Poaceae-specific cell wall-derived oligosaccharides activate plant immunity via OsCERK1 during Magnaporthe oryzae infection in rice

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Many phytopathogens secrete cell wall degradation enzymes (CWDEs) to damage host cells and facilitate colonization. As the major components of the plant cell wall, cellulose and hemicellulose are the targets of CWDEs. Damaged plant cells often release damage-associated molecular patterns (DAMPs) to trigger plant immune responses. Here, we establish that the fungal pathogen Magnaporthe oryzae secretes the endoglucanases MoCel12A and MoCel12B during infection of rice (Oryza sativa). These endoglucanases target hemicellulose of the rice cell wall and release two specific oligosaccharides, namely the trisaccharide 31-β-D-Cellobiosyl-glucose and the tetrasaccharide 31-β-D-Cellotriosyl-glucose. 31-β-D-Cellobiosyl-glucose and 31-β-D-Cellotriosyl-glucose bind the immune receptor OsCERK1 but not the chitin binding protein OsCEBiP. However, they induce the dimerization of OsCERK1 and OsCEBiP. In addition, these Poaceae cell wall-specific oligosaccharides trigger a burst of reactive oxygen species (ROS) that is largely compromised in oscerk1 and oscebip mutants. We conclude that 31-β-D-Cellobiosyl-glucose and 31-β-D-Cellotriosyl-glucose are specific DAMPs released from the hemicellulose of rice cell wall, which are perceived by an OsCERK1 and OsCEBiP immune complex during M. oryzae infection in rice.
Plants live in an environment that is exposed to diverse microorganisms, many of which are pathogenic. Phytopathogens typically carry conserved features known as pathogen-associated molecular patterns (PAMPs), including flagellin, elongation factor-Tu (EF-Tu), and peptidoglycan in bacterial pathogens, and chitin in fungal pathogens. Plants have evolved an elegant immune system to recognize these non-self PAMPs and to activate immune responses as a defense against pathogen invasion. Cell surface-localized pattern recognition receptors (PRRs) perceive PAMPs and initiate PAMP-triggered immunity (PTI), which involves a burst of reactive oxygen species (ROS), Ca^{2+} influx, mitogen-activated protein kinase (MAPK) activation, callose deposition, and the expression of defense-related genes.

PRRs are either plasma membrane-localized receptor-like kinases (RLKs) or receptor-like proteins, functioning as the first line of plant defense against pathogen infection. RLKs are typically composed of an extracellular domain, a transmembrane domain, and a cytosolic kinase domain. RLK extracellular domains are variable ectodomains featuring leucine-rich repeats (LRRs), the lysin motif (LysM), or lectin-type motifs, and are essential for PAMP recognition specificity. Two well-characterized PRRs with extracellular domains are FLAGELLIN-SENSING 2 (FLS2) and EF-Tu receptor (EFR), which perceive bacterial flagellin (or the epitope flag22) and EF-Tu (or the epitope efl18), respectively. Different plant species may have different PAMP perception modules. In Arabidopsis thaliana, the LysM-RLK ATLYK5 coordinates with another LysM-RLK, CERK1, to perceive chitin and mount immune responses. In contrast, in rice (Oryza sativa), OsCERK1 forms a heterodimer with OsCEBiP to perceive chitin.

Pathogen infection often damages host cells, resulting in the production of plant-derived damage-associated molecular patterns (DAMPs). These DAMPs function as “danger” alert signals and can activate plant immune responses. The term DAMP was originally coined in studies of mammalian immunity and was subsequently extended to the studies of plant immune systems. Currently identified plant DAMPs include proteins, carbohydrates, lipids, and nucleotides. The Arabidopsis AtPEPs are well-established peptide DAMPs that are recognized by PEP RECEPTOR1/2, a plasma membrane-localized LRR-RLK. PAMP-induced secreted peptides are another family of protein-derived elicitor peptides, and are suggested to be perceived by RLK7 in Arabidopsis. In addition to peptide DAMPs, ATP has been found to trigger immune responses in both animal and plant cells. Extracellular ATP (eATP) is released into the plant apoplast during cell damage and can be perceived by the PRR DORN1/LecRK-19, a lectin RLK.

The plant cell wall is comprised of a complex interconnected mixture of proteins and polysaccharides, in which cellulose microfibrils are cross-linked to the matrix polysaccharides, hemicellulose, and pectin. Pectin-derived oligogalacturonides (OGs) are known as a major DAMP from plant cell wall. Although cellulose and hemicellulose are the most abundant polysaccharides in the plant cell wall, the potential functions of cell wall-derived cellobioamylases in plant immunity are poorly understood. Many phytopathogen genomes harbor genes that encode cell wall-degradation enzymes (CWDEs), such as polygalacturonases, cellulase, and hemicellulase, which help to break down the plant cell wall during infection. Recently, the oligomers derived from cellulose are found to be perceived as DAMPs in Arabidopsis, but they cannot stimulate ROS burst and callose deposition; instead, they induce calcium influx and defense gene expression. The cellobioamylases from fungal cell walls could also trigger plant immune responses. Interestingly, fungal cell wall-derived cellobioamylases-triggered immunity largely depends on CERK1 or operates synergistically with chitin, suggesting the essential role of CERK1 in perceiving cellobioamylases.

M. oryzae is an economically important fungus that causes up to 30% annual losses in cereal yields. This fungus is predicted to secrete ~100 cellulose-depolymerization enzymes during infection on rice. However, whether M. oryzae infection results in the release of DAMPs from rice cell wall is unclear. In this study, we demonstrate that M. oryzae secretes glycosyl hydrolases family GH12 endoglucanase MoCell12A/B to degrade rice cell walls, leading to the release of specific oligosaccharides, namely the trisaccharide 3′-β-D-cellulobioyl-glucose (G4G3G) and tetrasaccharide 3′-β-D-cellotriosyl-glucose (G4G4G3G). These specific oligosaccharides can activate immune responses in rice plants. Notably, these oligosaccharides contain a β-1,3,1,4-glucan backbone, which derives from hemicellulose and primarily presents in Poaceae species. We found that G4G3G and G4G4G3G are perceived by the plasma membrane-localized LysM-RLK OsCERK1 in rice. Thus, we reveal a set of non-cellulosic, polysaccharide-derived oligosaccharide DAMPs that can activate rice immune responses during M. oryzae infection.

Results

MoCell12A and MoCell12B expression is induced during M. oryzae infection. Ten microbial cellulosytic glycosyl hydrolase (GH) families that catalyze the degradation of plant cell wall cellulose have been characterized to date. During M. oryzae genome, three homologous GH12 endoglucanase-encoded genes, namely MoCell12A, MoCell12B, and MoCell12C, were identified by Takeda et al. (Supplementary Fig. 1a). MoCell12A, MoCell12B, and MoCell12C are the only members of GH12 family in M. oryzae genome, where MoCell12A shares a similarity of 43.6% with MoCell12B and 32.4% with MoCell12C. MoCell12C transcription was significantly induced during early M. oryzae infection, as revealed by transcriptome assays (Supplementary Data 1). This sparked our interest because CWDEs function to break down host cell walls and are considered to proceed necrotrophic growth generally; however, these genes were induced during early infection (by 24 h post inoculation, hpi), a stage of biotrophic growth. To validate the transcriptome data, we examined MoCell12A, MoCell12B, and MoCell12C expression levels at several key infection stages. MoCell12A was induced by 8 hpi (Fig. 1a), the time during primary infection hyphae formed. MoCell12A expression peaked at 24 hpi and then decreased, and was hardly detectable in the mycelium and spores (Fig. 1a). Similarly, MoCell12B expression was elevated during the early infection stages (Fig. 1a). By contrast, MoCell12C expression was relatively high in spores and the infection hyphae at 8 hpi when compared to the expression levels in the mycelium and the infection hyphae of later infection stage, suggesting that MoCell12C was also involved in infection or pathogen growth.

MoCell12A and MoCell12B are secreted β-glucanases. Since MoCell12A, MoCell12B, and MoCell12C are predicted to be endoglucanases, we investigated their function during M. oryzae infection. We expressed the recombinant proteins fused to a 6×His epitope in Pichia pastoris (Supplementary Fig. 1c) and then investigated the enzymatic specificity by performing enzymatic activity assays using barley β-glucan and tamarind seed xyloglucan as substrates, respectively. As expected, MoCell12A and MoCell12B hydrolyzed barley β-glucan, specifically the β-1,3,1,4-glucan polymers, with MoCell12A showing much higher activity than MoCell12B (Fig. 1b). We also expressed MoCell12A in M. oryzae fused to a GFP epitope, and compared their enzymatic activity with MoCell12A-His purified from P. pastoris. The result showed that their enzymatic activities were comparable.
given that MoCel12A-GFP molecular weight is twice that of MoCel12A-His (Supplementary Fig. 1d). In addition, MoCel12A did not prefer carboxymethyl cellulose (CMC), microcrystalline cellulose (MCC), or chitin as substrates (Supplementary Fig. 1e, f). By contrast, MoCel12C showed strong enzymatic activity for xyloglucan, specifically the polysaccharides of β-1,4-linked D-glucan substituted with xylose, indicating that it is a xyloglucanase. As xyloglucan is not the primary matrix polysaccharide in rice cell walls, we focused on the function of MoCel12A/B hereafter. Okada et al. reported that the key residues for Trichoderma reesei Cel12A endoglucanase activity are E116 and E200, which are conserved in family-12 cellulases (Supplementary Fig. 1g)34. We, therefore, targeted the corresponding residues in MoCel12A with the mutations E153D and E238D, and a subsequent enzymatic activity assay showed that these residue substitutions completely abolished enzyme activity, indicating that E153 and E238 are critical residues for MoCel12A function (Fig. 1b).

MoCel12A/B are putative CWDEs that are assumed to target the plant cell wall for degradation29. Thus, these proteins should be secreted to the host apoplast during infection. We created a version of MoCel12A with a C-terminal GFP tag that was expressed in M. oryzae strain Guy11, and confirmed that it was secreted into the supernatant; however, removal of the signal peptide caused MoCel12A-GFP to remain in the pellet, suggesting that MoCel12A is a secreted protein (Supplementary Fig. 2a). In addition, we expressed MoCel12A in rice plants alongside a truncated MoCel12A version lacking the signal peptide. Both proteins were present in a crude protein extract of rice seedlings, but only MoCel12A was detected in the rice apoplastic fluids (Supplementary Fig. 2b). These results further indicate that MoCel12A is a secreted protein and that MoCel12B, by association, is also a secreted protein.

MoCel12A/B negatively contribute to M. oryzae pathogenesis. Since MoCel12A/B were shown to be induced during infection, we characterized their roles in fungal growth and development. Using a recombinant exchange technique, we created a M. oryzae MoCel12A knockout mutant (ΔMoCel12A; Supplementary Fig. 3a), which was analyzed alongside a MoCel12A overexpression (MoCel12A-OE) strain. In both of these strains, we observed normal growth and a normal cell wall stress response induced by treatment with Congo red and SDS (Supplementary Fig. 3b).
To further investigate the roles of MoCel12A and MoCel12B in *M. oryzae* pathogenesis, we created a MoCel12A/B double knockout mutant (ΔMoCel12A/B; Supplementary Fig. 5c). Analysis of cell wall stress and spore germination in the WT and ΔMoCel12A/B revealed comparable colony morphology and appressorium formation (Supplementary Fig. 3d, e). To examine their pathogenesis, rice plants were inoculated with ΔMoCel12A and ΔMoCel12A/B conidia. Compared to Guy11, ΔMoCel12A/B strain showed similar pathogenesis based on the resulting fungal biomass in infected plants (Fig. 1c); however, ΔMoCel12A/B double mutant exhibited enhanced pathogenicity compared to the WT, revealed by the more severe disease symptoms and greater fungal biomass (Fig. 1d). By contrast, MoCel12A-OE overexpressing strain displayed markedly impaired pathogenicity and did not efficiently cause infection in rice (Fig. 1e). Consistently, the mutant strains complemented with the native promoter-driven MoCel12A and MoCel12B restored the pathogenesis as that of wild-type strain (Supplementary Fig. 4a, b). These results imply that MoCel12A and/or MoCel12B negatively contribute to pathogen pathogenicity.

**Ecotopic expression of MoCel12A in rice enhances disease resistance.** Since MoCel12A and MoCel12B are secreted proteins, and MoCel12A exhibited considerably higher hydrolase activity than MoCel12B (Fig. 1b), we expressed MoCel12A in rice plants (Supplementary Fig. 5a) and explored whether expression of MoCel12A could enhance plant susceptibility to *M. oryzae* infection. Transgenic expression of MoCel12A in rice resulted in a dwarf phenotype (Fig. 2a). In addition, MoCel12A expression resulted in spontaneous lesions on the leaves of transgenic plants (Fig. 2a). By contrast, transgenic expression of MoCel12A-2ED (carrying the E153D and E238D mutations, an enzymatic activity inactive mutant) (Supplementary Fig. 5b) did not visibly affect plant growth (Fig. 2b), indicating that the phenotype was attributed to the enzymatic function of MoCel12A in Fig. 2a. Trypan-blue staining confirmed that leaf lesions were caused due to cell death (Fig. 2c), indicating a constitutive immune activation in the transgenic plants.

To verify the phenotype of MoCel12A-expressing plants, we examined the expression levels of several immune-responsive genes. Indeed, the expression levels of OsRbohA, OsRbohD, OsPR3, and OsPR10 were significantly upregulated in MoCel12A-expressing plants relative to the wild-type plants, but not in MoCel12A-2ED-expressing plants (Fig. 2d). Next, we inoculated the transgenic plants with *M. oryzae* spores to assess their disease resistance. Consistently, MoCel12A-expressing plants displayed enhanced disease resistance compared to WT plants (Fig. 2e). By contrast, plants expressing inactive MoCel12A-2ED exhibited similar disease symptoms and fungal biomass as WT plants following infection (Fig. 2f), implying that the transgenic expression of MoCel12A in rice plants activates immune responses, resulting in enhanced disease resistance to *M. oryzae*.

**MoCel12A/B-hydrolyzed rice cell wall activates an immune response.** MoCel12A-activated immune response in rice could be associated with MoCel12A as a PAMP or with MoCel12A-released oligosaccharides from the rice cell wall as DAMPs. In addition, MoCel12A-released fungal cell wall components may serve as DAMPs. As MoCel12A-2ED-expressing plants did not display an autoimmune phenotype (Fig. 2b) and β-1,3-1,4-glucan barely exists in fungal cell walls, the scenario of rice cell wall-derived DAMPs seems more likely. To test this hypothesis, we first examined whether MoCel12A activated immune responses by measuring ROS bursts in rice-suspension cells. Indeed, neither the recombinant MoCel12A nor MoCel12A/E153D/MoCel12A/E238D proteins triggered a ROS burst (Fig. 3a). Similar results were observed using the recombinant MoCel12B and MoCel12C (Supplementary Fig. 5c), indicating that MoCel12s are not PAMPs. In addition, MoCel12A-digested *M. oryzae* cell walls did not trigger immune responses in rice cells (Supplementary Fig. 5d). This result excluded the fungal cell wall-derived oligosaccharides as PAMPs. We then incubated MoCel12A with isolated rice cell walls and tested the ROS burst using the cell wall extract. The extract from MoCel12A-digested cell wall strongly activated a ROS burst, whereas extracts from inactive MoCel12A/E153D/MoCel12A/E238D digests did not (Fig. 3b). In addition, MoCel12A digestion resulted in a stronger ROS burst than MoCel12B digestion, whereas MoCel12C digestion did not trigger a ROS burst (Fig. 3c), suggesting that MoCel12A/B hydrolyze rice cell walls, thus releasing oligosaccharides to activate the immune response.

We validated the above results by examining MAPK activation and immune-responsive gene expression. MoCel12A-digested cell wall extract induced a strong MAPK activation in rice cells, whereas MoCel12A/E153D/MoCel12A/E238D digestion did not (Fig. 3d). In addition, both MoCel12A and MoCel12B digestion induced MAPK activation, whereas MoCel12C digestion did not (Fig. 3e). Similar to MAPK activation, the expression of PTI marker genes PBZ1 and PAL was markedly upregulated in rice cells exposed to MoCel12A-digested cell wall extract, but not in those digested with MoCel12A/E153D/MoCel12A/E238D (Fig. 3f). Moreover, MoCel12A- and MoCel12B-digested cell wall extracts induced PBZ1 and PAL transcription, whereas MoCel12C digestion did not (Fig. 3g). These results further indicate that specific oligosaccharides released from rice cell wall by MoCel12A and MoCel12B activate plant immune responses.

**MoCel12A/B release specific oligosaccharides from rice cell wall.** We subsequently attempted to identify the oligosaccharides released by MoCel12A- and MoCel12B-hydrolyzed rice cell walls. Based on the characteristic of MoCel12A enzyme which cleaves the β-1,4 bond in the sequence of Glc-(1,3)-Glc-(1,4)-Glc (−G3G4G−), and the results from MALDI-TOF-MS analysis, we established that a trisaccharide and a tetrasaccharide are probably the two major carbohydrates released by MoCel12A- and MoCel12B-hydrolyzed rice cell wall (Fig. 4a). To further verify the oligosaccharides, we used MoCel12A to hydrolyze the highly purified barley β-glucan which is a typical component of Poacea-specific hemicellulose and is composed of polymerized β-glucose embedded with β-1,3-1,4-glucan, and examined the molecules produced. We confirmed that MoCel12A-released oligosaccharides from barley β-glucan triggered a strong ROS burst (Supplementary Fig. 5e). As a negative control, maltose-binding protein (MBP)-incubated barley β-glucan did not trigger a ROS burst. These results suggest that the specific oligosaccharides from barley β-glucan indeed are able to activate plant immune responses. The data from MALDI-TOF-MS analysis further showed that, similar to the oligosaccharides from rice cell wall, there are two oligosaccharides, a trisaccharide, and a tetrasaccharide, responsible for the immune activation in rice (Supplementary Fig. 5f).

To characterize these specific oligosaccharides, we collected all the combinations of commercial products of trisaccharide and tetrasaccharide hydrolyzed from barley β-glucan with specific attention on the oligosaccharides containing β-1,3,1,4-glycosidic linkage. Although we hypothesized that 3′-β-D-celllobiosyl-glucose (BGTRIB, G4G3G) and 3′-β-D-cellotriosyl-glucose (BGTETB, G4G4G3G) were likely the two major oligosaccharides, as MoCel12A/B produce oligosaccharides with β-1,3 bonds next to the reducing end. The polymerized β-1,3,1,4-glucan could

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theoretically be hydrolyzed to several oligosaccharides by other members of the GH family. We thus examined whether 32-β-D-glucosyl-cellobiose (BGTRIA, G3G4G) and a mixture of 32-β-D-cellobiosyl-cellobiose and 33-β-D-glucosyl-cellotriose (BGTETC, G4G3G4G/G3G4G4G) could activate immune responses in rice cells. In addition, we also assessed the capability of cellotriose (CTR, G4G4G) and cellotetrose (CTE, G4G4G4G), which are based on the β-1,4-glucan backbone and can be generated from cellulose, in activating immune responses. We first verified these oligosaccharides by MALDI-TOF-MS analysis (Supplementary Fig. 6a). The trisaccharides BGTRIA and BGTRIB activated strong ROS bursts, similar to those observed in the presence of chitin, whereas CTR did not activate a ROS burst (Fig. 4b). Similarly, the tetrasaccharide BGTETB also activated a ROS burst and BGTETC activated an even stronger response than chitin, whereas CTE did not (Fig. 4c), revealing that the basic backbones of G3G4G or G4G3G are determining factors in activating ROS burst. These trisaccharide- and tetrasaccharide-triggered ROS bursts were dose-dependent (Supplementary Fig. 6b, c). In addition to the ROS assays, MAPK activation and immune-responsive gene expression assays further demonstrated that BGTRIB and BGTETB activate PTI responses in rice (Fig. 4d–f).
Similar to PAMPs, pretreatment with DAMPs can prime plant disease resistance\(^3\). We thus tested the priming effects of BGTRIB and BGTETC on disease resistance in rice. The result showed that BGTRIB and BGTETC pretreatment enhanced plant disease resistance, exhibiting similar effects as chitin in priming defense (Fig. 4g).

We then determined the actual content of BGTRIB and BGTETB in the rice apoplastic fluid during \(M. \text{o}ryzae\) infection. Using a high-resolution triple-quadrupole mass spectrometer, we characterized the parental and daughter M/Z as 527.3/365.1 and 689.4/347.2 for BGTRIB and BGTETB, respectively (Supplementary Fig. 7a). Based on the parameters, we were able to identify BGTRIB and BGTETB in \(M. \text{o}ryzae\)-infected rice apoplastic fluids (Supplementary Fig. 7b). \(M. \text{o}ryzae\) WT infection increased BGTRIB content about twofold, while the \(\text{MoCel12A}\)-OE strain infection resulted in over fourfold increase in BGTRIB; however, BGTETB could be only detected in \(\text{MoCel12A}\)-OE strain infection, which is about twofold increase than that of Mock treatment (Supplementary Fig. 7c). These results indicate that BGTRIB and BGTETB are DAMPs released during \(M. \text{o}ryzae\) infection. We were unable to detect BGTETC during the infection, probably because BGTETC either was not produced or the content was too low to be detected.

**Fig. 3** MoCel12A-hydrolyzed rice cell wall activates an immune response in rice. a MoCel12A and its variant proteins cannot active ROS burst in rice-suspension cells. Recombinant MoCel12A-His proteins were purified from \(Pichia \text{pastoris}\), and 2 \(\mu\)g/ml protein each was used for the ROS burst assay. Chitin (1 \(\mu\)g/ml) served as a positive control. MBP served as a negative control. Values are means ± SD (\(n=4\) biological replicates). RLU relative light units. b MoCel12A-hydrolyzed rice cell wall activated the ROS burst in rice. Aliquots (1 ml) of a 50 mg/ml suspension of isolated rice cell walls in 50 mM sodium acetate buffer, pH 5.5, were incubated for 2 h at 37 °C with 5 \(\mu\)g recombinant MBP-His, MoCel12A-His, MoCel12A(E153D)-His, and MoCel12A(E238D)-His, respectively. After incubation, the supernatants (1/100, v/v) were used for the ROS burst assay. CW cell wall. Others are as in (a). c MoCel12A and MoCel12B but not MoCel12C hydrolyzed rice cell wall activated the ROS burst. The suspension of isolated rice cell walls was incubated with 5 \(\mu\)g recombinant MBP-His, MoCel12A-His, MoCel12B-His, and MoCel12C-His, respectively. Others are as in (b). d MoCel12A-hydrolyzed rice cell wall activated MAP kinase cascades. The supernatants of the different proteins-hydrolyzed rice cell wall were incubated with rice-suspension cells. Activated MAPKs were detected by immunoblotting with the phospho-p38 MAPK antibody at the indicated time points. The corresponding bands indicate the phosphorylation of MPK3 and MPK6. Coomassie brilliant blue (CBB) staining of Rubisco indicates equal loading of samples in each lane. The experiment was repeated three times with similar results. e MoCel12A and MoCel12B but not MoCel12C hydrolyzed rice cell wall activated MAP kinase cascades in rice. Others are as in (d). f MoCel12A-hydrolyzed rice cell wall induced PTI marker gene expression. Treatments with the hydrolyzed rice cell wall and rice-suspension cells were same as in (d), and the samples were collected for RT-qPCR at the indicated time points. Values are means ± SD (\(n=3\) biological replicates). g MoCel12A and MoCel12B but not MoCel12C hydrolyzed rice cell wall induced PTI marker gene expression. Others are as in (f).
Fig. 4 The specific trisaccharides and tetrasaccharides released by MoCel12A from rice cell wall activate the immune response in rice. 

**a** The oligosaccharides of MoCel12A, MoCel12B, and MoCel12C hydrolyzed cell wall detected by MALDI-TOF-MS. The filtered supernatants of MoCels-hydrolyzed rice cell walls were used for the analysis, using 10 mg/ml 2,5-dihydroxybenzoic acid in 2:1 acetonitrile:water as the matrix. All indicated peaks are single-charged ions of oligosaccharides on the reducing end with Na\(^+\) using 3-aminoquinoline (3-AMQ) as ionic liquid matrix.

**b** The trisaccharides activate ROS burst in rice-suspension cells. 10\(\mu\)M each of CTR (cellotriose, G4G4G), BGTRIA (3\(^{2}\)-\(\beta\)-D-glucosyl-cellobiose, G3G4G), and BGTRIB (3\(^{1}\)-\(\beta\)-D-cellobiosyl-glucose, G4G3G) were used for the ROS burst assays. Chitin (10\(\mu\)g/ml) served as the positive control.

**c** The tetrasaccharides activate ROS burst in rice. CTE (cellotetraose, G4G4G4G), BGTETB (3\(^{1}\)-\(\beta\)-D-cellotriosyl-glucose, G4G4G3G), and BGTETC (3\(^{2}\)-\(\beta\)-D-cellobiosyl-cellobiose + 3\(^{3}\)-\(\beta\)-D-glucosyl-cellotriose, G4G3G4G + G3G4G4G) were used for the ROS burst assays (10\(\mu\)M each). Others are as in (b).

**d** The trisaccharides and tetrasaccharides activated MAP kinase cascades in rice. The rice-suspension cells were treated with oligosaccharides (10\(\mu\)M) and were used for MAPK activation assays. Treatments with oligosaccharides and rice-suspension cells were same as in (d), and the samples were collected for RT-qPCR at the indicated time points. Values are means ± SD (n = 3 biological replicates).

**e** Pretreatment with BGTRIB and BGTETC enhanced the blast disease resistance in rice. Two-week-old rice plants were treated with 50\(\mu\)g/ml chitin, 100\(\mu\)M BGTRIB and BGTETC for 3 h, then the leaves were spray-inoculated with conidial suspensions (1 \times 10^5 conidia per ml in 0.02% Tween-20) for 4 d. Bar = 1 cm. The right panel is the relative fungal biomass. Values are means ± SD (n = 3 biological replicates). * indicates significant differences from the Mock at \(P<0.05\) (two-sided Student’s t test).
BGTRIB- and BGTETC-triggered immune responses are compromised in oscerk1. Because BGTRIB and BGTETC could trigger the ROS burst in rice plants, we next sought to identify the LysM-RLK(s) which are known to perceive carbohydrates or oligosaccharides. In a screen of LysM-RLK RNAi rice-suspension cells and knockout cells, we found that BGTRIB- and BGTETC-triggered ROS bursts, as well as ROS bursts resulting from chitin, were almost undetectable in the oscerk1 mutant (Fig. 5a). These results suggest that OsCERK1 perceives BGTRIB and BGTETC, thereby activating immune responses. In addition, we discovered that BGTRIB and BGTETC-triggered ROS bursts were partially compromised in osc bib mutant plants (Fig. 5b). Moreover, we found that OsRac1 was involved in the BGTRIB- and BGTETC-triggered ROS burst using an OsRac1-RNAi line (Supplementary Fig. 8a), which showed compromised responses to treatment with chitin and the two oligosaccharides (Fig. 5c). Notably, LYP4/6 was reported to mediate chitin recognition46, but the mutant plants did not exhibit compromised ROS bursts following treatment with BGTRIB and BGTETC (Supplementary Fig. 8b). In addition, as OsCERK1 associates with multiple PRRs but does not seem to be involved in plant immunity47, we used it as a negative control. The result showed that the oscerk1 mutant did not exhibit compromised ROS bursts following BGTRIB and BGTETC treatments (Supplementary Fig. 8c). These results suggest that OsCERK1 specifically perceives BGTRIB and BGTETC and that OsRac1 further transduces these signals and activates immune responses in rice.

As OsCERK1 is a typical PRR, we examined the PTI response in the oscerk1 suspension cells treated with BGTRIB and BGTETC. The result shows that MAPK activation was markedly reduced (Supplementary Fig. 8d). Similarly, BGTRIB- and BGTETC-induced PBZ1 and PAL transcription was abolished in the oscerk1 mutant (Supplementary Fig. 8e), further indicating that OsCERK1 mediates BGTRIB and BGTETC recognition. To reinforce this conclusion, we pretreated the WT rice plants and the oscerk1 mutant with BGTRIB and BGTETC. We found that pretreatment with the oligosaccharides increased disease resistance in WT plants but not in the oscerk1 mutant (Supplementary Fig. 8f), implying that the oscerk1 mutant was compromised in the perception of these specific oligosaccharides. This conclusion was further confirmed by inoculating the oscerk1 mutant with MoCel12A overexpression strain, where we observed that overexpressing MoCel12A reduced the pathogenicity in WT plants but enhanced the pathogenicity in the mutants (Supplementary Fig. 8g).

To examine the recognition specificity of OsCERK1 for oligosaccharides, we also tested cellobiose, laminaritriose (LAM3), and xylotriose (XTR) in the activation of the ROS burst. The result showed that cellobiose and LAM3 did not trigger ROS burst in both WT and the oscerk1 mutant; however, XTR triggered a ROS burst in WT but not oscerk1, indicating that OsCERK1 also recognizes XTR (Supplementary Fig. 9a). In addition, we examined whether Arabidopsis CERK1 functions similarly to OsCERK1. Notably, all the tested oligosaccharides except BGTETB activated a weak ROS burst and this activation was independent of CERK1 in Arabidopsis (Supplementary Fig. 9b). Furthermore, MoCel12A-digested Arabidopsis cell wall extracts could not trigger immune responses in rice (Supplementary Fig. 9c). These results demonstrate that rice cell wall-derived oligosaccharides are specific and are perceived by OsCERK1.

BGTRIB and BGTETC bind to and activate OsCERK1. Because OsCERK1 perceives BGTRIB and BGTETC, it may directly bind to these oligosaccharides. We, therefore, expressed the ectodomain of OsCERK1 (OsCERK1-ECD) in insect cells and purified the recombinant protein for binding assays according to the method of Liu et al.48. Since OsCEBiP binds to chitin and forms a heterodimer with OsCERK1, we also expressed OsCEBiP-ECD using the same method48. Microscale thermophoresis (MST) assays showed that OsCERK1-ECD bound to BGTETB, BGTRIB, and BGTETC (Fig. 5d). OsCERK1-ECD also bound to BGTRIA and the β-1,4-glucose-based CTR and CTE; however, it did not bind to the disaccharide sucrose and the monosaccharide glucose (Supplementary Fig. 9d). Although OsCEBiP bound to chitin, it did not bind to BGTETB, BGTETB, and BGTETC (Fig. 5e). These results demonstrate that OsCERK1 binds to trisaccharides and tetrasaccharides, but not to disaccharides and monosaccharides.

In rice, chitin induces OsCERK1 and OsCEBiP to form a receptor complex, which transduces the chitin signal to downstream components for immune activation30,39,40. Therefore, we examined whether BGTRIB or BGTETC could induce OsCERK1 and OsCEBiP to form a receptor complex. Similar to chitin, the MoCel12A-digested rice cell extracts, BGTRIB, and BGTETC induced OsCERK1 and OsCEBiP to form a heterodimer (Fig. 5f), suggesting that these oligosaccharides share a similar immune response signaling pathway as chitin. In Arabidopsis, chitin binds to the LysM-RLK AtLYK5 and induces the formation of a complex with AtCERK1, thus inducing AtCERK1 phosphorylation and leading to immune activation49. We also observed that BGTRIB and BGTETC induced OsCERK1 homodimerization (Fig. 5g). Notably, CTR did not induce the dimerization of OsCERK1 (Supplementary Fig. 9e). These results imply that cell wall-derived specific oligosaccharides likely induce OsCERK1 and OsCEBiP tetramers to transduce immune signaling.

Discussion

DAMP-activated plant immune responses are an important branch of plant innate immunity1,40,41. The exogenous application of many DAMPs, such as eATP or cutin monomers, triggers typical PTI responses17,42. So far, many DAMPs have been identified, including proteins/peptides and carbohydrates41. DAMPs are presumably recognized by plasma membrane-localized PRRs, but very few of the DAMP-associated PRRs have been identified1,23,41.

Cellulose is the major component of the plant cell wall, and is formed by the polymerization of β-1,4-glucan43. In dicots, the non-cellulosic polysaccharides also contain a β-1,4-glucan backbone. However, in most cereal crop species, the non-cellulosic polysaccharides are heteroxylans and β-1,3-1,4-glucans20, which are specific to grass plants. β-1,4-glucose-based cellulose is capable of triggering calcium influx in plant cells and inducing defense gene expression44, as well as a weak ROS burst in Arabidopsis40. However, cellulose induced a strong defense gene expression44. In addition, the β-1,4-glucoside and linear β-1,3-glucan can also induce ROS burst and calcium influx in grapevine cells45. These lines of evidence suggest that cellulose or cellobiose is a potent DAMP. However, it is unknown whether these oligosaccharides are generated during pathogen infection. A conserved GH17 glycosyl hydrolase from the tomato pathogen Cladosporium fulvum (CIGH17) was proposed to release a tomato cell wall DAMP that triggers host immune responses via an unknown plant receptor46. In this study, we established that the M. oryzae-secreted proteins MoCel12A and MoCel12B are members of the GH12 family of glycosyl hydrolases, which accept β-1,3-1,4-glucan as substrate. The resulting digestion products serve as DAMPs that activate plant immune responses during infection (Fig. 6). However, MoCel12A/B themselves are not PAMPs (Fig. 3a). Notably, the β-glucanase XEGI produced by the soybean (Glycine max) pathogen Phytophthora sojae and the endocellulase EG1 from Rhizoctonia solani both exhibit cellulase activity, but they have established elicitors and their enzymatic activities are not required for elicitor activity47,48.
Using a MALDI-TOF-MS and a high-resolution triple-quadrupole MS analysis approach, we identified apoplastic oligosaccharide DAMPs from *M. oryzae*-infected rice (Supplementary Fig. 7). We established that BGTETB and BGTRIB are two major oligosaccharides released from rice cell wall during *M. oryzae* infection (Fig. 4). Enzymatic activity analysis showed that MoCel12A and MoCel12B are β-glucanases and are responsible for BGTRIB and BGTETB production during infection (Figs. 1 and 4). Furthermore, we demonstrated that hemicellulose-derived oligosaccharides BGTRIB and BGTETB strongly activate OsCERK1. The rice cell wall-derived specific oligosaccharides are perceived by OsCERK1. a ROS burst induced by rice cell wall-derived specific oligosaccharides is abolished in *oscerk1* mutant. The WT and *oscerk1* suspension cells were elicited with sterile water (Mock), filtered supernatants (1/100, v/v) of MoCel12A-hydrolyzed rice cell walls (CW), 10 μM BGTRIB, 10 μM BGTETC, and 5 μg/ml chitin in ROS burst assays, respectively. flg22 (100 nM) served as a positive control. Values are means ± SD (n = 8 biological replicates). RLU, relative light units. b ROS generation induced by rice cell wall-derived specific oligosaccharides in *oscebip* mutant. Others are as in (a). c ROS generation induced by rice cell wall-derived specific oligosaccharides in OsRac1-RNAi plants. Others are as in (a). d Microscale thermophoresis (MST) assays show the binding of OsCERK1-ECD with the specific oligosaccharides. The solid curve is the fit of the data points to the standard K_d-fit function. K_d, dissociation constant. Bars represent ± SD (n = 3 biological replicates). e MST assays show that OsCEBiP-ECD does not bind the oligosaccharides. Others are as in (d). f BGTRIB and BGTETC triggered heterodimerization of OsCERK1 and OsCEBiP in rice. OsCERK1-FLAG and OsCEBiP-HA were co-expressed in WT rice protoplasts. GUS-FLAG served as a negative control. The protoplasts were treated with 5 μg/ml chitin, filtered supernatants (1/100, v/v) of MoCel12A-hydrolyzed rice cell walls (CW), 10 μM BGTRIB and 10 μM BGTETC for 15 min. Co-immunoprecipitation was performed using anti-FLAG antibody and subjected to immunoblot analysis with anti-FLAG or anti-HA antibody as indicated. The experiment was repeated three times with similar results. g BGTRIB and BGTETC triggered OsCERK1 dimerization in rice. FLAG- and HA-tagged OsCERK1 were co-expressed in WT rice protoplasts. Others are as in (f).
cell wall-derived immune responses during root-bacteria interactions. Never-
theless, whether cellulose-derived cellooligomers are perceived by
OsCERK1–OsCEBiP, which transduces the signal to downstream OsRac1 for
immune activation.

plant immune responses (Fig. 4b–f). Moreover, enhanced M. oryzae disease resistance results from pretreating rice plant with
these oligosaccharides (Fig. 4g). Therefore, BGTTETB, BGTTrib, and likely BGTTETC are DAMPs that function during M. oryzae infection in rice.

The cell wall integrity system is essential for perceiving cell wall
damage and status. Souza et al. discovered that cellobiose markedly facilitated the ability of flg22 or OGs but not chitin to induce ROS burst in Arabidopsis. Similarly, a cellobiose from the fungal cell wall induced weak ROS production in Arabidopsis roots; however, the combination of cellobiose and chitin induced much stronger ROS production than cellobiose or chitin alone, demonstrating that cellobiose functions as an amplifier for PTI. These studies reveal that PRRs are a part of the cell wall integrity system that monitors cell damage during PTI. In Arabidopsis, cell damage activates the expression of multiple PRRs, including FLS2, EFR, CERK1, RLP23, and LORE, which facilitate stronger immune responses during root-bacteria interactions. Nevertheless, whether cellulose-derived cellooligomers are perceived by a host PRR and directly activate immune responses in plants is unknown.

Recently, the non-branched β-1,3-glucan laminarin and laminaribiohexaose (β-1,3-(Glc)₆) were shown to activate strong immune responses in two Poaceae species, namely Hordeum vulgare (barley) and Brachypodium distachyon, although the concentrations used were higher than those used here for BGTTETB/C (Fig. 4). Laminarin, but not laminaribiohexaose, also activates strong immune responses in the dicot species Nicotiana benthamiana, whereas laminaribiohexaose activates weak calcium influx and defense gene expression, and these responses are largely dependent on CERK1 in Arabidopsis. However, laminarin-induced immune responses in the dicot N. benthamiana and the monocots barley and B. distachyon are independent of CERK1. In Arabidopsis, CERK1 associates with LYK5 or other lysozyme motif PRRs such as LYK4 to perceive chitin oligosaccharides. Similar to the findings by the crystal structure analysis on rice OsCERK1, Arabidopsis CERK1 showed a weak binding affinity to chitin, suggesting that OsCERK1 or CERK1 interacts with chitin receptors to mediate immune activation. Using oscerk1 mutant plants, we established that OsCERK1 is essential for the BGTTETC- and BGTTETB-activated ROS burst, MAPK activation, and defense gene expression (Fig. 5a and Supplementary Fig. 8d–f). Furthermore, MST assays showed that OsCERK1 binds to BGTTETB/C and BGTTrib in vitro and, importantly, BGTTETC and BGTTETB can induce the homodimerization of OsCERK1–OsCEBiP (Fig. 5d–g), which indicate successful recognition and immune activation. Although OsCEBiP cannot bind BGTTETB/C and BGTTrib, the oscebiP mutant was partially compromised in the immune activation (Figs. 5b, e), which is likely because BGTTETC and BGTTrib can induce its homodimerization with OsCERK1 (Fig. 5f) and OsCEBiP coordinates with OsCERK1 to perceive BGTTETB/C and BGTTrib. Notably, BGTTrib and BGTTETB activate strong immune responses and are G4G3G-based oligosaccharides, whereas BGTRIA is G3G4G-based and triggers comparably weaker immune responses, implying that G4G3G-based oligosaccharides are the more active DAMPs. Collectively, our results suggest that β-1,4-glucose-based oligosaccharides are the potential ligands of OsCERK1. Although xyloglucan is not the substrate of MoCel12A/B (Fig. 1b), a recent study showed that the arabinoxylan-oligosaccharides can serve as DAMPs in Arabidopsis, suggesting that pathogen-secreted xyloglucanase could potentially digest plant cell wall hemicellulose to release DAMPs.

M. oryzae and many other fungal pathogens secrete a diverse array of CWDEs to target host cell walls for infection. These CWDEs either target host cell walls for nutrition or destroy them to initiate cell death, which eventually leads to the switch of infection to necrotrophic stage of this pathogen. Because it is hard to distinguish the exact time course of biotrophic stage, the early expression of MoCel12A/B could help for the acquisition of nutrition or the initiation of cell death (Fig. 1a). However, the DAMPs from these events are checked by the host PRRs (Fig. 6). Because there are so many CWDEs in the fungal genome and many of them are functionally substitutable, removing one or two CWDEs generally could not significantly affect their pathogenicity. In our case, overexpression of MoCel12A largely impaired the pathogenicity in WT plants (Fig. 1e). However, overexpression of MoCel12A enhanced the fungal pathogenicity in the oscerk1 mutant, suggesting that this endoglucanase is somewhat indispensable for infection when the host cannot or is compromised to perceive the released oligosaccharides (Supplemental Fig. 8g). In fact, some PAMPs or DAMPs have been found to be modified by either host or pathogen-secreted enzymes, and the modified PAMPs or DAMPs could escape the detection by PRRs. For example, chitin oligomers and flagellin could be deacylated and deglycosylated to evade host recognition, respectively. In addition, the Arabidopsis berberine bridge enzyme-like proteins have been found to specifically oxidize the well-known DAMPs, cellulose oligomers and oligogalacturonides, and impair their elicitor activity. Whether these mechanisms exist in the interaction between M. oryzae and rice to evade OsCERK1 recognition is worth further investigation.

Taken together, we have characterized an immune mechanism in which M. oryzae infection leads to the production of specific DAMPs in the rice apoplast, which in turn activate strong immune responses in rice plants but not in Arabidopsis. These species-specific DAMPs indicate the existence of a co-evolutionary plant–pathogen relationship between M. oryzae...
and rice. Whether this specific recognition existing in other plant and pathogen interactions is worth of further investigation. In addition, how M. oryzae overcomes OsCERK1-mediated DAMPs recognition and the subsequent immune responses also need further investigation. Furthermore, it would be interesting to investigate why Arabidopsis CERK1 perceives chitin but not these specific oligosaccharides derived from rice cell walls.

**Methods**

**Plant materials and growth conditions.** Rice (Oryza sativa subsp. Japonica cv Nipponbare) was used as the wild-type (WT) plant. The MoCel12A, MoCel12A2ED, and MoCel12A-6SP (nsp, no signaling peptide) overexpression constructs were generated as described by Li et al.58. The coding sequence (CDS) amplified from rice OsCERK1 was ligated with the overexpression vector pRTN-eGFP that was digested with EcoRI and BamHI. For the complementary constructs, were generated as described by Li et al.58. The rice (Oryza sativa subsp. Japonica cv Nipponbare) was used as the wild-type (WT) plant. The knockout or RNAi lines were described previously56,57. OsCERK1-ECD or OsCEBiP-ECD to the oligosaccharides were measured by QIAGEN) according to the manufacturer’s instructions. Briefly, 1 μg chitinase and recombinant MoCel12A were included in the reaction buffer. The enzyme activity was defined as one unit that 1 μg enzyme produces 1 nmol GlcNAc in 500 μl reaction buffer at 37 °C per hour.

**Gene expression analyses.** Total RNAs were isolated from rice seedlings or suspension cells using a TRIzol reagent according to the manufacturer’s instructions (Invitrogen). cDNA was synthesized using the HiScript III RT Gene expression analyses secreted and a His6-tag at the C-terminus for purification. The recombinant baculovirus was prepared according to the manufacturer’s manual (Invitrogen). Trichoplusia ni (High Five) cells (Invitrogen) were infected with recombinant baculovirus at a multiplicity of infection of 0.5–10 pfu/cell. The secreted soluble protein was recovered using a HisTrap HP 5 ml column (GE Healthcare).

**Enzyme activity assays.** Enzyme activities of MoCel12A and chitinase were determined following the DNS method, as previously described60. The barley β- glucan and tamarind seed xyloglucan (Megazyme, Ireland), as well as CMC (carboxymethyl cellulose), MCC (microcrystalline cellulose), and chitin (Solarbio, China) were used as substrates. Chitinase from Streptomyces griseus was used as the control. The standard reaction mixture (250 μl) containing 0.1 μg protein in 50 μl PBS buffer and 1.6 mg carbohydrates in 200 μl sodium acetate buffer (100 mM, pH 5.2). The mixture was incubated for 30 min at 37 °C, and then 500 μl DNS reagent was added. The mixture was boiled for 5 min and the color intensity was determined by a spectrophotometer at 540 nm. The chitinase activity was measured by Chitinase Assay Kit (Solarbio, China) according to the manufacturer’s instructions. Briefly, 1 μg chitinase and recombinant MoCel12A were included in the reaction buffer. The enzyme activity was defined as one unit that 1 μg enzyme produces 1 nmol GlcNAc in 500 μl reaction buffer at 37 °C per hour.

**Rice cell wall hydrolysis assays.** For the rice cell wall isolation, 4-week-old rice seedlings (10–20 g) were harvested, frozen immediately in liquid nitrogen, and then were ground into powder. The material was extracted with 70% ethanol three times at 70 °C for 1 h and then dried at 50 °C to a constant weight. Aliquots (1 ml) of a 50 mg/ml suspension of the isolated rice cell wall in 50 mM sodium acetate, pH 5.5 were incubated for 2 h at 37 °C with 5 μg MoCel12A2ED per aliquot, samples were boiled for 20 min and centrifuged at 12,000×g for 30 min. The supernatant was used as the elictors.

**Detection of the oligosaccharides in hydrolyzed cell wall.** The oligosaccharides released from MoCel12A, MoCel12B, and MoCel12C hydrolyzed cell wall were detected by MALDI-TOF-MS. MALDI-TOF-MS analysis was performed on an AB Sciex 5800 MALDI-TOF/TOF mass spectrometer. Mass spectra were obtained in the positive reflection mode. The filtered products of different proteins-hydrolyzed rice cell walls (1 μl) were mixed with 1 μl of 2.5-dihydroxybenzoic acid solution (20 mg ml⁻¹ in 50:50 acetonitrile/water containing 0.1% TFA). The mixture was then spotted on the target plate with 1 μl of 5-chloro-2-mercaptobenzothiazole (10 mg ml⁻¹) and dried for MALDI-TOF-MS analysis.

**Oxidative burst and MAPK assays.** Rice suspension cells were used for ROS burst and MAPK assays. The cells were used 3–4 days after subcultivation in a fresh medium. The ROS burst and MAPK assays were described previously53. Briefly, aliquots of 500 mg cells were suspended in 5 ml pre-incubation medium (3% sucrose and 10 mM MES in 5% culture medium, pH 5.8) and incubated under culture conditions for 4–5 h. Before the assay, the medium was replaced with 200 μl fresh medium containing 25 μM DMB (Wako Chemicals, Japan) and 10 μl 10 mM heme/hemin peroxidase (Sigma), and respective elictors. The luminescence was recorded by a Centro XS3 LB 960 Luminometer (Berthold Technologies). Phospho-MPK3/6 signals were detected by a phosphorylation-specific pS83 MAPK antibody (Cell Signaling Technology; 1:200 dilution). The highly purified oligosaccharides (CTE, CTR, BGTRIA, BGTBRI, BGTFETB, BGTETC, LAM3, and XTR) were used for ROS burst and MAPK activation assays were obtained from Megazyme (Ireland).

**Microscale thermophoresis (MST) analysis.** Binding reactions of recombinant OsCERK1-ECD or OsCEBiP-ECD to the oligosaccharides were measured by microscale thermophoresis in a Monolith NT.Label-Free (Nano Temper Technologies GMBH, Germany) instrument which detects changes in size, charge, and conformation induced by binding. OsCERK1-ECD or OsCEBiP-ECD (10 μM) was displaced by a buffer containing 50 mM Tris-HCl (pH 7.4), 150 mM NaCl, 10 mM MgCl₂, and 0.05% (v/v) Tween-20. A range of concentrations of oligosaccharides in the assay buffer (50 mM Tris-HCl pH 7.8, 150 mM NaCl, 10 mM MgCl₂, 0.05% Tween-20) was incubated with labeled protein (1.1, v/v) for 10 min. The sample was loaded into the NT. Label-Free stained capillaries and measured with 20% LED power and 80% MST power. The KD Fit function of the Nano Tempe Analysis Software (Version 1.5.41) was used to fit the curve and calculate the value of the dissociation constant (KD).

**Co-immunoprecipitation assays.** Plasmid constructs were amplified by PCR and introduced into a pcDNA3 vector with a HA or FLAG epitope tag at the C-terminus and cloned into the pCAGGS vector. The overexpression proteins were transfectected with OsCERK1-FLAG and OsCERK1-HA or OsCEBiP-FLAG and OsCEBiP-HA using extraction/wash buffer (50 mM Tris-HCl, pH 7.5, 150 mM NaCl, 0.1% TritonX-100, 0.2% Nonidet P-40, 1 mM DTT, and 1× complete protease inhibitor (Roche)). Anti-FLAG antibody-conjugated agarose beads (Sigma) were added to the supernatant. After incubation at 4 °C on an end-over-end shaker for 1.5 h, the beads were spun down at 15000×g for 2 min and washed at least five times. The bound proteins were eluted by 1.5× Laemmli loading buffer and resolved by 12% SDS-PAGE. The gels were stained with Coomassie blue (Merck, Germany). The gel images were quantitated using ImageJ software (NIH, USA).

**RESULTS**

**Overexpression pattern of MoCel12A in rice cell walls.** Conidial suspensions at a concentration of 1 × 10⁵ conidia/ml in 0.02% Tween-20. Inoculated plants were placed in a growth chamber (16 h light for inoculation assays.

**Fluorometric enzymatic activity assay of MoCel12A.** Bacterial strains were grown at 37 °C with shaking at 250 rpm on LB plates. The medium was inoculated with 0.1 μl of a 10-fold dilution that was prepared using a 10-μl loop. The bacterial colonies were incubated for 18 h, and then picked and used for the experiments.

**Lytic enzyme activity on cell walls.** Rice cell wall hydrolysis assays were performed on rice cell walls. Suspension cultures of bacteria were grown in LB medium for 18 h at 37 °C and 250 rpm. The bacterial cultures were collected and washed three times with PBS (137 mM NaCl, 2.7 mM KCl, 10 mM Na₂HPO₄, 1.8 mM KH₂PO₄). The washed bacteria were resuspended in 50 mM Tris-HCl buffer, pH 7.5, containing 1% Triton X-100 and 0.1 mg/ml of DNAse I (Roche). The bacterial suspensions were incubated for 2 h at 37 °C with shaking at 250 rpm. The bacterial suspensions were collected by centrifugation at 4°C, 13,000×g for 15 min. The supernatants were used as elicitors for rice seedlings.
SDS-PAGE and then subjected to immunoblot analysis using anti-FLAG antibody (Sigma, F3165, 1:10,000 dilution) or anti-HA antibody (Huxiang bio, HX1820, 1:5000 dilution).

**Phylogenetic analyses.** The amino acid sequences of the MoCel12A homologous proteins were downloaded from the UniProt website (https://www.uniprot.org/). Sequence alignment was performed with ClustalW. A neighbor-joining method was used to generate the phylogenetic tree. The bootstrap values indicated at the nodes in the phylogenetic tree are based on 1000 replications.

**Data availability**

The authors declare that all data supporting the findings of this study are available within the paper and the supplementary files or are available from the corresponding author upon request. The RNA-seq data that support the findings of this study have been deposited in the SRA database of the National Center for Biotechnology Information with the accession code PRJNA707110. Mass spectrometry data that support the findings of this study have been deposited in MetaboLights with the accession code MTBLS2548. Source data are provided with this paper.

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**References**

1. Bournot, F. & Zipfel, C. Function, discovery, and exploitation of plant pattern recognition receptors for broad-spectrum disease resistance. *Annu. Rev. Phytopathol.* 55, 257–286 (2017).

2. Cheng, Y. T., Zhang, L. & He, S. Y. Plant-microbe interactions facing environmental challenge. *Cell Host Microbe* 26, 183–192 (2019).

3. Zipfel, C. & Oldroyd, G. Plant signalling in symbiosis and immunity. *Nature* 543, 338–340 (2017).

4. Couto, D. & Zipfel, C. Regulation of pattern recognition receptor signalling in plants. *Nat. Rev. Immunol.* 16, 537–552 (2016).

5. Zipfel, C. et al. Bacterial disease resistance in Arabidopsis through flagellin perception. *Nature* 428, 764–767 (2004).

6. Zipfel, C. et al. Perception of the bacterial PAMP EF-Tu by the receptor EFR restricts *Agrobacterium*-mediated transformation. *Cell* 125, 749–760 (2006).

7. Gomez-Gomez, L. & Boller, T. FLS2: an LRR receptor-like kinase involved in the perception of the bacterial effector flagellin in Arabidopsis. *Mol. Cell* 5, 1003–1011 (2000).

8. Cao, Y. T. et al. The kinase LYK5 is a major chitin receptor in Arabidopsis and forms a chitin-induced complex with related kinase CERK1. *Eukaryot. Cell* 3, e00766 (2014).

9. Miya, A. et al. CERK1, a LysM receptor kinase, is essential for chitin elicitor signaling in Arabidopsis. *Proc. Natl Acad. Sci. USA* 104, 19613–19618 (2007).

10. Akamatsu, A. et al. An OsCEBIP/OsCERK1-OsRacGEF1-OsRac1 module is an essential early component of chitin-induced rice immunity. *Cell Host Microbe* 13, 465–476 (2013).

11. Bacte, L. et al. Plant cell wall-mediated immunity: cell wall changes trigger disease resistance responses. *Plant J.* 93, 614–636 (2018).

12. Boller, T. & Felix, G. A. Renaissance of elicitors: perception of microbe-associated molecular patterns and danger signals by pattern-recognition receptors. *Annu. Rev. Plant Biol.* 60, 379–406 (2009).

13. Seong, S. Y. & Matzinger, P. Hydrophobicity: an ancient damage-associated molecular pattern that initiates innate immune responses. *Nat. Rev. Immunol.* 4, 469–478 (2004).

14. Yamaguchi, Y., Pearce, G. & Ryan, C. A. The cell surface leucine-rich repeat receptor for AtPEP1, an endogenous peptide elictor in Arabidopsis, is functional in transgenic tobacco cells. *Proc. Natl Acad. Sci. USA* 103, 10104–10109 (2006).

15. Huffaker, A., Pearce, G. & Ryan, C. A. An endogenous peptide signal in Arabidopsis activates components of the innate immune response. *Mol. Plant–Microbe Interact.* 18, 511–517 (2005).

16. Hou, S. G. et al. Identification of a plant receptor for extracellular ATP. *Science* 343, 290–294 (2014).

17. Lüstig, K. D. et al. Expression cloning of an ATP receptor from mouse neuroblastoma cells. *Proc. Natl Acad. Sci. USA* 90, 5113–5117 (1993).

18. Carpi, N. C. Update on mechanisms of plant cell wall biosynthesis: how plants make cellulose and other (1->4)-beta-D-glycans. *Plant Physiol.* 155, 171–184 (2011).

19. Burton, R. A. & Fincher, G. B. Current challenges in cell wall biology in the cereals and grasses. *Front. Plant Sci.* 3, 130 (2012).
50. Zhou, F. et al. Co-incidence of damage and microbial patterns controls localized immune responses in roots. Cell 180, 440–453 (2020).
51. Melida, H. et al. Arabinoxylan-oligosaccharides act as damage associated molecular patterns in plants regulating disease resistance. Front. Plant Sci. 11, 1210 (2020).
52. Doehlemann, G. & Hemetsberger, C. Apoplastic immunity and its suppression by filamentous plant pathogens. N. Phytol. 198, 1001–1016 (2013).
53. Buscail, P. et al. Glycosidase and glycan polymorphism control hydrolytic release of immunogenic flagellin peptides. Science 364, eaav0748 (2019).
54. Gao, F. et al. Deacetylation of chitin oligomers increases virulence in soil-borne fungal pathogens. Nat. Plants 5, 1167–1176 (2019).
55. Benedetti, M. et al. Four Arabidopsis berberine bridge enzyme-like proteins are specific oxidases that inactivate the elicitor-active oligogalacturonides. Plant J. 94, 260–273 (2018).
56. He, J. et al. A Lyn4 receptor heteromer mediates perception of arbuscular mycorrhizal symbiotic signal in rice. Mol. Plant 12, 1561–1576 (2019).
57. Yang, C. et al. Activation of ethylene signaling pathways enhances disease resistance by regulating ROS and phytoalexin production in rice. Plant J. 89, 338–353 (2017).
58. Li, X. et al. MofEnd3 regulates appressorium formation and virulence through mediating endocytosis in rice blast fungus Magnaporthe oryzae. PLoS Pathog. 13, e1006449 (2017).
59. Kim, S. G. et al. In-depth insight into in vivo apoplastic secretome of rice-Magnaporthe oryzae interaction. J. Proteom. 78, 58–71 (2013).
60. Haslam, R. P. et al. The assessment of enriched apoplastic extracts using proteomic approaches. Ann. Appl. Biol. 143, 81–91 (2003).
61. O’Leary, B. M. et al. The infiltration-centrifugation technique for extraction of apoplastic fluid from plant leaves using Phaeosol vulgaris as an example. J. Vis. Exp. 19, 52113 (2014).
62. Ghorai, T. Measurement of cellulase activities. Pure Appl. Chem. 59, 257–268 (1987).

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Author contributions
J.L., C.Y., and R.L. conceived and designed the experiments; C.Y. and R.L. performed most of the experiments; J.P., R.R., G.W., H.Z., and E.W. provided technical assistance; J.L. and C.Y. wrote the article.

Competing interests
The authors declare no competing interests

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