Direct Phosphorylation and Activation of a Mitogen-Activated Protein Kinase by a Calcium-Dependent Protein Kinase in Rice

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The mitogen-activated protein kinase (MAPK) is a pivotal point of convergence for many signaling pathways in eukaryotes. In the classical MAPK cascade, a signal is transmitted via sequential phosphorylation and activation of MAPK kinase kinase (MAPKKK or MKKK), MAPK kinase (MAPKK or MKK), and MAPK. The activation of MAPK is dependent on dual phosphorylation of a TXY motif by an MKK, which is considered the sole kinase to phosphorylate and activate MAPK. Here, we report a novel regulatory mechanism of MAPK phosphorylation and activation besides the canonical MAPK cascade. A rice (Oryza sativa) calcium-dependent protein kinase (CDPK), CPK18, was identified as an upstream kinase of MAPK (MPK5) in vitro and in vivo. Curiously, CPK18 was shown to phosphorylate and activate MPK5 without affecting the phosphorylation of its TXY motif. Instead, CPK18 was found to predominantly phosphorylate two Thr residues (Thr-14 and Thr-32) that are widely conserved in MAPKs from land plants. Further analyses reveal that the newly identified CPK18-MPK5 pathway represses defense gene expression and negatively regulates rice blast resistance. Our results suggest that land plants have evolved an MKK-independent phosphorylation pathway that directly connects calcium signaling to the MAPK machinery.

INTRODUCTION

Plants have evolved sophisticated signaling pathways to transduce biotic or abiotic stimuli into proper cellular responses in order to adapt to ever-changing environmental challenges. These signaling pathways often require the activation of protein kinases to orchestrate diverse cellular processes through the phosphorylation of various substrates. The eukaryotically conserved mitogen-activated protein kinase (MAPK) cascade is a pivotal phosphorylation pathway to convey external or internal signals to downstream effectors such as transcription factors. The prototypic MAPK cascade consists of three sequentially phosphorylated and activated protein kinases: MAPK kinase kinase (MAPKKK or MKKK), MAPK kinase (MAPKK or MKK), and MAPK (Seger and Krebs, 1995). Following the dual phosphorylation at the threonine (T) and tyrosine (Y) residues in its TXY motif (where X is Asp or Glu in plants), a MAPK is activated and becomes capable of phosphorylating downstream transcription factors and other effectors to reprogram the transcriptome and metabolome (Seger and Krebs, 1995; Chang and Karin, 2001; Pitzschke et al., 2009; Rodriguez et al., 2010; Meng and Zhang, 2013). As a result, MKK is generally considered the sole kinase to phosphorylate and activate MAPKs (Andreasson and Ellis, 2010; Rodriguez et al., 2010; Meng and Zhang, 2013).

Land plants typically encode more MKKKs, MKKs, and MAPKs than yeasts or mammals. Arabidopsis thaliana and rice (Oryza sativa) contain 20 and 17 MAPKs (MAPK Group, 2002; Reyna and Yang, 2006), respectively, which can be divided into four phylogenetic groups (Supplemental Figure 1). Among them, Arabidopsis MPK3/4/6 and their orthologs from group A and B MAPKs play a key role in stress signal transduction associated with reactive oxygen species homeostasis, stomata development, hormone signaling, disease resistance, and abiotic stress tolerance (Pitzschke et al., 2009; Rodriguez et al., 2010; Meng and Zhang, 2013). For example, rice MPK5, the ortholog of Arabidopsis MPK3 (Supplemental Figure 1 and Supplemental Data Set 1), was previously shown to positively regulate abiotic stress tolerance but negatively modulate rice disease resistance (Xiong and Yang, 2003). During the past decade, many studies have focused on deciphering the components of the classical MAPK cascade involved in developmental or stress signal transduction, such as plant immunity signaling. Two extensively studied canonical MAPK cascades in Arabidopsis, MKK4/5-MPK3/6 and MEKK1-MKK1/2-MPK4, were activated by pathogen-associated molecular patterns or effectors to regulate downstream signaling components. Activated MPK3/6 was shown to phosphorylate transcription factors like WRKY33, ETHYLENE-RESPONSE FACTOR104 (ERF104), and VIRE2-INTERACTING PROTEIN1 to activate defense gene expression and induce ethylene (ET) camalexin biosynthesis (Djamei et al., 2007; Bethke et al., 2009; Mao et al., 2011). However, how receptors/sensors transmit different signals to MAPKs remains a major gap in our understanding of plant MAPK cascade signaling (Meng and Zhang, 2013).
Besides the MAPK cascade, calcium (Ca\(^{2+}\)) signaling is another critical pathway triggered by environmental stimuli and developmental cues. Ca\(^{2+}\) is a ubiquitous secondary messenger in cellular signal transduction and is decoded by various Ca\(^{2+}\) binding proteins and kinases such as calcium-dependent protein kinase (CDPK) (Harper et al., 2004). CDPK is an ancient family of Ser/Thr kinases that has evolved independently in green algae and land plants (Hamel et al., 2014). There are 34 and 29 CDPKs in Arabidopsis and rice (Cheng et al., 2002; Asano et al., 2005), respectively. Many CDPKs contain a myristoylation motif (Hrabak et al., 2003), and a few of them were confirmed to be associated with the plasma membrane (Dammann et al., 2003). Biochemical and genetic studies have shown that CDPKs play important roles in numerous signaling pathways and biological processes such as hormone signaling, oxidative burst, innate immunity, and abiotic stress response (Cheng et al., 2002; Harper et al., 2004; Lecourieux et al., 2006; Reddy et al., 2011).

Since the CDPK and MAPK pathways are often activated in response to the same environmental stimuli, it has been speculated that a potential crosstalk may occur between the two important families of protein kinases (Ludwig et al., 2005; Wurzinger et al., 2011). However, recent studies in Arabidopsis indicated that stress-responsive CDPKs and MAPKs acted differentially in innate immunity and salt signaling without direct crosstalk (Boudsocq et al., 2010; Mehmer et al., 2010). Therefore, whether and how Ca\(^{2+}\)/CDPK signaling may regulate the MAPK cascade in plants is largely unknown. In this study, a plasma membrane–associated rice CDPK, CPK18, was found to directly phosphorylate and activate MPK5 in vitro and in vivo. Genetic and functional analyses demonstrate that the CPK18-MPK5 pathway negatively regulates rice immunity and represents a novel regulatory mechanism for the MKK-independent activation of MAPKs in land plants.

RESULTS

CPK18 Is Both a Kinase and a Substrate of MPK5 in Vitro

In a screening assay of MPK5 substrates by in situ solid-phase phosphorylation (Fukunaga and Hunter, 1997), CPK18 was initially identified as a potential MPK5 substrate. These two kinases were then found to phosphorylate each other when in vitro kinase assays were performed using the polyhistidine (His)-tagged recombinant proteins (His-CPK18 and His-MPK5). To confirm their phosphorylation, kinase-defective recombinant proteins, in which CPK18 Asp-178 was substituted with Ala (His-CPK18DA) and MPK5 Lys-65 was substituted with Arg (His-MPK5KR), were used as substrates. His-CPK18DA retained barely detectable autophosphorylation activity and His-MPK5KR had no detectable autophosphorylation (Figure 1A). An in vitro kinase assay showed that His-CPK18 phosphorylated His-MPK5KR, whereas His-MPK5 phosphorylated His-CPK18DA (Figure 1A). The in vitro interaction between MPK5 and CPK18 was also confirmed by the glutathione S-transferase (GST) pull-down assay (Figure 1B). These results demonstrated that direct interaction and phosphorylation occurred between CPK18 and MPK5 in vitro.

The existence of a CDPK interacting with and phosphorylating MAPK (Figure 1) is highly intriguing, because the eukaryotically conserved MAPK is typically known to be only phosphorylated by MKK. Based on the protein kinase phylogeny, CDPK is a plant- and protist-specific group in the CAMK kinase superfamily and is distinct from MKK, which belongs to the STE kinase superfamily (Dardick et al., 2007). CPK18 is a typical group IV CDPK (Supplemental Figure 2) with four domains: the N-terminal variable domain (amino acids 1 to 40) with a myristoylation site for membrane association, the protein kinase domain (amino acids 40 to 312), the autoinhibitory junction domain (amino acids 313 to 350), and the calmodulin-like domain containing four EF-hand motifs (amino acids 351 to 512). Consistent with the CDPK and MKK phylogeny, the in vitro kinase assay showed that CPK18 did not affect the phosphorylation level of the MPK5 TEY motif, which was phosphorylated by the rice MKK MKK4/6 (Asano et al., 2005) or by itself (autophosphorylation) in the canonical MAPK cascade (Supplemental Figure 3).

CPK18 Interacts with and Phosphorylates MPK5 in Vivo

We then sought evidence that CPK18 interacts with MPK5 in vivo. Their interaction was first tested by coimmunoprecipitation. To this end, a constitutively activated CPK18 (in which the autoinhibitory and calmodulin domains were removed) with a FLAG tag at its C terminus (CPK18C-FLAG) was expressed in rice plants.

Figure 1. Direct Interaction and Phosphorylation between CPK18 and MPK5 in Vitro.

(A) Recombinant His-CPK18 and His-MPK5 phosphorylated kinase-defective His-MPK5KR (K65R; labeled as KR) and His-CPK18DA (D178A; labeled as DA), respectively. The size difference between CPK18 and CPK18DA was due to autophosphorylation. Recombinant His-MPK5 purified from E. coli was capable of autophosphorylation and possessed the catalytic activity. Asterisks indicate bacterial protein contaminants with His-MPK5KR. CBB, Coomassie Brilliant Blue staining.

(B) MPK5 interacted with CPK18 in the GST pull-down assay. WB, immunoblotting.
Immunoblotting, using the MPK5-specific antibody (Xiong and Yang, 2003), showed that MPK5 was specifically bound to CPK18AC-FLAG (Figure 2A). A bimolecular fluorescence complementation (BiFC) assay was also performed to investigate the subcellular localization of CPK18, MPK5, and their complex. Transient expression of green fluorescent protein (GFP)-tagged MPK5 and CPK18 in onion (Allium cepa) epidermal cells and rice protoplasts showed that MPK5 was localized in the nucleus and cytoplasm, whereas CPK18 was predominantly associated with the plasma membrane (Figure 2B). These observations are in agreement with previously reported locations of CPK18 (Campos-Soriano et al., 2011) and MPK5 (Singh et al., 2012). When MPK5-nYFP (the N-terminal half of YFP [yellow fluorescent protein]) and CPK18-cYFP (the C-terminal half of YFP) were coexpressed in onion cells or rice protoplasts, the complemented YFP signal was observed along the plasma membrane, as was CPK18-GFP (Figure 2B). These results confirmed that CPK18 interacted with MPK5 in vivo.

**Figure 2.** CPK18 Interacted with and Phosphorylated MPK5 in Vivo.

(A) MPK5 was coimmunoprecipitated (IP) with FLAG-tagged CPK18 in vivo. MPK5 and CPK18 were detected in input control or coimmunoprecipitated samples by immunoblotting (WB) with anti-MPK5 and anti-FLAG antibodies, respectively.

(B) Subcellular localization of CPK18-GFP, MPK5-GFP, and their complex (BiFC) in onion epidermal cells or rice protoplasts. Bars = 100 µm (onion cells) and 10 µm (rice protoplast).

(C) In-gel kinase assay shows that CPK18 phosphorylated MPK5KR and MPK5AEF in a calcium-dependent manner. Total rice proteins from wild-type and CPK18-RI plants were resolved by SDS-PAGE containing His-MPK5KR or His-MPK5AEF. The in-gel kinase reaction was performed in the presence of Ca^{2+} (2 mM) or EGTA (5 mM) as indicated. A gel without a substrate was used as a control to monitor autophosphorylation (right panel). The kinase was visualized by autoradiography, and the relative activity toward phosphorylating His-MPK5KR or His-MPK5AEF was quantified with normalization to protein loading and shown at the bottom of the gel. Protein loadings are indicated by Coomassie Brilliant Blue (CBB) staining. MW, molecular mass marker in kD.

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To confirm that native CPK18 phosphorylates MPK5, we generated transgenic plants (CPK18-RI lines) in which CPK18 expression was specifically suppressed by RNA interference (Supplemental Figure 4). In three independent CPK18-RI lines (lines 9, 10, and 13), CPK18 mRNA was reduced by 70 to 80% in comparison with the wild type, but the expression of CPK4, which is the closest homolog of CPK18 in rice (Supplemental Figure 2), was not affected (Supplemental Figure 4). By contrast, the mRNA levels of MKK4 and MKK6, two MKKs upstream of MPK5 in the classical MAPK cascade, were increased slightly (~1.2- to 1.5-fold) in CPK18-RI lines (Supplemental Figure 4). To examine native kinases that phosphorylate MPK5 in CPK18-RI and wild-type plants, in-gel kinase assays were performed using the highly purified His-MPK5KR or His-MPK5AEF (where Thr-194, Glu-195, and Tyr-196 were replaced by Ala, Asp, and Phe) as a substrate. A protein kinase, with a molecular mass (~58 to 60 kD) comparable to rice CDPKs but distinct from two 40-kD MKKs (MKK4 and MKK6), was the only kinase we identified to phosphorylate His-MPK5KR and His-MPK5AEF in the presence of Ca²⁺ (Figure 2C). We barely detected any kinases phosphorylating His-MPK5KR in the presence of EGTA (a Ca²⁺ chelator), nor did we detect any autophosphorylation without substrates in the gel (Figure 2C). We reasoned that the detected CDPKs included CPK18, because their ability to phosphorylate His-MPK5KR or His-MPK5AEF was specifically reduced in CPK18-RI lines (Figure 2C). In comparison with the wild type, the overall activity of CDPKs toward phosphorylating His-MPK5KR was reduced approximately 70% in the strongest CPK18-RI line (line 13), indicating that CPK18 is likely a predominant CDPK that phosphorylates MPK5 in rice. Taken together, these results suggest that CPK18 phosphorylates MPK5 at a site(s) out of its TEY motif in vivo.

Substrate Specificity of CDPK-MAPK Phosphorylation

We further examined the specificity of CPK18 toward phosphorylating different MAPKs. Five additional MAPKs, rice MPK4/6, *Arabidopsis* MPK3, tobacco (*Nicotiana tabacum*) salicylic acid-induced protein kinase (SIPK) and wounding-induced protein kinase (WIPK) (Zhang and Klessig, 1998), and two rice CDPKs (CPK4 and CPK7), were fused with His or GST tags and purified from *Escherichia coli*. An in vitro kinase assay showed that rice CPK18 phosphorylated four MAPKs (Os MPK6, At MPK3, Nt WIPK, and Nt SIPK) from groups A and B in addition to MPK5 (Figure 3A). On the other hand, both rice group IV CDPKs, CPK4 and CPK18, phosphorylated MPK5KR (Figure 3B).

**Figure 3.** Phosphorylation Specificity between CDPKs and MAPKs.  
(A) CPK18 phosphorylated tobacco WIPK and SIPK, rice MPK6, and *Arabidopsis* MPK3 but hardly phosphorylated rice MPK4. The autophosphorylation of MAPKs was blocked by the mutation of essential residues (SIPK-KR and WIPK-KR) or by adding the MAPK inhibitor SB202190 (right panel; Os MPK6 and At MPK3, except for rice MPK4, which had barely detectable autophosphorylation. At, *Arabidopsis*; MW, molecular mass marker; Os, rice. Bacterial protein contaminants copurified with MAPKs are indicated with asterisks on the Coomassie Brilliant Blue (CBB)-stained gels.  
(B) Rice CPK4 phosphorylated MPK5KR but CPK7 did not. The common CDPK substrate Histone III (Histone type III-S) was used as a positive control to show GST-CPK7 activity. The asterisk indicates a protein contaminant in Histone III.
whereas the group I rice CDPK, CPK7, did not (Figure 3B). We speculate that phosphorylation between CDPKs and MAPKs likely exists between specific subgroups of CDPK and MAPK families.

**CPK18 Predominantly Phosphorylates Thr-14 and Thr-34 in the N Terminus of MPK5**

Next we determined which Thr/Ser on MPK5 was phosphorylated by CPK18. A total of 16 Ser/Thr residues (Supplemental Figure 5) were predicted as potential CPK18 phosphorylation sites according to the substrates’ specificity (Figure 3) or common CDPK phosphorylated motifs (Hegeman et al., 2006). Using site-directed mutagenesis, these candidate sites were substituted by Ala in the kinase defect MPK5KR, and 13 mutated recombinant proteins were made to cover all 16 substitutions (Supplemental Figure 6). By comparing the phosphorylation levels of these mutated proteins with MPK5KR, two Thr residues (Thr-14 and Thr-32) emerged as CPK18 phosphorylated sites (Supplemental Figure 6). In comparison with His-MPK5KR, T14A and T32A substitutions removed ∼37% and 75% of CPK18 phosphorylation, respectively (Figure 4).

**Figure 4. CPK18 Phosphorylated MPK5 at Thr-14 and Thr-32.**

(A) In vitro kinase assay shows that the T14A and T32A mutations drastically impaired MPK5KR phosphorylation by CPK18. The relative phosphorylation level of MPK5 mutants is indicated at the bottom. The asterisks indicate bacterial protein contaminants.

(B) Analysis of MPK5 phosphorylation using a Phos-tag gel. Two phosphorylated forms of MPK5KR (forms 1 and 2) were detected on the Coomassie Brilliant Blue (CBB)-stained Phos-tag gel. The asterisks indicate bacterial protein contaminants.

(C) In vivo phosphorylation profiles of MPK5 mutants. FLAG-tagged MPK5KR, MPK5KR-AA (T14A-T32A-K65R), and MPK5KR-AEF (K65R-T194A-Y196F) were expressed in rice protoplasts, and their phosphorylation patterns were analyzed using a Phos-tag gel and immunoblotting. The percentage of differentially phosphorylated forms (no-, low-, and high-phos) was quantified and indicated. WB, immunoblotting.

(D) CPK18 phosphorylation sites (indicated with arrows) are highly conserved in plant group A and B MAPKs but exist neither in group C plant MAPKs nor in MAPKs of humans and C. reinhardtii. N-terminal sequences of all group A/B/C MAPKs from Arabidopsis (At MPK) and rice (Os MPK) were aligned. Two tobacco MAPKs (WIPK and SIPK), human ERK2 and P38, P. patens (Pp) MAPK, and C. reinhardtii (Cre) MAPK8 were also included for comparison. The number at the end of each line indicates the coordinate of the last residue. The high- and medium-level conserved residues among MAPKs are indicated in the alignment with black and gray shading, respectively.

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However, mutated MPK5 proteins, which had both Thr-14 and Thr-32 replaced by other amino acids, were completely insoluble in *E. coli*. As a result, we used Phos-tag SDS-PAGE (Phos-tag gel) to confirm the phosphorylation sites. A Phos-tag gel could separate the phosphorylated proteins according to the number of phosphate groups attached (Kinoshita-Kikuta et al., 2007), and a protein containing more phosphorylated residues would have a lower migration rate in the Phos-tag gel. After phosphorylation by His-CPK18, two distinct forms of phosphorylated His-MPK5KR (phos-MPK5) were detected according to their migration rate in a Phos-tag gel (Figure 4B), suggesting that CPK18 phosphorylated at least two sites on His-MPK5KR. In agreement with the initial phosphorylated site mapping data, both T14A and T32A mutations drastically reduced the phos-MPK5 level by CPK18 (Figure 4B). Because we still observed two faint phosphorylated forms of His-MPK5KR-T14A, it is possible that CPK18 may weakly phosphorylate another site on MPK5 besides Thr-14 and Thr-32. These results suggest that MPK5 Thr-14 and Thr-32 are two major sites, and Thr-32 is the predominant site, for CPK18 phosphorylation in vitro.

We further sought evidence that Thr-14 and Thr-32 were phosphorylated in vivo using the Phos-tag gel. We compared the in vivo phosphorylation pattern of MPK5KR, MPK5KR-AEF (K65R-T194A-Y196F), and MPK5KR-AA (K65R-T14A-T32A), which were expressed in rice protoplasts with a FLAG tag. MPK5 with these mutations showed identical migrating rates in normal SDS-PAGE, although MPK5KR-AA showed much lower protein expression than the others (Figure 4C). Consistent with multiple phosphorylation sites, multiple phosphorylation forms of MPK5KR were detected in the Phos-tag gel and are referred to as no-, low-, and high-phosphorylated forms according to their migration rates (Figure 4C). The percentage of high-phosphorylated MPK5KR, which had the lowest migrating rate, was substantially reduced when the TEY motif was replaced by AEF (Figure 4C). Although MPK5KR-AA had a similar phosphorylation pattern to that of MPK5KR, ~30% of MPK5KR-AA remained nonphosphorylated, while only ~10% of MPK5KR or MPK5KR-AEF remained nonphosphorylated (Figure 4C). The T14A-T32A mutations mainly impaired the low-phosphorylated form, which was likely attached with one phosphate group. We noticed that the T14A-T32A mutations appeared to increase the percentage of the high-phosphorylated form compared with MPK5KR (Figure 4C). This is probably due to compensatory phosphorylation in a protein with multiple phosphorylation sites, as observed previously (Bauer et al.,...)

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**Figure 5.** CPK18 Positively Regulates MPK5 Activity.

(A) MPK5-T32D, containing the T32A mutation to mimic phosphorylation, exhibited increased kinase activity toward MBP. The relative radioactivity is shown at the bottom.

(B) and (C) Detection of overall MAPK activity and MPK5 activity in CPK18-RI and wild-type plants using an in-gel kinase assay. Total protein (B) and immunoprecipitated MPK5 (C) were resolved on 10% SDS-PAGE gels containing MBP. The in-gel kinase assay was performed to analyze MAPK activity. The relative MAPK activity is shown at the bottom of (B). Protein loading is shown by Coomassie Brilliant Blue (CBB) staining.

(D) MPK5 protein level and overall MAPK TEY phosphorylation level in wild type and CPK18-RI lines. MPK5 and phos-TEY levels were examined by immunoblotting (WB) with anti-MPK5 and anti-phos-TEY antibodies, respectively.

(E) RT-qPCR examination of MPK4/5/6 expression in the wild type and CPK18-RI lines. Data are presented as means ± so (n = 3). [See online article for color version of this figure.]
Taken together, both in vitro and in vivo data have shown that CPK18 phosphorylated MPK5 mainly at the Thr-14 and Thr-32 sites.

Thr-14 and Thr-32 Residues Are Conserved in Group A and B MAPKs of Land Plants

Sequence alignment of Arabidopsis and rice MAPKs indicates that all group A/B MAPKs have Ser/Thr aligned with MPK5 Thr-14 and Thr-32, with highly conserved context sequences except Arabidopsis MPK5/10/11 (Figure 4D). Although the context sequence–flanked MPKS Thr-32 is conserved in all MAPKs, MAPKs in other groups do not have Ser/Thr at the position aligned with the Thr-32 site (Figure 4D). We also examined the sequences of putative MPK5 orthologs in tobacco, moss (Physcomitrella patens), green algae (Chlamydomonas reinhardtii), and human (Homo sapiens). As is the case for Arabidopsis and rice, both Thr-32 and Thr-14 were also conserved in tobacco and P. patens MAPKs but absent in C. reinhardtii and human MAPKs (Figure 4D). These results suggest that CPK18 phosphorylated sites are conserved in MAPKs from both lower and higher plants and might have emerged during the evolution of land plants.

CPK18 Positively Regulates MPK5 Activity

To examine the effect on MPK5 activity of CPK18 phosphorylation, MPK5-T32D was made to mimic phosphorylated Thr-32, which is a predominant phosphorylation site. Upon expression in E. coli, His-MPK5T32D showed comparable protein yield and solubility.
to His-MPK5. An in vitro kinase assay showed that recombinant MPK5-T32D exhibited twice as much catalytic activity toward phosphorylating the common MAPK substrate myelin basic protein (MBP) than wild-type MPK5, but their autophosphorylation levels were comparable (Figure 5A).

Then, we further investigated whether CPK18 regulates MPK5 activity in rice plants. We compared overall MAPK activity in CPK18-RI and wild-type plants by in-gel kinase assays using MBP as a substrate. The activity of ~43-kD MAPKs was reduced 40 to 80% in three CPK18-RI lines compared with the wild type (Figure 5B). The result also showed that the activity of another kinase(s) with a size of ~55 to 65 kD was reduced in CPK18-RI plants, but its identity was undefined (Figure 5B). Because most rice group A/B/C MAPKs have similar molecular masses (43 to 45 kD), the activity of native MPK5 was further measured after immunoprecipitation. As shown by the in-gel kinase assay, immunoprecipitated MPK5 from CPK18-RI plants also exhibited a lower kinase activity than that from wild-type plants (Figure 5C). By contrast, the MPK5 protein level was increased slightly in CPK18-RI plants compared with the wild type, and the overall phos-TEY level of MAPKs was comparable between CPK18-RI and the wild type (Figure 5D). Further expression profiling of three rice MAPKs reveals that the transcripts of MPK5 and MPK6, which encode CPK18 substrates (Figure 3A), were increased, but MPK4 expression was not significantly changed in CPK18-RI plants compared with the wild type (Figure 5E). Thus, we conclude that CPK18 positively regulates MPK5 activity through phosphorylating Thr-14/Thr-32 without affecting MPK5 TEY phosphorylation.

The CPK18-MPK5 Pathway Negatively Regulates Rice Blast Resistance

To explore the physiological function of the CPK18-MPK5 pathway, the phenotype of CPK18-RI lines was compared with that of MPK5-RI lines, in which MPK5 was suppressed by RNA interference (Xiong and Yang, 2003). Similar to MPK5-RI plants, three CPK18-RI lines exhibited enhanced resistance to the rice blast fungus (Magnaporthe oryzae) (Figures 6A and 6B). After fungal infection, the size and number of disease lesions or relative fungal amounts in leaves of CPK18-RI lines were significantly lower than those in wild-type plants. The enhanced resistance phenotype is consistent with the observations that the expression of rice defense-related genes (pathogenesis-related genes PR5 and PR10, Chitinase, and Harpin-induced protein1 [Hin1]) was increased in CPK18-RI and MPK5-RI lines (Figure 6C). Although MPK5 also phosphorylated CPK18 in vitro (Figure 1A), an in-gel kinase assay suggested that the CDPK activity (including CPK18) was comparable between wild-type and MPK5-RI plants (Supplemental Figure 7). To confirm that phosphorylation at MPK5 Thr-14/Thr-32 contributes to the repression of defense genes, we used the β-glucuronidase (GUS) reporter assay to examine the PR5 and Hin1 promoter activities after expressing FLAG-tagged MPK5 or MPK5-DD (T14D-T32D) in rice protoplasts. Consistent with the quantitative RT-PCR (RT-qPCR) results (Figure 6C), PR5 and Hin1 promoter activities were repressed in rice protoplasts expressing MPK5 in comparison with the vector control (Figure 6D). MPK5DD that mimicked the Thr-14 and Thr-32 phosphorylation exhibited stronger repression; even its protein level was much less than that of MPK5 (Figures 6D and 6E). These results suggest that the CPK18-MPK5 pathway negatively regulates rice immunity by repressing defense gene expression.

DISCUSSION

In the classical MAPK cascade, MKK has been recognized as the sole kinase to transmit signals from upstream MAPKKKs to MAPKs in plant, animal, and yeast systems. Three exceptions have been
reported for the activation of the mammalian MAPK P38α (Ge et al., 2002; Salvador et al., 2005) or the Arabidopsis MAPK MPK8 (Takahashi et al., 2011) via an MKK-independent pathway. P38α could be activated after TAK1 Binding Protein1 binding (Ge et al., 2002) or through Tyr-323 phosphorylation by T cell receptor-proximal Tyr kinases (Salvador et al., 2005), and both pathways activated P38α by triggering autophosphorylation of the TDY motif. The full activation of Arabidopsis MPK8 was found to require a calmodulin protein binding to its TDY motif in addition to MKK phosphorylation (Takahashi et al., 2011). The CPK18-MPK5 pathway described in this study is distinct from these reported examples of MKK-independent activation of MAPKs (Ge et al., 2002; Salvador et al., 2005; Takahashi et al., 2011). CPK18 triggered MPK5 activation by phosphorylating its N-terminal Thr residues (Thr-14 and Thr-32) without affecting the phos-TEY level (Figure 5; Supplemental Figure 3). Hence, the CPK18-MPK5 pathway represents a novel mechanism to directly modulate MAPK activity in addition to the canonical MAPK cascade.

It is intriguing that the N-terminal Thr-14/Thr-32 residues and their context sequence in MPK5 are conserved in almost all Arabidopsis and rice group A and B MAPKs (Figure 4D). We have demonstrated that at least five MAPKs from three plant species could be directly phosphorylated by CPK18 (Figure 3A). In addition, both group IV rice CDPKs, CPK4 and CPK18, are capable of phosphorylating MPK5 (Figure 3B). Thus, the CPK18-MPK5 pathway may not be an exception but likely represents a conserved regulatory mechanism for a specific subset of CDPKs to modulate group A and B MAPKs in plants. Interestingly, group IV CDPKs appear to have evolved only in land plants (Hamel et al., 2014), whereas Thr-14/Thr-32 residues are conserved in the MAPKs of land plants but are absent in green algae (Figure 4D). This evolutionary coincidence suggests that the CDPK-MAPK pathway might have emerged during the evolution of land plants as an adaptation to environmental changes.

A major effort has been made to study MAPK cascades and Ca²⁺/CDPK signaling in plants, but little is known about the cross-talk between these two important kinase pathways. Although both pathways may share downstream components, previous studies in Arabidopsis and tobacco suggest that CDPKs and MAPKs work in parallel without direct interaction or crosstalk (Boudsocq et al., 2010). The phosphorylation of MPK5 and other MAPKs by CPK18 shown in this study indicates that the Ca²⁺ signaling pathway can directly activate MAPKs through specific CDPKs. Ca²⁺ is one of the earliest secondary signals to sense microbial infection, abiotic stresses, or phytohormones such as abscisic acid (ABA) (McAinsh et al., 1992; Kurusu et al., 2005). The CPK18-MAPK pathway potentially provides a shortcut or an alternative route to activate MAPKs without the multiple phosphorylation steps involved in the canonical MAPK cascade. As shown in our proposed working model (Figure 7), the Ca²⁺ signal activates the membrane-anchored CPK18, which in turn phosphorylates MPK5 to repress defense gene expression. Ca²⁺ signals triggered by environmental stimuli (e.g., pathogen infection, drought, or ABA) could be rapidly decoded by CPK18 and transmitted to transcriptional machinery through the CPK18-MAPK pathway by taking advantage of the fact that MAPKs target numerous transcription factors (Popescu et al., 2009). Since CPK18 does not affect MPK5 TEY phosphorylation (Figures 5A and 5D; Supplemental Figure 3), multiple phosphorylated forms of MPK5 likely coexist (Figure 4C), and the CPK18-MAPK pathway may work independently or cooperatively with the classical MAPK cascade.

Our observations suggest that the newly identified CPK18-MPK5 pathway represses defense gene expression, thereby increasing host susceptibility to rice blast fungus (Figure 6). Recent advances indicate that MPK5 likely mediates the interplay of plant hormones to fine-tune immunity and growth (Yang et al., 2013). Among the phytohormones, ET appears to play a prominent role in rice blast resistance (Helliwell et al., 2013). In Arabidopsis, MPK3 and MPK6 positively regulate the ET pathway by phosphorylating and stabilizing 1-aminocyclopropane-1-carboxylic acid synthase, which catalyzes ET production (Liu and Zhang, 2004), or transcription factors (ETHYLENE-INSENSITIVE3 and ERF104) that are essential components of ET signaling (Yoo et al., 2008; Bethke et al., 2009). A similar phosphorylation event likely exists in rice, but it may lead to a different biochemical and physiological consequence. Indeed, elevated ET production was observed in MPK5-RI plants (Bailey et al., 2009), suggesting the repression of the ET pathway by MPK5. By mediating antagonistic crosstalk between ET and ABA, MPK5 is a positive regulator of the ABA pathway and abiotic stress tolerance (Xiong and Yang, 2003). Interestingly, MPK5 is required for ABA-mediated brown spot resistance in rice (De Vleesschauwer et al., 2010). Taking into account the fact that abiotic stresses and ABA rapidly trigger cellular Ca²⁺ signals that could be decoded by CDPK, the CPK18-MPK5 pathway likely mediates the crosstalk among abiotic stress, ABA, and ET pathways that fine-tune rice growth, immunity, and abiotic stress tolerance.

Moreover, in vitro experiments suggest that phosphorylation between CPK18 and MPK5 may occur in both directions (i.e., CPK18 phosphorylates MPK5 while MPK5 feedback phosphorylates CPK18) (Figure 1A). However, the molecular and physiological consequences of the MPK5-CPK18 phosphorylation were not demonstrated or elucidated in this study (Figure 7). Bidirectional phosphorylation between two kinases is rarely reported but has been observed between specific MKKs and MAPKs in human, Xenopus, and yeast (Matsuda et al., 1993; Mansour et al., 1994; Jiménez-Sánchez et al., 2007). Further studies are needed to reveal if such bidirectional phosphorylation exists in plants to provide a simple mechanism to mediate crosstalk between the CDPK and MAPK pathways. In conclusion, our results demonstrate that plants have evolved a CDPK-mediated pathway for MAPK activation in addition to the linear model of the classical MAPK cascade.

METHODS

Plant Materials and Growth Conditions

Rice (Oryza sativa) cv Kitaake was used to generate CPK18-RI lines in this study, whereas cv Nipponbare was used previously to generate MPK5-RI lines (Xiong and Yang, 2003). The rice plants were grown in a greenhouse or a growth chamber at 28°C day/23°C night with 12 h of light.

In Situ Solid-Phase Phosphorylation Screen

To screen putative MPK5 substrate, a rice cDNA library was constructed from Magnaporthe oryzae–infected rice leaves using the ZAP Express cDNA Synthesis Kit and the ZAP Express cDNA Gigapack III Gold Cloning
Kit (Stratagene). The in situ solid-phase screen was performed according to the published method that was used to identify mammalian MAPK substrates (Fukunaga and Hunter, 1997).

**Gene Constructs for Recombinant Protein Expression**

The full-length cDNA of CPK18 in the pBK-CMV phagemid vector (Stratagene) was cut with EcoRI and XhoI and inserted into pGEX-5x-3 and pET28c, which were used to express GST-CPK18 and His-CPK18, respectively, in *Escherichia coli* which were used to express GST-CPK18 and His-CPK18, respectively, in the published method that was used to identify mammalian MAPK substrates (Fukunaga and Hunter, 1997).

To express and purify substrates for an in-gel kinase assay, His-MPK5KR or His-MPK5AEF was induced with 1 mM IPTG at 37°C for 4 h. After induction, bacteria were harvested and resuspended in lysis buffer (25 mM Tris, pH 7.4, 10 mM EDTA, and 1% Triton X-100) and then disrupted by sonication in ice (Branson Sonifier). Insoluble inclusion bodies were collected by centrifugation at 10,000g for 15 min. After washing three times with lysis buffer, inclusion bodies were dissolved in solubilized buffer (50 mM Tris, pH 9.5, and 0.3% sarkosyl) at 10 mg/mL. The insoluble cell debris was removed by centrifugation. The dissolved His-MPK5KR or His-MPK5AEF was then dialyzed against 20 mM Tris, pH 8.8, with three changes of buffer. The purity of His-MPK5KR and His-MPK5AEF was higher than 95% based on estimation by SDS-PAGE.

**Gene Constructs for Rice Transformation**

To express CPK18 and MPK5 in plants, the complete open reading frames without a stop codon of MPK5 and CPK18 were amplified (primers are listed in Supplemental Table 1) and then cloned into pDEST17 or pDEST15 by the LR reaction for recombinant protein expression with a His or GST tag (Life Technologies). Arabidopsis thaliana MPK3 in a pENTR vector was obtained from the ABRC (http://abrc.osu.edu/) and cloned into pDEST17 to express His-MPK3.

**In Vitro Kinase Assay**

For the in vitro kinase assay, 0.5 μg of recombinant kinase (His-CPK18 or His-MPK5) and 2 μg of substrate were added to a 20-μL reaction containing 25 mM Tris, pH 7.5, 10 mM MgCl2, 1 mM DTT, 0.2 mM ATP, and 0.1 μCi of [γ-32P]ATP (MP Biomedicals) supplemented with 5 mM CaCl2 (for the CDPK assay), 5 mM MnCl2 (for the MAPK assay), or 50 μM SB202190 (LC Laboratories). The reaction was incubated at room temperature (25°C) for 1 h and terminated with SDS sample buffer. After separation with SDS-PAGE and staining, gels were washed thoroughly with gel washing buffer (5% trichloroacetic acid and 1% sodium pyrophosphate) and then washed thoroughly to remove 32P-labeled protein. Protein phosphorylation was analyzed by autoradiography with the Storage Phosphor Screen and Storm 820 scanner (GE). The relative phosphorylation level was quantified using ImageJ (http://rsb.info.nih.gov/ij/).

**In-Gel Kinase Assay**

For the in-gel kinase assay to detect CDPK, total protein was extracted from rice plants using extraction buffer (25 mM Tris, pH 7.6, 150 mM NaCl, 50 mM glyceraldehyde phosphate, 5 mM NaF, 5 mM Na3VO4, 1 mM DTT, 1% Triton X-100, 10% glycerol, 1 mM phenylmethylsulfonyl fluoride (PMSF), and 1% Protease Inhibitor Cocktail (Sigma-Aldrich)). Total protein extracts (50 μg) were separated by 10% SDS-PAGE supplemented with 1 mg/mL substrate (His-MPK5KR or His-MPK5AEF) in a resolving gel. After electrophoresis, SDS was removed from the gel by thorough washing in SDS removal buffer (25 mM Tris, pH 7.6, 0.5 mM DTT, 0.1 mM Na3VO4, 5 mM NaF, 0.5 mg/mL BSA, and 0.1% Triton X-100). Then, the native kinases were renatured in gel by incubation at 4°C for 20 h with five changes of renaturation buffer (25 mM Tris, pH 7.6, 1 mM DTT, 0.1 mM Na3VO4, and 5 mM NaF). After renaturation, gels were soaked in cold CDPK kinase buffer (25 mM Tris, pH 7.6, 12 mM MgCl2, 1 mM DTT, and 0.1 mM Na3VO4) for 10 min, and then the cold CDPK kinase buffer was replaced with fresh CDPK kinase buffer containing 200 mM ATP and 2 μCi/mL [γ-32P]ATP (MP Biomedicals) to start the kinase reaction. The CDPK kinase buffer contained 2 mM CaCl2 or 5 mM EGTA, as indicated in Figure 2C. The reaction was performed at room temperature for 1 h and stopped by replacing kinase buffer with gel washing buffer (5% trichloroacetic acid and 1% sodium pyrophosphate) and then washed thoroughly to remove free radioactive ATP. The gel was stained with Coomassie Brilliant Blue R 250 and vacuum dried to 3MM filter paper. The autoradiographic detection was performed using the Storage Phosphor screen and Storm 820 scanner (GE).

For the in-gel kinase assay to detect MAPK activity, total protein (30 μg) or immunoprecipitated MPK5 was separated on 10% SDS-PAGE gels containing 0.25 mg/mL MBP (Life Technologies). The immunoprecipitation...
and MAPK in-gel kinase assay were performed as described previously by Xiong and Yang (2003).

**In Vitro GST Pull Down**

GST-CPK18 and GST (5 µg) were mixed with 10 µg of His-OsMAPK5 in 0.5 mL of binding buffer (25 mM Tris, pH 7.5, 150 mM NaCl, and 1% Triton X-100). Then, 20 µL of equilibrated Glutathione Sepharose 4B (GE) was added. After incubation at 25°C for 2 h, the Glutathione Sepharose was washed five times with binding buffer and resuspended in 0.2 mL of SDS sample buffer. The pull-down product was analyzed by immunoblotting using anti-MPK5 and anti-GST (Sigma-Aldrich) antibodies.

**RT-qPCR**

Total RNAs were extracted from rice tissues and treated with DNase I (New England Biolabs) before reverse transcription. The cDNA was synthesized using the Multiscript Reverse Transcription Kit (Life Technologies). Real-time PCR was performed in StepOnePlus (Life Technologies) using the GoTag qPCR Master Mix (Promega). The relative quantification of gene expression was calculated as described previously (Kino-shita-Kikuta et al., 2007)."
**Supplemental Figure 6.** Mapping CPK18 Phosphorylated Sites on MPK5.

**Supplemental Figure 7.** Native CPK18 Activities in Wild-Type and MPK5-RI Lines.

**Supplemental Table 1.** List of Genes and DNA Oligonucleotides Used in This Study.

**Supplemental Data Set 1.** Sequence Alignment of MAPKs to Construct the Phylogenetic Tree in Supplemental Figure 1.

**Supplemental Data Set 2.** Sequence Alignment of CDPKs to Construct the Phylogenetic Tree in Supplemental Figure 2.

**ACKNOWLEDGMENTS**

We thank the late Ko Shimamoto at the Nara Institute of Science and Technology and Tsuyoshi Nakagawa at Shimane University for providing the pANDA and pUGW vectors, respectively, and Shuqun Zhang at the University of California San Diego for carefully reviewing the article and the Genomics Core Facility at Pennsylvania State University for providing the DNA sequencing service. This work was supported by the USDA/NIFA (Grant 2008-35301-19028 to Y.Y.) and the National Science Foundation Plant Genome Research Program (Grant DBI-0922747 to Y.Y.).

**AUTHOR CONTRIBUTIONS**

K.X., J.C., and Y.Y. designed the research. K.X., J.C., and Q.W. performed the experiments. K.X., J.C., and Y.Y. analyzed the data. K.X., J.C., and Q.W. designed the research. K.X., J.C., and Q.W. wrote the article.

Received April 9, 2014; revised May 18, 2014; accepted June 16, 2014; published July 17, 2014.

**REFERENCES**


