

Proteomic Analysis to Identify Tightly-Bound Cell Wall Protein in Rice Calli

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Rice is a model plant widely used for basic and applied research programs. Plant cell wall proteins play key roles in a broad range of biological processes. However, presently, knowledge on the rice cell wall proteome is rudimentary in nature. In the present study, the tightly-bound cell wall proteome of rice callus cultured cells using sequential extraction protocols was developed using mass spectrometry and bioinformatics methods, leading to the identification of 1568 candidate proteins. Based on bioinformatics analyses, 389 classical rice cell wall proteins, possessing a signal peptide, and 334 putative non-classical cell wall proteins, lacking a signal peptide, were identified. By combining previously established rice cell wall protein databases with current data for the classical rice cell wall proteins, a comprehensive rice cell wall proteome, comprised of 496 proteins, was constructed. A comparative analysis of the rice and Arabidopsis cell wall proteomes revealed a high level of homology, suggesting a predominant conservation between monocot and eudicot cell wall proteins. This study importantly increased information on cell wall proteins, which serves for future functional analyses of these identified rice cell wall proteins.

INTRODUCTION

In higher plants, the cell wall, also known as the extracellular matrix, functions as a means of mechanical support as well as establishing an apoplasmic matrix for signaling and defense. The initial cell wall is deposited during cytokinesis and is termed the primary wall; it is composed of polysaccharides, such as

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cellulose, hemicelluloses, and pectins (Keegstra, 2010; Pettolino et al., 2012; Tan et al., 2013; Yang et al., 2011). In the case where wall strengthening is required, this is achieved through the formation of secondary walls and, here, lignin is also incorporated into the polysaccharide matrix. These wall constituents are thought to play important roles in wall maintenance, metabolic and developmental regulation, as well as responses to environmental cues (Jamet et al., 2008; Rose and Lee, 2010). As a prerequisite to developing a comprehensive understanding of the functional importance of cell wall proteins, it will be necessary to compile cell wall proteomes for a number of model plant species. To this end, cell wall proteomics studies have been performed on alfalfa, Arabidopsis, chickpea, maize, rice, soybean, tobacco, tomato, and wheat (Bayer et al., 2006; Bhushan et al., 2006; Chen et al., 2009; Cho et al., 2009; Dahal et al., 2010; Komatsu et al., 2010; Kong et al., 2010; Millar et al., 2009; Watson et al., 2004; Zhu et al., 2007). Of these plant species, the most extensive database has been developed for Arabidopsis; here, some 500 proteins have been identified using a combination of 2D-gel/MudPIT and genome/bioinformatics tools (Bayer et al., 2006; Borderies et al., 2003; Charmont et al., 2005; Chivasa et al., 2002; Ndimba et al., 2003).

To date, rice cell wall proteomics studies have focused on identifying non-bound apoplasmic and weakly-bound proteins, but not tightly-bound cell wall proteins. Other studies have investigated the induction of wall proteins in response to cold, dehydration, pathogen challenge and chemical stress (Chen et al., 2009; Cho et al., 2009; Cui et al., 2005; Ge et al., 2008; Jung et al., 2008; Kim et al., 2009; Pandey et al., 2010; Zhou et al., 2011). Previously, our group employed a rice callus culture system to identify non-redundant classical cell wall proteins containing conventional signal peptides. Here, 249 and 153 cell wall proteins were characterized as weakly-bound and secreted proteins, respectively (Chen et al., 2009; Cho et al., 2009). Given that the number of identified rice classical cell wall proteins was significantly below that observed for Arabidopsis, it would seem that this database is not yet comprehensive in nature.

Cell wall proteins have various affinities to the extracellular matrix, ranging from fully mobile proteins to those that are tightly integrated into the matrix, via covalent linkages; these latter proteins cannot be extracted by strong salt solutions (Jamet et al., 2008; Rose and Lee, 2010). Destructive methods, which require extensive grinding of the plant material, CaCl₂ extraction and SDS/ dithiothreitol extraction, have previously been employed to extract tightly-bound cell wall proteins (Feiz et al.,

2006; Jamet et al., 2008; Rose and Lee, 2010). In the present study, an expanded rice cell wall proteome, based on callus cultured cells, was developed using mass spectrometry and bioinformatics methods. A combination of destructive and sequential extraction protocols was employed to purify tightlybound wall proteins; this approach led to the identification of 1,999 candidate proteins. Based on bioinformatics analyses, 389 classical rice cell wall proteins, possessing a signal peptide, and putative 334 non-classical cell wall proteins, lacking a signal peptide, were identified. The classical rice callus cell wall proteins were compared to previously established rice callus cell wall protein databases, and a comprehensive rice callus cell wall proteome containing 496 proteins was constructed. A comparative analysis of the rice and Arabidopsis cell wall proteomes revealed a high level of homology, suggesting a predominant conservation between monocot and eudicot cell wall proteins.

MATERIALS AND METHODS

Callus induction

Rice callus cultures were established using the following procedure. Rice seeds (*Oryza sativa* L. cv. 'Dongjin') were first dehusked and washed in tap water to remove dust and other surface contaminants. Washed seeds were then surface-sterilized for 30 min in 2% NaOCI solution, rinsed extensively (3 times) with sterilized water and then inoculated on Nitsch's basal (NB) callus induction medium (N6 major salts, N6 minor salts, N6 vitamins, 1 g/L casamino acids, 30 g/L sucrose, 2 mg/L 2,4-D, 2 g/L Gelrite, pH 5.8), as previously described (Hiei et al., 1994). Callus formation was induced by culturing seeds at 30°C in darkness for three weeks. Proliferating calli were sub-cultured on NB medium every two weeks.

Isolation of cell walls from rice calli

Aliquots (20 g) of cultured rice calli (Fig. 1A) were first frozen in liquid nitrogen and cells were then disrupted using several rounds of vortexing in a commercial blender that was prechilled. Disrupted rice calli were further homogenized using a mortar and pestle. Wall preparation buffer (WPB; 50 mM Tris, pH 8.0, 100 mM KCl, 10% v/v glycerol, 10 mM EDTA, 1 mM DTT and 1 mM phenylmethanesulfonylfluoride [PMSF]) was added to homogenized calli (4 ml/g of wall preparation) and the mixture was then centrifuged, at 400x g, for 5 min using a 5810 R centrifuge (Eppendorf AG, Germany). The supernatant was discarded and the pellet (Fig. 1B) was resuspended in 400 ml WPB, without PMSF. This suspension was then further homogenized using three rounds of French press treatment (13 MPa minimal outlet aperture pressure). Next, 200 ml of WPB was added, followed by sonication (1 min × 10 cycles). Aliquots of the suspended pellet (Fig.1C) were equally distributed into four 250 ml tubes. Each tube was centrifuged at 428× g for 3 min, the pellet was then washed with 50 ml of WPB containing 0.1% triton X-100 and recentrifuged at 260x g for 3 min; this step was repeated twice, followed by centrifugation at 115× g for 3 min. After each centrifugation, the supernatant was removed. Finally, the pellet was washed five times with 50 ml of WPB without triton X-100 and then centrifuged at 115× g for 3 min, yielding a clear supernatant.

Protein extraction from rice cell walls

To extract proteins from purified rice callus cell walls, two volumes of 0.2 M $CaCl_2$ solution were added to the final cell wall pellet and the mixture was incubated for 2 h with stirring at $4^{\circ}C$.

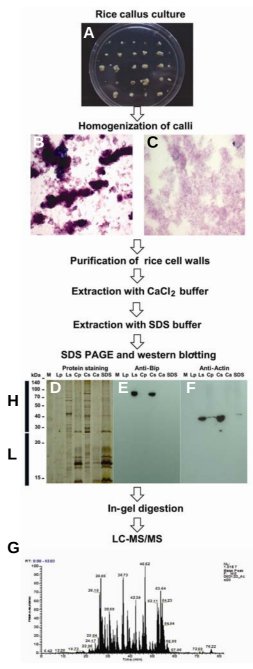


Fig. 1. Experimental scheme for extraction of tightly-bound rice cell wall proteins. (A) Rice callus culture was maintained on NB medium. (B) Rice calli after homogenization in a blender, followed by grinding using a mortar and pestle; note the presence of remaining intact tissue. (C) Complete cellular disruption achieved following French press treatment. (D) Extracted cell wall proteins separated on 1-D SDS-PAGE: M, molecular marker; Lp, leaf pellet; Ls, leaf soluble fraction; Cp, callus pellet; Cs, callus soluble fraction (cytosolic proteins in supernatant); Ca, CaCl₂-extracted cell wall proteins derived from Cp; SDS, SDS-extracted proteins from pellet after CaCl₂ extraction. (E, F) Western blot analyses performed to evaluate the purity of extracted cell wall proteins. Actin (E): cytosolic and PD marker, BiP (F): ER marker. (G) An MS/MS spectrum conducted to identify rice cell wall proteins.

After centrifugation at $15000\times$ g for 3 min, the supernatant was collected and extracted proteins were incubated with four volumes of cold acetone for 2 h at 4°C. The mixture was centrifuged at $15000\times$ g for 15 min, and the resultant pellet was dried and resuspended in a minimal volume of sample buffer. The remaining pellet was further sequentially washed with 0.2 M CaCl₂ solution and TBS buffer. Residual proteins were extracted from the pellet by boiling for 5 min in two volumes of 2X sodium dodecyl sulfate (SDS) extraction buffer (62 mM Tris-HCl, pH 6.8, 2% SDS, 10% v/v glycerol, 0.005% bromophenol blue, 100 mM dithiothreitol).

In-gel digestion and peptide sample preparation

Extracted proteins were loaded onto a 12% polyacrylamide mini-gel (5 \times 8 cm) for 1-D SDS-PAGE (100 V for 45 min in SDS-PAGE running buffer; 25 mM Tris, pH 8.3, 192 mM glycine, 0.1% SDS). Analytical and preparative gels were visualized with silver-staining (Yan et al., 2000) and Coomassie brilliant blue (CBB) staining buffer (50% methanol, 10% glacial acetic acid, 40% H₂O, 0.5 g/L CBB), respectively. Gels were washed thoroughly with destaining buffer (30% methanol, 10% glacial acetic acid in H₂O) and then cut into two parts to yield high and low molecular weight portions. These gel pieces were trypsin-digested and processed for LC-MS/MS analysis.

MudPIT analysis

MudPIT analysis was conducted as described previously (Cho et al., 2009). In brief, peptides were pressure-loaded onto a fused silica capillary column (100 μm i.d.) that contained 7 cm of 5- μ m Aqua C18 material (Phenomenex, USA), 3 cm of 5 μ m Partisphere strong cation exchanger (Whatman, USA), and 3 cm of 5-µm Aqua C18 reversed phase column material (Phenomenex). Peptides were eluted from the microcapillary column and electrosprayed into a LTQ linear ion trap mass spectrometer (ThermoElectron), with a 2.3-kV spray voltage used distally and applied to the waste of the HPLC split. A cycle of a full-scan mass spectrum (400-1400 m/z) followed by nine data-dependent tandem mass spectrometry (MS/MS) spectra at 35% normalized collision energy was repeated continuously throughout the multidimensional separation. The MS/MS spectra obtained from the LC-ESI-MS/MS analyses were used to search against a rice protein database from the NCBI website (http://www.ncbi.nlm.nih.gov/) using TurboSequest (SageN, CA), while tandem mass spectra were extracted using BioWorks version 3.0. Charge state deconvolution and deisotoping were not carried out. Sequest was searched with a fragment ion mass tolerance of 1.0 Da and a parent ion tolerance of 3.0 Da. The iodoacetamide derivative of cysteine was allowed in Sequest as a fixed modification (cysteines +57). Oxidation of methionine was assigned in Sequest searches as a variable modification (methionine +16), with a maximum of three modifications allowed per peptide (the maximum number of modification per type was five), and maximum of two missed cleavage sites for trypsin digestion. Proteins that contained similar peptides and could not be differentiated, based on MS/MS analysis alone, were grouped to satisfy the principles of parsimony. Bioworks v. 3.0 was used to filter the search results, and the following Xcorr values and a delta Cn value of 0.1 were applied to different charge states of peptides: 1.8 for singly charged peptides, 2.5 for doubly charged peptides, and 3.5 for triply charged peptides. A requirement of at least one tryptic-digested end was used in the filtering process.

Bioinformatics analyses

SignalP 3.0 program including two different algorithms (Signal-

NN and SignalP-HMM; http://www.cbs.dtu.dk/services/SignalP) was used to predict the presence and location of signal peptide cleavage sites in each identified protein sequence (Emanuelsson et al., 2007). The 0.43 value was applied as a default cutoff in both NN and HMM algorithms. If the score was higher than the selected cutoff, the comment would be 'Yes', otherwise the comment would be 'No'. To predict transmembrane domains, in each identified protein sequences, the TMHMM v. 2.0 program was applied (http://www.cbs.dtu.dk/services/TMHMM) (Möller et al., 2001). The cutoff value for the TMHMM program was 0.5, indicating the presence of predicted transmembrane helices.

The TargetP 1.1 was used to predict proteins localized to plastids and mitochondria with appropriate default parameters (Emanuelsson et al., 2007). Non-classical secreted proteins were predicted by the SecretomeP program and proteins with a neural network-score (NN-score) more than 0.5 were considered as non-classical secreted proteins (Bendtsen et al., 2004). Three different web-based programs, PredGPI prediction server (http://gpcr2.biocomp.unibo.it/gpipe/index.htm), GPI-SOM (http:// gpi.unibe.ch/) and Big-PI Predictor (http://mendel.imp.ac.at/gpi/ gpi_server.html), were used to identify glycosylphosphatidylinositol (GPI) anchored signals (Eisenhaber et al., 2003; Fankhauser and Mäser, 2005; Pierleoni et al., 2008). In general, proteins with GPI-anchoring signals should possess a signal peptide for the secretory pathway; therefore, proteins having both N- and C-terminal signals were regarded as GPI-anchored proteins by the GPI-SOM program. The specificity cutoff for the PreGPI program should be more than 99% and the Big-PI program produced the quality of the site with P or S.

Conserved protein domains were identified by performing InterProScan (http://www.ebi.ac.uk/Tools/InterProScan) implemented in Blast2GO (Götz et al., 2008; Mulder and Apweiler, 2007). When a protein contained endoplasmic reticulum (ER) retention signals, like KDEL or HDEL sequences at the C-terminus, it was regarded as an ER protein (Denecke et al., 1992). To remove redundant rice proteins, all identified proteins were reassigned with corresponding rice accession numbers, using BLASTP (http://blast.ncbi.nlm.nih.gov/Blast.cgi) (Altschul et al., 1997). In order to find homologous *Arabidopsis* cell wall proteins, all rice proteins were blasted against the whole *Arabidopsis* proteome. For the BLASTP analysis, an E-value of 1E-6 was used as a threshold.

Immunoblot analyses

To perform immunoblot analyses of the extracted proteins, two different antibodies were used. One was anti-actin known to detect proteins localized in the cytoplasm and plasmodesmata (PD). The second was anti-BiP, which detects the molecular chaperone, BiP, located in the ER lumen. Proteins were separated on 10% SDS-PAGE gels and electro-transferred onto PVDF membrane. After blocking with 7% skim milk, the membrane was incubated with rabbit polyclonal anti-actin (LabFrontier) or mouse monoclonal anti-BiP antibodies (Stressgen). Then horse radish peroxidase-conjugated donkey anti-rabbit, or goat anti-mouse, secondary antibodies (Santa Cruz Biotechnology) were used to react with anti-actin and anti-BiP, respectively. The antibodies were detected by chemiluminescence with an ECL kit (Amersham Biosciences).

Cloning, agroinfiltration and confocal imaging

To validate cell wall localization of non-classical secreted proteins, two candidate proteins, LOC_Os02g37710.1 and LOC_Os01g71860.1, were selected. The respective full length CDS was amplified without its stop codon by PCR with gene specific

primers (LOC_Os02g37710.1_FW [AAAAAGCAGGCTTTATG-GAGCTCGTGGAGACAG], LOC_Os02g37710.1_RE [AGAAA-GCTGGGTTCGGGCTAGTTCCGCTC], LOC_Os01g71860.1_ FW [AAAAAGCAGGCTTTATGGGAGCTGTCAATGGTGT] and LOC_Os01g71860.1_RE [AGAAAGCTGGGTTATTCGAAAA-GGCAATCTGATAGAC]) containing attB gateway sites using cDNA prepared from rice seedlings. Purified PCR products were cloned into the pDONR207 entry vector by the BP reaction, according to the manufacturer's instructions (Invitrogen, Germany). LR clonase reactions were next performed to transfer each cDNA from the entry clone into a pMDC83 gateway compatible destination vector containing an N-terminal GFPhis₆-tagged fusion driven by the 35S promoter, according to the manufacturer's instructions (Invitrogen; Curtis and Grossniklaus, 2003). Purified plasmids were finally transformed into competent Agrobacterium strain GV3101 by electroporation.

Agrobacterium cells harboring each GFP fusion construct were grown to an OD600 of 1.0 in liquid yeast extract peptone (YEP) medium (1% peptone, 1% yeast extract, and 0.5% NaCl) supplemented with 50 μg/ml kanamycin, 50 μg/ml rifampicin and 25 μg/ml gentamicin overnight at 28°C. Cells were pelleted, resuspended at OD600 = 1.0 in infiltration buffer [10 mM MgCl2, 10 mM MES (pH 5.6), 100 μM acetosyringone], incubated for 2-3 h at room temperature, and infiltrated into the abaxial side of 6-week-old Nicotiana benthamiana leaves using a 1-ml needleless syringe. Agrobacterium harboring the Tomato bushy stunt virus P19 silencing suppressor was also co-infiltrated in order to minimize the gene silencing effects on heterologous gene expression in tobacco.

After 36 h post-infiltration, green fluorescent signals were observed in leaves using an Olympus (Japan) confocal laser scanning microscope (model FV1000). Excitation and emission wavelengths for GFP were 488/510-540 nm. To perform plasmolysis experiments, cut leaf sections from *N. benthamiana* leaves transiently expressing GFP fusion constructs were vacuum-infiltrated in 0.8 M mannitol for 1 h. After a short incubation period, GFP signals were observed using a CLSM.

RESULTS

Extraction and identification of rice callus cell wall proteins

To extract tightly-bound cell wall proteins, two extraction buffers were sequentially employed. The highly purified cell wall fraction was first subjected to 0.2 M CaCl2 extraction solution that has been well characterized as an excellent agent to remove cell wall proteins (Kim et al., 2013; Robertson et al., 1997). Next, an SDS extraction buffer, including 100 mM dithiothreitol, was used to remove residual tightly-bound cell wall proteins from the cell wall pellet. The CaCl₂ and SDS extracted cell wall proteins, as well as cytosolic proteins, were loaded onto 1-D SDS-PAGE gels. These cell wall-extracted proteins displayed quite different profiles as compared to the cytosolic proteins (Fig. 1D). In both cell wall protein fractions extracted by the CaCl2 and SDS treatments, strong protein bands were detected predominantly in the low molecular weight regions below 20 kDa (Fig. 1D). Note that the band intensity for cell wall proteins extracted with SDS was stronger than that for the CaCl2 treatment. Furthermore, a number of high molecular weight bands were present in the SDS extracted cell wall proteins (Fig. 1D), indicating that SDS/dithiothreitol-based extraction is much effective compared to CaCl₂-based extraction.

To evaluate the purity of these cell wall-extracted proteins, western blot analysis was performed using antibodies specific for actin, a cytosolic/plasmodesmal marker, and BiP, an endo-

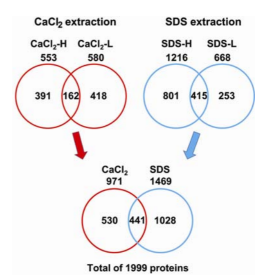


Fig. 2. Venn diagrams illustrating the populations of rice proteins identified using $CaCl_2$ and SDS extraction methods. (A) $CaCl_2$ -extracted proteins were separated and processed based on the scheme presented in Fig. 1. $CaCl_2$ -H and $CaCl_2$ -L; total number of proteins identified within the high (H) and low (L) molecular weight fractions. (B) SDS-extracted proteins were separated and processed based on the scheme presented in Fig. 1. SDS-H and SDS-L; total number of proteins identified within the high (H) and low (L) molecular weight fractions. (C) Comparison between rice proteomes extracted by $CaCl_2$ and SDS methods. A total of 1999 non-redundant rice callus proteins were identified using this sequential two-step extraction protocol.

plasmic reticulum (ER) lumen-localized soluble protein. Strong BiP signals were detected in only the leaf (Ls) and callus (Cs) soluble fractions (Fig. 1E), but not in pellet fractions (Lp and Cp). Actin was detected strongly in the leaf (Ls), callus (Cs) soluble fractions and weakly in callus pellet (Cp) (Fig. 1F). Furthermore, actin was also detected in the SDS-extracted cell wall fraction (Fig. 1F). These results are consistent with the hypothesis that the proteins contained within the CaCl2 and SDS extracted cell wall fractions are likely highly enriched with extracellular proteins. In this regard, actin detected in the SDS fraction may reflect protein derived from PD (Baluska et al., 2004). To prepare the isolated cell wall proteins for interrogation by mass spectrometry, samples were first separated on 1-D SDS-PAGE gels which were subsequently divided into high and low molecular weight sections (Fig. 1D). Following in-gel trypsin digestion, samples were analyzed by LC-MS/MS and protein identification achieved using a TurboSEQUEST program in conjunction with the rice protein database (Fig. 1G). Full details regarding peptide sequences are contained in Supplementary Tables S1-S4.

All identified rice proteins with GenBank identifier (GI) numbers were converted to the corresponding rice protein locus, based on the rice genome annotation (http://rice.plantbiology.msu.edu/) (Ouyang et al., 2007). After removal of redundant proteins, some 553 and 580 proteins were identified from the high and low molecular weight gel sections, respectively, based on the CaCl₂ extraction protocol (Fig. 2A). Statistics for SDS extracted proteins were 1216 and 668 proteins contained within the high and low molecular weight gel fractions, respectively (Fig. 2B). More proteins (1469 in total) were contained

Table 1. Representative rice classical cell wall proteins containing signal peptides predicted by SignalP 3.0

Gl ^a	Function	Molecular mass	pl	Sequence count	Sequence coverage
21104619	Thaumatin family	26189	7.7	104	92%
62733218	Glycosyl hydrolase	33947	9.3	64	79%
113683	Alpha-amylase	48708	5.6	62	73%
55168113	Glycosyl hydrolase	32549	6.5	59	70%
50918709	Periplasmic beta-glucosidase	67778	7.5	45	43%
34895554	Glycosyl hydrolase family 31	102551	6.5	40	43%
50508397	Glucan endo-1, 3-beta-glucosidase	59363	5.4	33	64%
50940911	Cysteine-rich repeat secretory protein 55	30046	7.9	33	67%
50905163	Retrotransposon protein	70761	7.7	28	35%
51963488	Polygalacturonase inhibitor	35465	7.4	28	76%
50940909	Cysteine-rich repeat secretory protein 55	29829	7.9	26	62%
53792759	Retrotransposon protein	56264	7.9	23	41%
113682	Alpha-amylase	47911	6.4	23	59%
51964350	Dehydrogenase	42702	7.8	21	55%
50940901	Cysteine-rich repeat secretory protein 55	29695	7.9	21	57%
113766	Alpha-amylase	47756	5.3	20	59%
57900682	Uncharacterized protein At4g06744	44330	6.5	19	36%
55168101	Cellulase	57788	6.4	19	42%
34898314	GDSL-like lipase/acylhydrolase	40958	8.2	18	47%
20160767	Xylanase inhibitor	43787	9.2	15	43%

^aGI (GenBank identifier) number indicates the protein accession number in NCBI.

Table 2. Representative rice non-classical cell wall proteins that lack a signal peptide as predicted by SecretomeP 2.0

Glª	Function	Molecular mass	pl	Sequence count	Sequence coverage	NN-score ^b
51963380	Unknown protein	67355	6.7	27	0.576	0.524
50509692	Beta-1,3-glucanase	53753	6.1	26	0.585	0.725
51964830	Unknown protein	67154	7.2	20	0.386	0.599
22830913	Endo-1,3-beta-glucanase	67635	5.5	18	0.37	0.683
460989	Beta tubulin	43002	4.9	14	0.386	0.55
38345164	Unknown protein	71231	8.8	11	0.255	0.566
50912401	Aldehyde dehydrogenase	45483	6.3	11	0.333	0.543
50938789	Early nodulin 8 like protein	22215	8.2	9	0.47	0.582
3023713	Enolase	47987	5.6	8	0.215	0.53
38346903	Unknown protein	52786	8.3	8	0.176	0.503
50510001	Prohibitin	31955	9.8	8	0.343	0.607
51964530	Unknown protein	46830	8.2	8	0.208	0.702
730456	40S ribosomal protein S19	16387	10	7	0.507	0.703
34893994	40S ribosomal protein S5	22227	9.7	7	0.25	0.827
38344200	Unknown protein	55227	7.9	7	0.153	0.539
542153	Translation initiation factor eIF-4A	46932	5.5	6	0.232	0.573
37533060	Unknown protein	96740	5.8	6	0.085	0.589
42408023	Phosphoribosyl pyrophosphate synthetase	36149	6.8	6	0.265	0.823
50907169	Unknown protein	90986	6.5	6	0.129	0.587
51854281	Unknown protein	90130	5.5	6	0.117	0.525

^aGI (GenBank identifier) number indicates the protein accession number in NCBI. ^bCutoff value of the NN-score for non-classical cell wall proteins is greater than 0.5.

Table 3. GPI-anchored proteins isolated from the rice cell wall preparation identified by three prediction programs

Glª	Function ^b	Rice locus	Evalue	Length	Molecular mass	pl	Omega-site ^c
62733403	Os3bglu8 - beta-glucosidase	LOC_Os03g49610.1	0	603	67245	7.4	573
55733786	Glucan endo-1,3-beta-glucosidase-like protein 3	LOC_Os05g43690.1	5.00E-52	228	21760	5.7	204
51964118	Glycerophosphoryl diester phosphodiesterase family	LOC_Os02g37590.1	0	749	81084	6.1	723
51963864	Glucan endo-1,3-beta-glucosidase	LOC_Os02g04670.1	0	489	53375	5.5	459
51243456	LysM domain-containing GPI-anchored protein 1	LOC_Os06g10660.1	0	409	40488	4.9	387
50943329	Plastocyanin-like domain containing protein	LOC_Os08g17160.1	3.00E-69	193	19274	8.3	169
50941247	Monocopper oxidase	LOC_Os08g05820.1	0	600	66229	6.8	576
50939031	X8 domain containing protein	LOC_Os07g40940.1	7.00E-43	191	18902	5.7	167
50932835	X8 domain containing protein	LOC_Os05g50490.3	1.00E-136	281	28222	6	255
50919115	Glucan endo-1,3-beta-glucosidase	LOC_Os03g57880.3	0	491	52783	5	467
50918839	X8 domain containing protein	LOC_Os03g54910.1	1.00E-68	175	17825	4.8	149
50912821	Eukaryotic aspartyl protease domain containing protein	LOC_Os02g51540.1	0	520	56272	6	494
50907029	Pectinesterase	LOC_Os02g18650.1	0	554	58764	9.5	536
38605955	Fasciclin-like arabinogalactan protein	LOC_Os04g48490.1	1.00E-170	431	44536	6.7	410
37805880	Retrotransposon protein	LOC_Os08g16810.1	8.00E-51	130	13441	10.4	102
37531036	Thaumatin-like protein 1	LOC_Os10g05600.1	1.00E-135	389	38581	4.8	362
34897712	Monocopper oxidase	LOC_Os06g01490.1	0	593	65750	6.5	570

^aGI (Gene Identifier) number indicates the protein accession number in NCBI.

Table 4. A comparison between the classical cell wall proteins identified by four different cell wall proteome studies. Criterion for classical cell wall proteins is that they contain a signal peptide predicted by SignalP3.0

Description	Weakly bound proteins ^a	Purely secreted proteins ^b	Tightly bound proteins (CaCl ₂) ^c	Tightly bound proteins (SDS) ^d
No. of classical cell wall proteins	205	153	251	312
No. of proteins without a signal peptide	1456	402	722	1157
No. of total proteins	1705	555	973	1469
Ratio of classical cell wall proteins to total proteins	14.6%	27.6%	25.8%	21.2%

^aStudy on weakly-bound rice cell wall proteome (Chen et al., 2009)

within the SDS-extracted fraction when compared to the 971 proteins extracted by $CaCl_2$ (Fig. 2C). The combined total of all rice proteins identified in this study was 1999; 530 and 1028 proteins were unique and 441 proteins were common for the $CaCl_2$ - and SDS-extracted fractions, respectively (Fig. 2C).

Proteins predicted to contain a signal peptide

In general, protein targeting to the extracellular space requires an N-terminal signal peptide that mediates in the translocation of secreted proteins through the ER/Golgi pathway (Rabouille et al., 2012). The presence of a signal peptide in the N-terminus of identified cell wall proteins was investigated using the SignalP

^bPutative functions of 17 GPI-anchored proteins based on the annotation of the Rice Genome Annotation Project (http://rice.plantbiology.msu.edu/).

^cOmega-site for GPI-anchored signal in protein sequences was predicted by the Big-PI program.

^bStudy on rice secretome (Cho et al., 2009)

[°]Current study on tightly-bound rice cell wall proteome extracted by CaCl₂

^dCurrent study on tightly-bound rice cell wall proteome extracted by SDS

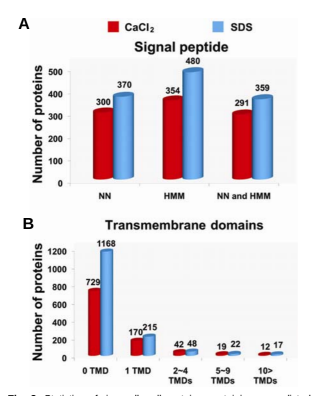


Fig. 3. Statistics of rice cell wall proteins containing a predicted signal peptide and transmembrane domains. (A) Number of proteins having a predicted signal peptide based on SignalP 3.0 using the NN and HMM algorithms. (B) Number of proteins having predicted transmembrane domains (TMDs) based on the TMHMM v. 2.0 program.

3.0 algorithms, SignalP-NN and SignalP-HMM (Emanuelsson et al., 2007). Based on the NN program, some 300 CaCl₂- and 370 SDS-extracted proteins were predicted to contain signal peptides (Fig. 3A). As compared to the NN algorithm, the number of secreted proteins predicted by the HMM method to have a signal peptide was slightly higher, being 354 and 480 from CaCl₂ and SDS treatments, respectively (Fig. 3A). Finally, to ensure a low error rate, only rice proteins (458 unique proteins) containing a signal peptide predicted by both algorithms (291 and 359 from CaCl₂ and SDS treatments) were categorized for further consideration to identify classical cell wall proteins (Emanuelsson et al., 2007).

Identification of proteins containing predicted transmembrane helices

Given the intimate association between the plant cell wall and the plasma membrane, it is technically difficult, if not impossible, to separate completely the proteins from these two cellular domains. Furthermore, PD are plant-specific intercellular organelles that are inserted within the matrix of the cell wall and contain both ER and plasma membrane components. Thus, it is likely that our cell wall proteome will include PD-localized plasma membrane and ER_proteins. To assess this situation, the TMHMM v. 2.0 program was used to identify candidate proteins containing transmembrane helices (Möller et al., 2001). As expected, 243 cell wall proteins were predicted to contain one or more transmembrane domains (Supplementary Table

S5). For example, 25% and 21% of the proteins within the CaCl₂ and SDS extracted fractions, respectively, were predicted to be membrane proteins (Fig. 3B).

The number of transmembrane helices for any given cell wall protein was highly variable, ranging from one to 20 domains. For instance, there were 170 CaCl₂ and 215 SDS extracted proteins that contained a single transmembrane domain (Fig. 3B). Here, some 27 predicted receptor-like kinase proteins were identified that contained a single transmembrane domain (Supplementary Table S6). In contrast, two proteins, callose synthase 1 (LOC_Os03g03610.3) and embryogenesis transmembrane protein (LOC_Os06g18880.1), have 15 and 20 predicted transmembrane domains, respectively (Supplementary Table S6).

Identification of classical cell wall proteins based on bioinformatics

A further analysis of the signal peptide-containing cell wall proteins (Fig. 3A) identified 458 non-redundant proteins, which were predicted as a signal peptide-containing cell wall protein by both the NN and HMM algorithms (Supplementary Table S6). Among them, 66 were eliminated as they contained more than one predicted membrane domain. However, in other cases, the transmembrane domain within the N-terminus was often predicted as a signal peptide because of the hydrophobic characteristics. Hence, when the predicted transmembrane helix overlapped with a predicted signal peptide, such proteins were regarded as classical cell wall or secreted proteins. Based on this criterion, predicted transmembrane domains in 178 proteins were classified as a signal peptide and these were included in the secreted protein category (Supplementary Table S6).

Proteins contained within the ER possess classical retention signals, such as an HDEL or KDEL sequence located at the C-terminus. Interestingly, we identified only two DnaK family proteins (LOC_Os05g35400.1 and LOC_Os02g02410.1) and a hydrolase (LOC_Os01g37960.1) that possessed ER retention motifs (Supplementary Table S6). This finding supports the notion that our extracted rice proteins are specifically enriched for cell wall proteins. After exclusion of 66 plasma membrane and 3 ER proteins, based on the above criteria, we identified some 389 classical rice cell wall proteins predicted to be secreted into the extracellular matrix, via a default secretory pathway (Table 1 and Supplementary Table S7).

Prediction of non-classical secreted proteins

In comparison to previous 2-DGE-based cell wall proteomics, the number of proteins lacking a signal peptide was quite high. Thus, the possibility existed that, of the remaining 1610 identified proteins, a significant number might well reflect cytosolic and membrane contaminants. However, some of these proteins may be secreted via a non-classical secretory pathway.

In order to remove proteins targeted to organelles, such as plastids and mitochondria, the TargetP 1.1 program was employed to screen the remaining rice proteins (Emanuelsson et al., 2007). Based on this analysis, 205 and 226 proteins were identified as likely being targeting to plastids and mitochondria, respectively (Fig. 4). We next used the SecretomeP program to identify potential non-classical secreted proteins (Bendtsen et al., 2004). Although this program was initially designed for mammalian cells, we assumed that secreted proteins in eukaryotic cells might well have common properties. In any event, using this approach 334 putative non-conventional secreted proteins were identified (Table 2, Fig. 4 and Supplementary Table S8A). The cell wall localization of such proteins was ex-

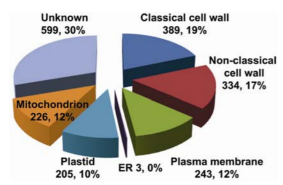


Fig. 4. Pie diagram illustrating the potential sub-cellular protein localization for the rice callus proteome. Classical cell wall proteins contain a signal peptide (predicted by both NN and HMM algorithms of SignalP 3.0), but lack transmembrane domains (predicted by the TMHMM v. 2.0 program). Plasma membrane proteins were predicted by the TMHMM v. 2.0 program. Proteins possessing both a predicted signal peptide and a transmembrane domain were scored as classical cell wall proteins. Plastid and mitochondrial proteins were predicted by the TargetP 1.1 program and considered as contaminated proteins. Non-classical cell wall proteins were predicted by the SecretomeP 2.0 program. Proteins containing an ER retention motif (KDEL or HEDL) at C-terminal region were identified as ER proteins.

perimentally validated by randomly picking up two proteins to be fused with fluorescent reporter green fluorescent protein (GFP) using tobacco leaf infiltration system. As shown in Supplementary Fig. S3, green fluorescent signals associated with LOC_Os02g 37710.1 belonging to lecithin cholesterol acyltransferase-like protein family, and LOC_Os01g71860.1 belonging to endo-1,3-beta-glucanase family were found not only in cytoplasm, but also in extracellular space as shown by plasmolysis. Collectively, we identified 389 classical cell wall proteins (19%) and 334 putative non-classical (17%) cell wall proteins, leaving 599 proteins (30%) unassigned to any predicted subcellular localization (Fig. 4).

Identification of GPI-anchored proteins

In plants, many proteins can be anchored to the extracellular surface of the plasma membrane following glycosylphosphatidylinositol (GPI) posttranslational modification. In general, GPI-anchored proteins have two highly conserved domains, composed of an N-terminal signal peptide and a C-terminal hydrophobic signal sequence, termed the omega site (Eisenhaber et al., 2003). This site is located between 20 and 30 residues upstream of the C-terminus. During posttranslational processing in the ER, the omega site in GPI-anchored proteins is cleaved and a glycolipid is then attached.

An earlier genome-wide analysis, using computational methods, identified some 198 and 180 GPI-anchored proteins in rice and *Arabidopsis*, respectively (Eisenhaber et al., 2003). To identify putative GPI-anchored proteins, we next used the Pre-GPI, Big-PI and GPI-SOM programs (Eisenhaber et al., 2003; Fankhauser and Mäser, 2005; Pierleoni et al., 2008). Among these programs, the Big-PI program employs experimental data, likely making it the more reliable method (Eisenhaber et al., 2003). The GPI-SOM and Pre-GPI programs predicted 62 and 73 GPI-anchored proteins, respectively, whereas the Big-PI identified only 20 such proteins (Fig. 5).

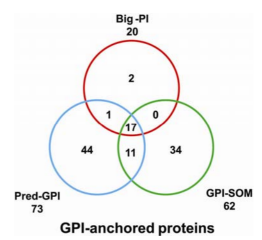


Fig. 5. Venn diagram illustrating GPI-anchored proteins predicted by the Big-PI, Pred-GPI, and GPI-SOM programs

As mentioned above, the presence of a signal peptide is a prerequisite for GPI-anchoring. 18 of 20 proteins predicted by the Big-PI predictor program contained a signal peptide (Supplementary Table S6). However, only 66 of 73 and 52 of 62 proteins predicted by the Pred-GPI and GPI-SOM programs, respectively, had a signal peptide (Supplementary Table S6). Of the identified GPI-anchored proteins, the 17 proteins predicted by all three programs are likely to be GPI-anchored to the plasma membrane (Fig. 5). Here, three glucan endo-1,3-beta-glucosidases, a LysM domain-containing protein, three X8 domain containing proteins, and a monocopper oxidase were identified as GPI-anchored proteins (Table 3). These GPI-anchored proteins are included in the list of classical cell wall proteins, since extracellular domain of these proteins can be cleaved by diverse phosphatases (Brewis et al., 1994).

Distribution of molecular mass and isoelectric points for the rice cell wall proteome

The molecular mass distribution of the 1999 putative cell wall proteins was highly variable, ranging from 8 to 359 kDa, with the average being 60 kDa (Supplementary Table S6). Due to the unbiased characteristics of MudPIT technology, the molecular mass of these proteins exhibited a normal distribution (Supplementary Fig. S1A). The molecular mass distribution of the 389 cell wall and 243 membrane proteins was also examined; profiles similar to that of the total proteome were obtained (Supplementary Figs. S1B and S1C).

The isoelectric points (pls) of proteins in the rice cell wall proteome ranged from 4.3 to 11.9, with 7.5 being the average value (Supplementary Table S6). Here, only 49 and 34 proteins had values less than 5 and higher than 11, respectively (Supplementary Fig. S1D). The distribution patterns of the 1999 proteins and the 389 cell wall proteins were comparable, without any bias towards acidic or alkali proteins (Supplementary Figs. S1D and S1E). This result is similar to that of an earlier *Arabidopsis* cell wall proteome study (Bayer et al., 2006). Interestingly, as shown in Supplementary Fig. S1F, the 243 membrane proteins displayed a dissimilar pl pattern to that of the secreted proteins.

Functional characterization of the rice cell wall proteome To gain insights into the role of the proteins contained within the

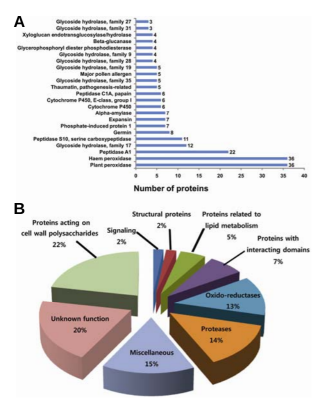


Fig. 6. Conserved domains and functional categories for the 389 classical rice cell wall proteins. (A) Distribution of the most frequently identified conserved domains based on the InterProScan database implemented in Blast2GO. Conserved domains with more than two proteins are displayed. (C) Functional categories identified based on the WallProtDB database (http://www.polebio.scsv.upstlse.fr/WallProtDB/).

rice cell wall proteome, the Blast2Go program was used to assign putative functions, based on gene ontology (GO) (Götz et al., 2008; Jensen and Bork, 2010). Using this approach, some 1200 proteins were annotated in two GO categories, namely biological processes and molecular functions. Based on biological processes, many of these proteins were predicted to be involved in a wide range of metabolic pathways (890 proteins), from response to various stimuli (455 proteins) to developmental processes (413 proteins) and gene expression (263 proteins) (Supplementary Fig. S2A). These findings are consistent with previous studies which have established that cell wall proteins participate in numerous metabolic processes and responses to environmental cues (Jamet et al., 2008). Similarly, based on molecular function, many proteins were assigned to binding (969 proteins), catalytic (842 proteins), hydrolase (414 proteins), transferase (214 proteins), structural (108 proteins) and kinase (106 proteins) activities (Supplementary Fig. S2B). To further analyze the functions likely associated with the 389 classical rice cell wall proteins, we next examined the conserved domains of each protein using the InterProScan database implemented in the Blast2Go program (Götz et al., 2008; Mulder and Apweiler, 2007). Proteins in the following families were found to be abundant: peroxidases (36), peptidase A1 (22) and glycoside hydrolases (36) (Fig. 6A). Protein families containing fewer members included the peptidase S10/serine

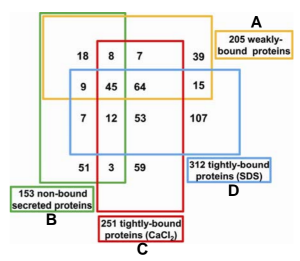


Fig. 7. Distribution of rice cell wall proteins identified by four methods. (A) Weakly-bound cell wall proteome (Chen et al., 2009). (B) Rice proteins identified by secretome analysis (Cho et al., 2009). (C) Tightly-bound cell wall proteins extracted by CaCl₂ treatment. (D) Tightly-bound cell wall proteins extracted by SDS treatment.

carboxypeptidase (11), germin (8), phosphate-induced protein 1 (7), expansin (7) and alpha-amylase (7).

We next categorized these 389 conventional cell wall proteins into eight functional groups, according to the previous annotation of WallProtDB (http://www.polebio.scsv.ups-tlse.fr/WallProtDB/). Here, some 80% of the identified proteins could be assigned into functional groups: among them were proteins acting on cell wall polysaccharides (22%), proteases (14%), oxido-reductases (13%), but various miscellaneous proteins (15%) were also present (Fig. 6B).

Of the identified 334 non-classical rice cell wall proteins, a large fraction (approx. 150 proteins) was assigned as being of unknown function. To further analyze this group, they were examined for conserved domains using the InterProScan database: a total of 793 redundant domains were identified. Domains for armadillo-type fold, RNA recognition motif, nucleotide-binding, F-box domain, protein kinase-like domain, glycoside hydrolase, NAD(P)-binding domain, and tubulin were frequently found (Supplementary Table S8B).

Comparison of rice and Arabidopsis cell wall proteomes

To develop a comprehensive rice cell wall proteome database, we integrated data from our current and previous studies. For this purpose, we used only classical cell wall proteins that possess a signal peptide. As shown in Fig. 7, four independent rice cell wall proteomes have been established using rice callus cultured material. The first study used a non-destructive mannitol/CaCl₂ method to identify some 205 weakly-bound cell wall proteins (Chen et al., 2009). The second study focused on proteins secreted into the rice callus culture medium; here, 153 cell wall proteins were detected (Cho et al., 2009). The third and fourth studies used CaCl₂- and SDS-based protein extraction protocols; 251 and 312 tightly-bound cell wall proteins were extracted, respectively (current study).

Then, we established an expanded rice cell wall proteome database comprised of 496 classical secreted proteins (Supplementary Table S9). This rice database was next used to

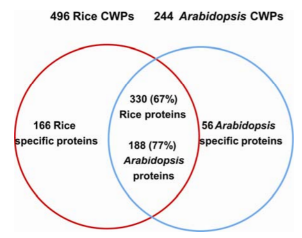


Fig. 8. Comparison of rice and *Arabidopsis* classical cell wall proteins. BLASTP was used to analyze 496 rice and 244 *Arabidopsis* cell wall proteins (CWPs; cutoff value of 1E-6).

analyze the level of conservation with the current *Arabidopsis* cell wall proteome, containing some 244 conventional secreted proteins (Bayer et al., 2006). To this end, the BLASTP program was used with an 1E-6 cutoff value; here, 166 and 56 cell wall proteins were identified that appeared to be specific for rice and *Arabidopsis*, respectively (Fig. 8, Supplementary Tables S10A and S10B). However, 330 rice (67%) and 188 *Arabidopsis* (77%) proteins were common to both proteomes, consistent with a high level of conservation between the cell wall proteomes of the monocots and eudicots (Fig. 8). Future functional analyses of these identified rice cell wall proteins will offer important insights into the roles they perform in wall chemistry, signaling and defense.

DISCUSSION

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In the present study, a large scale proteomics experiment was performed to explore the nature of tightly-bound proteins present in the rice callus cell wall. To this end, we employed a combination of destructive methods, two protein extraction protocols and a MudPIT interrogation approach. Total 1999 proteins were identified by a combination of destructive methods (Fig. 2C). This total is relatively high as compared to 2-DGE-based proteomics, in which around 54-300 cell wall proteins have been identified (Jung et al., 2008; Kim et al., 2013; Zhou et al., 2011), indicating that this current study led to the identification of a significant number of additional plant cell wall-associated proteins.

In our current rice cell wall proteome, we identified 389 classical cell wall proteins possessing a conventional signal peptide, as well as 334 putative non-classical cell wall proteins that lacked such signal peptides. Approximately 80% of the classical cell wall proteins have functions related to cell wall polysaccharide chemistry, as well as to protease and oxido-reductase activities; however, some 40% of the identified non-classical cell wall proteins have unknown functions (Supplementary Tables S7 and S8A) and two candidate proteins LOC_Os02g 37710.1 and LOC_Os01g71860.1 have been experimentally validated for their secretion into extracellular space (Supplementary Fig. S3). This suggests that these non-classical proteins, which utilize unconventional export mechanisms, are likely to repre-

sent important components of the extracellular matrix. To advance plant cell wall biology, it will be important to unraveling the molecular mechanisms that evolved to mediate unconventional protein secretion. In addition, it is important to note that a number of proteins identified in this study are homologous to known non-classical secreted proteins in other organisms, including bacteria, yeast, and mammals for which experimental evidence exists. Examples are annexin, enolase and elongation factor (Edwards et al., 1999; Marcilla et al., 2012; Marques et al., 1998). One such protein, elongation factor 1α that lacks a signal peptide was isolated from tobacco and has been shown to localize to the cell wall by immunogold localization (Zhu et al., 1994). In addition, GAPDH is another known non-classical cell wall protein in bacteria and yeast (Eichenbaum et al., 1996; Gozalbo et al., 1998). The GAPDH domain containing protein (LOC_Os04g40950.1) identified in our study was assigned as an unknown protein without predicted localization. Regardless of proteins that are secreted, via either the classical or nonconventional secretory pathway, it is necessary to stress that other proteins identified in the rice cell wall proteome could well function in unknown biological processes by interacting with various components of the cell wall. Molecular methods will be needed to probe the cell wall localization of these proteins (Groover et al., 2003).

In conclusion, we have analyzed the tightly-bound cell wall proteins in rice calli using a combination of destructive methods. This sensitive and relatively unbiased approach identified a range of proteins involved in metabolic processes and responses to environmental cues. A comparative analysis of the *Arabidopsis* cell wall proteome and comprehensive rice cell wall proteome generated by combining our current and previous studies (Chen et al., 2009; Cho et al., 2009), suggests a predominant conservation between monocot and eudicot cell wall proteins. This rice cell wall proteome dataset could increase information on cell wall proteins, thus it will act as a valuable resource for further functional analysis of cell wall proteins.

Note: Supplementary information is available on the Molecules and Cells website (www.molcells.org).

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