Histogram showing fold changes in abundance (following 200 mM NaCl treatment for 1, 3, and 6 h) for each of the protein spots in three independent experiments. Values plotted are mean values with s. The significance of the differences between treated and untreated groups is indicated by asterisks: * P < 0.05, ** P < 0.01.
locally in the callus and was high in the roots and leaves of transgenic seedlings (Fig. 5A, a–c). In the mature rice plants, the stems and nodes, except the flag leaf, were stained blue (Fig. 5A, d and e). In the inflorescence, we could see GUS expression in the palea, lodicule, stamen, pistil, and especially in the anther (Fig. 5A, f and g). The seeds of transgenic rice were also stained (Fig. 5A, h and i). Expression of GUS was under the control of the OsRMC promoter in the transgenic plant, so the transcriptional induction of the OsRMC gene in response to salt treatment was determined by monitoring GUS activity. The GUS activity of OsRMCpro::GUS plants in response to salt stress was measured over a 24-h period. Each data point is an average of three independent experiments. Treating the plants with 200 mM NaCl resulted in rapidly and continuously increased GUS activity (Fig. 5C).

To further confirm the protein level change of OsRMC in response to salt stress, we generated polyclonal antibodies against OsRMC. A western blot was used to confirm the induction of OsRMC at the translational level by salt stress. The results showed that the level of OsRMC protein increased when plants were treated with 200 mM NaCl for 1 h and kept increasing after 12 h under NaCl treatment (Fig. 5D). These results indicate that OsRMC is a salt stress-responsive gene and that salt stress promotes the expression of OsRMC at the transcriptional and the translational levels.

Knockdown of OsRMC Leads to Improved Salt Tolerance of Transgenic Plants

To assess the function of OsRMC in the salt stress response in vivo, an RNA interference (RNAi) construct (pTCK303) containing part of the coding sequence fragment (428 bp) to knock down OsRMC expression and an overexpression construct (pCAMBIA1300) containing the OsRMC full-length ORF (without the stop codon) fused to a GUS reporter gene driven by a ubiquitin promoter (Ubi-1pro::OsRMC-Gus) were introduced into rice by Agrobacterium tumefaciens EHA105-mediated transformation. We examined the T-DNA insertions of select transgenic lines by Southern blot (Fig. 6A). Two independent RNAi homozygous lines (R1 and R12) and an overexpression homozygous line (OS1i) were used for further analysis. Real-time Q-PCR indicated the OsRMC mRNA expression levels in the
RNAi lines were significantly knocked down. However, gene silencing of OsRMC also occurred in the overexpression line OXSi, resulting in a low OsRMC expression level, which was similar to that in the RNAi lines (Fig. 6B). To confirm the protein expression level for the RNAi lines, a western-blot assay was performed using an anti-OsRMC antibody. The results of the western blots matched those of the real-time Q-PCR: the protein expression level of OsRMC in the knock-down lines, including the overexpression silenced line, OXSi, was obviously reduced (Fig. 6C).

To analyze the functions of OsRMC under salinity stress, we tested the effect of NaCl on seed germinations of OsRMC knockdown lines. There were no differences in seed germination rates between the transgenic lines and the wild type under normal condition. We tried gradients of NaCl concentrations from 100 to 200 mM to treat seeds for 7 d and found that the seed germination rates were higher than that of the wild type, and in 175 mM NaCl treatment the difference was most distinct (Fig. 7, A and B). In 175 mM NaCl treatment, the seeds of Ri1, Ri2, and OXSi were soaked in NaCl solution on a plate containing a filter for 9 d. Seeds were scored for germination from the 4th d, based on whether or not the shoot length exceeded half of the seed length. In statistical analysis, the germination rates of the Ri1, Ri2, and OXSi transgenic lines were higher than that of the wild type in the presence of 175 mM NaCl (Fig. 7C).

We also put the seeds of knockdown lines Ri1 and OXSi in half-strength Murashige and Skoog (1/2MS) medium containing 50 to 175 mM NaCl or without NaCl as a control to germinate and grow into seedlings. Ten-day-old seedlings were measured for shoot length, fresh weight, and dry weight of the shoot. The knockdown transgenic rice seedlings exhibited no difference in growth compared with wild-type rice seedlings under normal conditions, whereas in the presence of NaCl, the knockdown lines showed inhibited growth. In the 100 mM NaCl treatment, the difference was most distinct (Fig. 8A). Statistical analysis revealed that the knockdown plants exhibited distinct reduction in shoot length, fresh weight, and dry weight compared with control, wild-type plants (Fig. 8, B–D). In 175 mM NaCl, both the germinated seedlings of knockdown plants and control plants exhibited growth reduction with no apparent difference. The Ri2 knockdown line, which has a relatively higher expression level of OsRMC than Ri1 and OXSi (Fig. 6C), was also used for the growth assay in salt treatments and showed a similar phenotype of growth inhibition (data not shown).

Growth rate reduction in the OsRMC-silenced plants could reflect either an adaptive response or an injury response of the plants under the stress (Zhu, 2001). In order to delineate these possibilities, we measured the seedling survival rates of OsRMC knockdown lines under salt stress conditions. For the survival assay, 10-d-old knockdown seedlings cultured in Hoagland liquid medium were transferred to Hoagland medium with gradient NaCl concentrations from 100 to 200 mM for 4 d and then allowed to recover under normal conditions for 7 d, after which the survival rate was calculated. Plants that could not grow any more after the recovery period were considered to be dead. After recovering for 1 week, more transgenic seedlings survived than wild-type plants, which appeared mostly withered (Fig. 9A). The sur-
vival rates of Ri1 and OXSi were higher than that of the wild type after 7 d of recovery from 150 and 200 mM salt stress, whereas in recovery after 100 mM NaCl treatment both the knockdown lines and the wild type all survived (Fig. 9B). We measured the lipid peroxidation level in rice seedlings under salt stress, and the end product MDA contents of leaves were determined by thiobarbituric acid test (Heath and Parker, 1968). As shown in Figure 9C, MDA levels were elevated in both wild-type and transgenic plants within gradient NaCl treatments, but the degree of lipid peroxidation was significantly lower in knockdown plants than in wild-type plants, which means that the transgenic plants are more tolerant to stress injury.

The expression levels of two salt-responsive genes, OsDREB2A and Rab16A, also were monitored by real-time Q-PCR analysis. As shown in Figure 9D, both OsDREB2A and Rab16A showed higher expression in knockdown lines than in the wild type under normal conditions. When treated with 200 mM NaCl for 6 h, significant inductions of these two salt-responsive genes happened in all salt-treated plants, with higher expression levels in transgenic plants than in wild-type plants. Thus, knockdown of OsRMC results in up-regulated expression of OsDREB2A and Rab16A, which are involved in plant tolerance to salt stress.

DISCUSSION

To understand the signaling network underlying plant salt adaptation, a two-dimensional electrophoresis approach was used to identify the apoplastic components involved in this process. Despite the difficulties encountered in extracting and identifying proteins, apoplast proteomics has become an active field in recent years (Jamet et al., 2006, 2008). For a differential display analysis of apoplast proteins responding to salt stress, the quality of the apoplast-specific extract is crucial. There are several methods commonly used to extract the apoplast solution from the plant, including the application of pressure using a Scholander bomb, the vacuum perfusion method, the elution method, and the vacuum infiltration method (Lohaus et al., 2001; Jamet et al., 2008). Of all these methods, the vacuum infiltration method has been most widely used in obtaining apoplastic fluid from different plant species (Husted and Schjoerring, 1995; Lohaus et al., 2001; Dani et al., 2005). To avoid the cytoplasmic contamination of apoplast extracts, an MDH assay and a western-blot assay were used here to evaluate cytoplasmic contamination. MDH activity was detected in total protein extract controls but not in apoplastic extracts (Table I). The immunoblotting against tubulin indicated that tubulin was not present in apoplastic extracts but was present in total protein extracts (Fig. 1). Furthermore, all identified protein sequences were analyzed using the TargetP program (www.cbs.dtu.dk/services/TargetP) to predict their subcellular localization (Emanuelsson et al., 2000), since the sensitivity of mass spectrometry is 100 to 1,000 times greater than that of the biochemical assay (Jamet et al., 2006). As we can see from Supplemental Table S1, eight of 10 spots were predicted to be typical secretory proteins with a signal peptide sequence. These data indicate that our preparations of apoplastic proteins from the rice root did not show detectable intracellular contamination and therefore were appropriate for use in proteomic analysis.
Based on the 2-DE analysis, it was observed that the apoplastic proteins changed in response to salt stress during the initial phase. We found that protein expression pattern changes within 1 h (Figs. 2 and 3), which is quite a swift response. Previous studies had indicated that salt, drought, and heavy metals alter the apoplastic protein composition of leaves (Fuhrer, 1982; Brune et al., 1994; Dietz, 1997; Dani et al., 2005). According to the previous [35S]Met incorporation experiment (Dietz, 1997), apoplast protein was slightly changed after 1 h and continued to increase after 10 h of treatment with nickel. Previous work had also indicated that mRNA levels were altered in response to salt stress within 15 min in rice (Kawasaki et al., 2001). These results indicated that plants can rapidly respond and adapt to salt stress.

Ten significantly salt stress-responsive spots were subjected to in-gel digestion and analyzed by MALDI-TOF MS, and six kinds of protein were identified: OsRMC, a peroxidase, a glucanase, rab5B, a thioredoxin, and a putative pathogenesis-related protein. Based on their potential physiological functions, the salt stress-regulated apoplastic proteins identified in this study are all involved in the plant stress response.

In plants, β-1,3-glucanase belongs to the same family of pathogenesis-related proteins as PR-2 and is strongly induced when plants respond to wounding or infection by fungal, bacterial, or viral pathogens (Trudel et al., 1998). Plant peroxidases are also well-studied PR proteins belonging to the PR-9 family (Swapan and Muthukrishnan, 1999). During the pathogenesis resistance response, apoplastic peroxides polymerize pro-
teins and lignin or suberin precursors into plant cell walls (Young et al., 1995). This leads to the formation of a physical barrier that prevents the pathogen from penetrating the cell walls and spreading. Prb1 (a putative pathogenesis-related protein) contains a conserved domain named SCP, which is often seen in plant pathogenesis-related proteins (Marchler-Bauer et al., 2007) such as PR-1, a homolog of Prb1, and a salicylic acid (SA)-response marker gene widely used in SA and jasmonic acid (JA) response studies (Koornneef and Pieterse, 2008). Rab5B belongs to the Rab subfamily of the small G-protein superfamily. Rab GTPases are implicated in vesicle trafficking and are involved in multiple physiological processes in plants (Ueda et al., 2001; Preuss et al., 2004). In Mesembryanthemum crystallinum, a protein belonging to the plant rab5 family was regulated at the transcriptional level by early salt stress (Bolte et al., 2000). Also, thioredoxin reductase is part of the plant redox-regulating system, and there is evidence showing that thioredoxin participates in a number of physiological processes in plants, including abiotic and biotic stress responses, seed germination, self-incompatibility, cell division, and translation (Besse and Buchanan, 1997; Vieira Dos Santos and Rey, 2006).

It was interesting that spots 1 and 6 were both identified as an OsRMC, containing a predicted N-terminal signal peptide and two DUF26 domains (Supplemental Fig. S3C). Proteins with the DUF26 domain belong to a large protein family with many members in higher plants. All of these share similarities in the DUF26 domain (C-X8-C-X2-C, a Cys-rich repeat motif). These conserved Cys residues may function to maintain the three-dimensional structure and form a zinc finger motif to mediate protein-protein interactions or sense the redox changes in the extracellular space during plant defense responses (Chen, 2001). The majority of the DUF26 protein family consists of two types. One type is receptor-like kinases with the extracellular domain, the transmembrane domain, and the intracellular kinase domain, whereas the other type is secretory proteins, which have only the extracellular domain with Cys-rich motifs (DUF26). In the previous study of Jiang et al. (2007), the subcellular localization of an OsRMC-GFP fusion protein was analyzed in the epidermal cells of hypo-
cotyls of transgenic Arabidopsis, and the results showed that the fusion protein was localized in the plasma membrane. As to our work, we have generated an OsRMC-Citrine fusion protein that was transiently expressed in onion epidermal cells via particle bombardment. Confocal microscopy analysis indicated that the OsRMC-Citrine fusion protein was localized not only in the plasma membrane region but also in the extracellular cell wall region of onion epidermal cells after plasmolysis (Fig. 4). This result confirms that OsRMC, as a member of the secretory DUF26 proteins with only an extracellular domain and without a transmembrane domain or a kinase domain, is a secreted apoplastic protein.

The sequences of various cis-acting elements, including ABREs, DREs, LTREs, MYBRS, and MYCRS, were predicted in the 1,777-bp promoter region of OsRMC (Supplemental Fig. S3D). These cis elements and their respective transcription factors have important roles in abiotic stress responses (Nakashima and Yamaguchi-Shinozaki, 2006). We performed real-time Q-PCR, relative GUS activity analysis of OsRMC promoter::GUS transgenic rice plants, and western blotting and detected the up-regulated expression profile of OsRMC in response to salt stress (Fig. 5, B–D). These results suggested that OsRMC has some functions in the response to salt stress.

In the salt stress experiments with transgenic plants, knocking down OsRMC led to growth inhibition upon 50, 100, 150, and 175 mM NaCl treatment for 10 d (Fig. 8, A–D). Although the knockdown lines show higher germination rates than the wild-type control in 175 mM NaCl treatment, in the germinated seedlings there was no apparent difference in growth reduction between the knockdown plants and control plants, both seedlings showing severe growth inhibition in such salt conditions. This result is consistent with the previous work of Jiang et al. (2007), in which knocking down OsRMC enhanced the response of transgenic rice plants to JA treatments and led to the growth inhibition of shoots and root coiling. Growth inhibition is commonly observed in transgenic plants overexpressing stress-related transcription factors that have been reported to improve tolerance to stress conditions (Ciftci-Yilmaz et al., 2007; Nakashima et al., 2007). In the plants under stress conditions, the transcript levels of certain photosynthesis-related genes and genes for carbohydrate metabolism have been found to be reduced (Sakamoto et al., 2004). Down-regulated expression of these proteins may lead to a suitable energy balance for plants under stress conditions. As a result, plant growth is suppressed and stress tolerance is increased. In the salt tolerance test assay, knockdown lines showed improved salt tolerance, as manifested in higher seed germination rates (Fig. 7, A–C), and higher seedling survival rates (Fig. 9, A and B) than the wild type.

Under salt stress conditions, the MDA contents in leaves of transgenic rice seedlings were lower than in the wild type, which indicates that the degree of lipid peroxidation was significantly reduced in transgenic plants when treated with salt stress (Fig. 9C). Meanwhile, the rice salt response genes OsDREB2A and Rab16A were up-regulated in the OsRMC knockdown transgenic plants (Fig. 9D). Expression of OsDREB2A was induced by dehydration and high-salt stresses (Dubouzet et al., 2003). Overexpression of constitutively active Arabidopsis DREB2A or maize (Zea mays) ZmDREB2A resulted in enhanced tolerance to drought stress in transgenic Arabidopsis plants (Sakuma et al., 2006; Qin et al., 2007). Rab16A belongs to the group 2 lea gene family, which was distinctly induced by salt and abscisic acid treatment (Mundy and Chua, 1988). Transgenic tobacco that overexpressed Rab16A displayed increased salt tolerance (RoyChoudhury et al., 2007). These results indicate that OsRMC negatively regulates the salt tolerance of rice plants and may be involved in the injury response of plants (Zhu, 2002; Koornneef and Pieterse, 2008).

Interestingly, it was found that the OsRMC protein was also regulated by pathogen inoculation (Kim et al., 2003) and the wound response (Shen et al., 2003), except for JA and salt stress in rice. Similarly, other DUF26 receptor protein kinases have been reported to be involved in pathogen infection, wound responses, and responses to reactive oxygen species and SA in Arabidopsis (Chen et al., 2003, 2004; Acharya et al., 2007) and rice (Nakashima et al., 2007). Therefore, the DUf26 protein family may play a broad-spectrum role in regulating the perception of injury caused by biotic and abiotic stress signals.

MATERIALS AND METHODS

Plant Material and Salt Stress Treatments

Rice (Oryza sativa L. japonica ‘Nipponbare’) seedlings were grown in the greenhouse at 28°C/25°C, 16 h of light/8 h of dark, and 50% humidity (Kawasaki et al., 2001). Ten-day-old rice plants were used for salt stress treatments by 200 mM NaCl solution and treated for 1, 3, and 6 h. Roots were harvested immediately for apoplastic protein extraction. For germination assays, seeds were soaked in NaCl solution and scored for germination based on whether or not the shoot length exceeded half of the seed length. For growth assay, plants were grown in 1/2MS solid medium containing NaCl for 10 d and used for growth measurements (Hu et al., 2006). For survival experiments, 10-d-old rice plants were treated with NaCl solution for 4 d and allowed to recover for 1 week. Seedlings that could not grow were considered dead (Xiong and Yang, 2003). All of these salt tolerance experiments were repeated at least three times.

Rice Transformation

Rice seeds were sterilized and cultured on Murashige and Skoog plus (4 mg L⁻¹ 2,4-dichlorophenoxyacetic acid) medium for 4 weeks in the dark at 28°C to induce embryogenic callus for Agrobacterium tumefaciens-mediated transformation, as described by Yang et al. (2004). The positively transformed calli were selected by hygromycin B (50 mg L⁻¹) and differentiated on differentiation medium (0.5 mg L⁻¹ naphthalacetic acid and 3 mg L⁻¹ 6-benzyladenine). The positive calli that generated roots and shoots were transferred to 1/2MS medium to develop into T0 seedlings.

Apoplast and Total Soluble Protein Extraction

Roots were cut into approximately 5-cm segments and washed with deionized water as rapidly as possible. Segments were vacuum infiltrated for...
15 min in deionized water, producing a reduced pressure of 80 kPa. The vacuum was gradually released for 5 min. Segments were then dried and carefully arranged in a bundle in a Centriplus concentrator tube (Amicon). The tubes were centrifuged at 900 g for 15 min at 4°C to extract the apoplastic solution. The apoplast solution was lyophilized and stored at −80°C before analysis.

After the apoplastic solution had been extracted, the root tissue was ground to powder and extracted with 50 ml Tris-HCl buffer plus protease inhibitors, stirred for 15 min at 4°C, and centrifuged at 25,000g for 30 min at 4°C. The supernatant was used as total soluble protein for further analysis.

Testing Apoplastic Extract for Cytoplasmic Contamination

MDH activity was assayed by a modified method from Husted and Schjoerring (1995) in a reaction mixture containing 0.17 mM oxaloacetic acid, 0.094 mM NADH disodium salt, and 0.1 M Tris buffer, pH 7.4. Enzyme activity was assayed using a spectrophotometer at 340 nm by adding a 100-μl extract to 3 ml of assay medium at 25°C.

For immunoblots against tubulin, apoplastic proteins and total soluble proteins were separated by 12.5% SDS-PAGE and stained with Coomassie Brilliant Blue R-250 or transferred to a polyvinylidene difluoride membrane. The membrane was probed with anti-α-tubulin monoclonal antibody (Sigma, 1:2,000 dilution). Alkaline phosphatase-conjugated secondary antibody (Sigma, 1:10,000) was used to develop the blot.

2-DE

After quantification, 250 μl of buffer containing 150 μg of protein was loaded onto 13-cm isoelectric focusing strips, pH 3 to 10 linear gradient (Amersham Biosciences). 2-DE was carried out as described (Blum et al., 1987; Gorg et al., 1998) with minor modifications. Isoelectric focusing was conducted at 20°C using an IPGphore (Amersham Biosciences). The running conditions were as follows: 30 V for 12 h, 500 V for 1 h, 1,000 V for 1 h, and finally 8,000 V for 2 h. For the second dimension, the strips were incubated in an equilibration buffer containing 15 ms dithiothreitol for 20 min as the first step and then replaced by 2.5% iodoacetamide as the second step. The strips were placed on top of 12.5% polyacrylamide gels for SDS-PAGE according to Laemmli (1970) and sealed with 0.5% agarose. The electrophoresis was carried out at 10 mA per gel for 20 min and then 20 mA per gel by Hoefer SE 600 (Amersham Biosciences). The gels were stained with Coomassie Brilliant Blue R-250.

Image Analysis

The stained gels were scanned using Labscan software with image scanner (Amersham Biosciences). All experiments were repeated at least three times. Then, the gels were analyzed using Image Master 2D Elite software version 4.01 (Amersham Biosciences). After spot detection and normalization (in the total spot volume mode), the protein spots were matched and their volume or abundance was determined. A criterion of P < 0.05 was used to define significant differences when analyzing parallel spots between groups with one-way ANOVA using GraphPad Prim4 (GraphPad Software).

Protein Identification Using MALDI-TOF MS

In-gel digestion was performed according to Fulk et al. (2000). The mass spectra were recorded using a MALDI-TOF MS system (Bruker Autoflex). All obtained peptide mass fingerprints were searched against the National Center for Biotechnology Information database using Mascot (http://www.matrixscience.com) with a mass accuracy of 50 ppm in the parent ion mass. To denote a protein as an unambiguous identification, the following criteria were used: coverage of the mature protein by the matching peptides must reach a minimum of 15%, and at least five independent peptides should match within the protein sequence.

Molecular Cloning and Vector Construction

To construct 35S::OsRMC-YFP transient expression vector, the OsRMC 774-bp coding sequence containing a full-length ORF without a stop codon was amplified by reverse transcription (RT)-PCR, digested by NolI, and ligated to the pAVA321 vector (von Arnim et al., 1998). To construct OsRMCh::GUS vector, the 1,777-bp DNA sequence from upstream of the OsRMC ATG (start codon) was amplified, digested by HindIII and Xhol, and ligated to the pCAMBIA1300 vector. To construct the RNAi vector, the 428-bp coding sequence was amplified, digested by SacI and SpeI and then BamHI and KpnI, and ligated to the pCTK303 vector (Wang et al., 2004). To construct 3bi:: OsRMC-GUS overexpression vector, the 774-bp coding sequence was amplified, digested by BamHI and ligated to the pCAMBIA1300 vector. All primer sequences are listed in Supplemental Table S2.

Histochemical Analysis of OsRMC Promoter::GUS Activity

Histochemical analysis of GUS activity was according to Jefferson et al. (1987). Samples from different tissues of OsRMCh::GUS T1 transgenic lines were vacuum infiltrated for 15 min in 5-bromo-4-chloro-3-indolyl-β-D-glucuronic acid buffer and then incubated at 37°C for 12 h. After staining, the organs were destained in 70% ethanol several times until the chlorophyll was removed. GUS-positive samples were examined with a stereoscopic zoom microscope (Nikon), and digital images were recorded. Quantitative analysis of GUS activity was performed as described by Nakashima and Yamaguchi-Shinozaki (2002). The results represent averages from three independent experiments.

Subcellular Localization of OsRMC-Citrine Fusion Protein

The plasmids of the OsRMC-Citrine fusion protein construct and Citrine empty vector, which was used as a control, were transformed into the onion (Allium cepa) epidermal cells by particle bombardment in the Bio-Rad PDS-1000/He system according to the protocol. The transformed epidermal cells were cultured on Murashige and Skoog medium for 22 h and then treated with 0.9 M mannitol for plasmolysis. Observation was performed with a confocal microscope (Zeiss 510), and digital images were recorded.

Real-Time Q-PCR

The shoots of 10-d-old rice seedlings were collected for total RNA extraction (Trizol reagent; Invitrogen). Five hundred nanograms of total RNA was used as template for the RT reaction to synthesize the first-strand cDNA, which was used for SYBR Green-monitored real-time Q-PCR as described in the protocol. The analysis was performed with the use of an ABI Prism 7000 real-time PCR system (Applied Biosystems).

Southern Blot

The shoots of 10-d-old rice seedlings were collected for genomic DNA isolation and purification as described by Sambrook et al. (1989). Digested by HindIII and EcoRI, the fractionated DNA was electrophoresed on a 0.8% agarose gel at 60 V for 6 h and denaturized for blotting. The Hydrocyanic B-resistant gene from the vectors pCAMBIA1300 and pCTK303 was amplified and labeled with [α-32P]dCTP as a probe for hybridization. The image was scanned by Typhoon 9200.

Protein Purification and Antibody Preparation

The OsRMC coding sequence containing a full-length ORF without a stop codon was amplified by RT-PCR, digested by BamHI and EcoRI, and ligated to the pGEX-2T vector. Transformed into Escherichia coli strain BL21, the recombinant proteins extracted from the bacteria were purified on a GSTrap FF column according to the manufacturer’s (Amersham Biosciences) protocol. The antibody of OsRMC-GST was prepared by the Tailun Biological Technology company and tested by western blot.

Determination of MDA Content

The leaves of 10-d-old rice seedlings were weighed and homogenized in 1 ml of 10% TCA solution. The homogenate was centrifuged, and the supernatant was added to 0.6% thiobarbituric acid in 10% TCA. The mixture was incubated in boiling water for 15 min, and the reaction was stopped in an

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ice bath. Then, the samples were centrifuged and the absorbance of the supernatant was measured at 450, 532, and 600 nm. MDA contents (nmol g⁻¹ fresh weight) were calculated by the following formula: \[0.56(\text{A}_{532} - \text{A}_{600})/0.56\text{A}_{450}/\text{fresh weight} \].

Sequence data from this article can be found in the GenBank/EMBL data libraries under accession number AAL87185.

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

**Supplemental Figure S1.** MALDI-TOF mass fingerprint of OsRMC.

**Supplemental Figure S2.** LC-MS/MS results showing that spot 1 and spot 6 are identical proteins.

**Supplemental Figure S3.** Bioinformatic analysis for OsRMC.

**Supplemental Table S1.** Salt stress-regulated rice root apoplastic proteins.

**Supplemental Table S2.** Primer sequences used in gene construction and real-time Q-PCR.

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