Identification of Differentially Expressed Proteins and Phosphorylated Proteins in Rice Seedlings in Response to Strigolactone Treatment

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Abstract

Strigolactones (SLs) are recently identified plant hormones that inhibit shoot branching and control various aspects of plant growth, development and interaction with parasites. Previous studies have shown that plant D10 protein is a carotenoid cleavage dioxygenase that functions in SL biosynthesis. In this work, we used an allelic SL-deficient d10 mutant XJC of rice (Oryza sativa L. spp. indica) to investigate proteins that were responsive to SL treatment. When grown in darkness, d10 mutant seedlings exhibited elongated mesocotyl that could be rescued by exogenous application of SLs. Soluble protein extracts were prepared from d10 mutant seedlings grown in darkness in the presence of GR24, a synthetic SL analog. Soluble proteins were separated on two-dimensional gels and subjected to proteomic analysis. Proteins that were expressed differentially and phosho-proteins whose phosphorylation status changed in response to GR24 treatment were identified. Eight proteins were found to be induced or down-regulated by GR24, and a different set of 8 phosphoproteins were shown to change their phosphorylation intensities in the dark-grown d10 seedlings in response to GR24 treatment. Analysis of these proteins revealed that they are important enzymes of the carbohydrate and amino acid metabolic pathways and key components of the cellular energy generation machinery. These proteins may represent potential targets of the SL signaling pathway. This study provides new insight into the complex and negative regulatory mechanism by which SLs control shoot branching and plant development.

Introduction

Strigolactones, a group of carotenoid-derived terpenoid lactones, are recently identified endogenous plant hormones that inhibit shoot branching [1,2]. Tillers in rice (Oryza sativa L.) are derived from vegetative shoot branching, and the growth of tillers is one of the most important agronomic traits for rice grain production. It has been shown that monocot and dicot species share a conserved pathway for the biosynthesis of SLs [3–6]. In the last decade, remarkable progress has been made in understanding the molecular basis of shoot branching through studies of a series of increased branching mutants in the SL pathway, including more auxillary growth (max) of Arabidopsis, ramulos (rms) of pea (Pisum sativum), decreased apical dominance (dad) of petunia (Petunia hybrida), and dwarf (d) or high-tillering dwarf (hid) of rice [3–5,7–20]. Three rice genes, D10, D17/HTD1 and D27, have been implicated in the SL pathway. D27 is an iron-binding protein with the  \( \alpha/\beta \)-hydrolase superfamily [3–5,15,18]. Recently, protein structure analysis has revealed that D14 forms an \( \alpha/\beta \)-fold that contains a canonical catalytic triad with a large internal cavity capable of binding SLs [23–25].

SLs have been demonstrated to be important endogenous regulators of plant growth and development both underground and aboveground [26]. Other notable functions of SLs include seed germination [27,28], root growth and architecture [13,29,30], adventitious root formation [31], flower development [13], leaf senescence and development [8,11,13,32]. Genes implicated in the SL biosynthesis and signaling pathways have been shown to express in mature seeds and young seedlings as well as adult plants, suggesting that they play various roles in a spectrum of physiological and developmental processes ranging from germination to leaf development [33].
The mesocotyl of young rice seedlings is a tissue located between the coleoptilar node and a basal part of the seminal root. Previous studies have suggested that the elongation of mesocotyl is controlled by multiple genetic, developmental and environmental signals [34–37]. Rice mutants defective in SL-related genes (d5, d10, d14, d17 and d27) have been shown to produce longer mesocotyl than the wild type seedlings when grown in darkness [30]. Exogenous application of a synthetic SL analog, GR24, can restore the normal mesocotyl phenotype in the SL-deficient mutant (d10, d17 and d27). Regardless of the remarkable progress that has been made, the molecular mechanisms by which SLs control shoot branching and other developmental and physiological processes still remain to be elucidated.

Two-dimensional gel electrophoresis (2-DE) is a powerful proteomics technique for differential display of proteins with posttranslational modifications. It can separate thousands of proteins based on their differences in charge and size [39,40], and has been used to monitor molecular responses induced by the activation or inhibition of specific signaling pathways [41]. Quantitative proteomics analyses have identified a large number of brassinosteroid (BR)-regulated proteins, including the BR signaling components, which are regulated by phosphorylation at the posttranslational level [42–44]. Two of these BR signaling components, BSK1 and BSK2, have been identified through proteomic characterization of phosphorylated proteins of the BR-deficient mutant dsk-2 in response to BR treatment [44]. The identification of these additional components has filled the last gap in the BR signal pathway [44].

Protein phosphorylation and phosphorelay have been recognized as an important mechanism for signal transduction. Among the several ways to detect phosphoproteins, the application of antibodies specific to phosphorytrosine, phosphothreonine, or phosphoserine is advantageous [45]. In this study, we found that mesocotyl elongation of dark-grown rice seedlings was higher in the d10 mutant than in the wild-type plant, and this dark-hypersensitivity could be rescued by exogenous application of GR24. To understand the molecular mechanism underlying the function of SLs in dark-grown d10 seedlings, we analyzed the differential expressed proteins and phosphoproteins in d10 in response to SL treatment, and identified several candidates that may play a role in the SL responses in rice.

Results

Rescue of the tillering phenotype in XJC by GR24

D10 gene expression is controlled by feedback regulation [4], indicating that the level of D10 mRNA accumulation might be a critical step in the regulation of SL biosynthesis. Our previous study revealed that a 39 bp deletion at the second exon of D10 (LOC_Os01g0746400, OsCCD8b) results in the frame-shift mutation in the indica rice mutant XJC, exhibiting a high-tillering dwarf phenotype [46]. Quantitative real-time reverse transcription-PCR (qRT-PCR) analysis also confirms the dramatic up-regulation of D10 expression in XJC compared to that in the wild type GC13 [46]. By applying 1.0 μM GR24, a synthetic strigolactone analog, to the wild-type (GC13) and d10 (XJC) seedlings in a hydroponic culture, we found that the exogenous supplement of GR24 was able to fully inhibit tiller bud outgrowth of 2-week-old d10 seedlings (Figure 1), suggesting that the tillering phenotypes of d10 mutant plants could be rescued by GR24 treatment. These findings further confirm that XJC is indeed defective in SL biosynthesis, while the perception and signaling pathway of SLs in XJC is apparently normal.

Identification of SL-responsive proteins

To examine the molecular mechanism of inhibition of GR24 on mesocotyl elongation in dark-grown d10 mutant seedlings, we applied proteome analysis to the analysis of differentially expressed proteins and phosphoproteins in the d10 mutant seedlings in response to GR24 application. Etiolated XJC seedlings after removal of residual seeds (Figure 2D) with or without GR24 treatment were harvested and their proteomes were resolved by 2-DE. Protein profiles of the gels were visualized with silver staining (Figure 3A–B). A total of 957 ± 3(n = 3) protein spots were detected on each of the 2-DE maps. Ten proteins showed more than 1.5-fold reproducible changes in abundance (Figure 3C). After GR24 treatment in the XJC plants, two proteins (G1 and G9) were up-regulated, five (G2, G4, G6, G7 and G8) down-regulated, and three (G3, G5 and G10) became

Figure 1. Response of d10 mutant and wild type seedlings to the application of GR24. Wild type (WT) GC13 and mutant XJC (d10) seedlings were grown in the presence of 1.0 μM GR24 for two weeks. Tillers were present in the axil of the first (red arrow) and second (white arrow) leaves of the mutant (a), but absent in WT plants and the GR24-treated d10 mutant (b). The tiller outgrowth phenotype in d10 was suppressed by application of GR24. Scale bars = 1 cm.
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Rescue of the mesocotyl elongation phenotype in XJC by GR24

To examine the effects of SLs in mesocotyl elongation, seedlings were germinated and grown on agar plates containing 0 and 1.0 μM of GR24 for 6 days in darkness. The length of mesocotyl of d10 mutant XJC was 2.3 fold longer than that of the wild-type GC13 seedlings (Figure 2A–C). GR24 did not affect mesocotyl elongation of wild-type seedlings but decreased the length of mesocotyl of d10 mutant XJC. At a concentration of 1.0 μM GR24, the length of mesocotyl between XJC and GC13 was indistinguishable (Figure 2A–C). This result suggests that mesocotyl length is negatively regulated by SLs in d10 mutant seedling under dark-growth conditions.
undetectable (Figure 3C–D). By means of mass spectrometry, seven protein spots (G1, G2, G3, G4, G6, G8 and G10) were successfully identified as eight different proteins (Table 1). Spot G4 was identified as the mixture of putative late embryogenesis abundant (LEA) protein and putative adenosine kinase (AK). Sucrose synthase 2 (SUS2) of spot G1 was up-regulated by GR24 treatment. The protein levels of putative vitamin B12-independent methionine synthase (MetE), LEA, AK, actin-7 (ACT7) and L-ascorbate peroxidase 2 (APX2) were down-regulated in XJC seedlings by exogenous SL application. Two protein spots, 26S proteasome regulatory subunit A (OsL_09330, G3) and mitochondrial phosphate translocator (MPT, G10) disappeared in XJC grown in darkness (Figure 3C–D). By means of mass spectrometry, seven protein spots (G1, G2, G3, G4, G6, G8 and G10) were successfully identified as eight different proteins (Table 1). Spot G4 was identified as the mixture of putative late embryogenesis abundant (LEA) protein and putative adenosine kinase (AK). Sucrose synthase 2 (SUS2) of spot G1 was up-regulated by GR24 treatment. The protein levels of putative vitamin B12-independent methionine synthase (MetE), LEA, AK, actin-7 (ACT7) and L-ascorbate peroxidase 2 (APX2) were down-regulated in XJC seedlings by exogenous SL application. Two protein spots, 26S proteasome regulatory subunit A (OsL_09330, G3) and mitochondrial phosphate translocator (MPT, G10) disappeared in XJC seedlings treated with GR24.

Identification of SL-responsive phosphoproteins

Phosphoserine- and phosphothreonine-specific antibodies have been used to identify and characterize phosphoproteins via immunoblotting analysis in rice [47,48]. In our research, total proteins were extracted from 6 day-old XJC seedlings that were grown in darkness in the presence of 1.0 μM GR24. Seedlings of the d10 mutant XJC grown in the absence of GR24 served as a control. Proteins were resolved by 2-DE, and probed with antibodies against phosphoserine (pS) or phosphothreonine (pT). A total of 9 proteins (spots S1 to S9) reacted with the pS antibody exhibited differential phosphorylation intensity between the control and GR24 treatment (Figure 4A–B). Two pT-positive proteins (spots T1 and T2) showed differential phosphorylation intensity between the control and GR24 treatment (Figure 4C–D). As shown in Figure 4E and Table 2, three differential patterns of phosphorylation intensity were distinguished: induced phosphorylation by GR24 treatment (S1, S3, S6, S8 and S9), induced de-phosphorylation by GR24 (S8 and T1) and reduced phosphorylation intensity by GR24 (S2, S7 and T1). A total of eight proteins were identified by mass spectrometry from seven spots (Table 2). Spot T1 was identified as a mixture of two proteins. Disappointedly, the identities of four phosphoprotein spots (S2, S7, S9 and T2) were not made successfully by mass spectrometry largely due to their low abundance (Table 2). Among the identified phosphoproteins, phosphoglycerate kinase (PGK, spot S1), triosephosphate isomerase (TPI, spot S3), phosphomannose mutase (PMM, spot S4), NADH:ubiquinone oxidoreductase (Complex I, spot S5) and cytoplasmic aconitase hydratase (ACO, spot S6) were found to be phosphorylated in the d10 mutant XJC seedlings induced by GR24 treatment (Figure 4A–B, Table 2). The aminotransferase AGD2 (AGD2, spot S8), and methylmalonate semi-aldehyde dehydrogenase (MSAH) and UDP-glucose pyrophosphorylase (UGP) of spot T11 were found to be dephosphorylated in the d10 mutant XJC seedlings after treatment with GR24 (Figure 4, Table 2).

Discussion

Inhibition of mesocotyl elongation of dark-grown d10 seedlings by SLs

We observed that mesocotyl elongation in darkness was greater in the d10 mutant than in the wild-type GC13. Exogenous application of GR24 restored the normal short mesocotyl phenotype in the d10 mutant XJC seedlings grown in darkness, which is consistent with the previous observation by Hu et al [38], showing that SLs negatively regulate cell division, but not cell elongation, in the mesocotyl during germination and growth of rice in darkness. This negative regulation mechanism of SLs in mesocotyl development might be related to the role of SLs in the axillary buds, and might explain the observations that shoot branching of the d mutants is enhanced. To understand the general roles of SLs in mesocotyl elongation and axillary branching, we applied the proteomic approach to analyze the total protein and phosphoprotein profiles differentially expressed in dark-grown d10 mutant XJC. A total of eight SL-responsive proteins were identified. They are mainly involved in carbohydrate metabolism, energy supply, defense responses, amino acid metabolism and cytoskeleton maintenance.
Differential expression of key enzymes of metabolic pathways in response to SL treatment

Sucrose synthase 2 (SUS2, spot G1 Table 1) was the only protein that was up-regulated in dark-grown XJC seedlings by the application of GR24. The expression levels of all other differentially regulated proteins were down-regulated or even became undetectable. Sucrose synthase is a key enzyme in sucrose degradation and in directing sucrose to different metabolic pathways. Our observation was consistent with a previous report on the up-regulation of sucrose synthase 2 in Arabidopsis in response to BR treatment [44]. It is likely that sucrose synthase 2 is an important target of phytohormones.

The protein level of adenosine kinase (ADK, spot G4 Table 1) was down-regulated by GR24 application. ADK catalyzes the salvage synthesis of adenine monophosphate from adenosine and ATP. In Arabidopsis, the role of ADK in metabolism has been investigated in transgenic plants expressing either sense or antisense ADK1 cDNA. Transgenic plants with less than 10% ADK activity show developmental abnormalities including reduced hypocotyl and root elongation [49]. ADK1 has also been shown to be involved in the development of root hairs and to negatively regulate systemic acquired resistance (SAR) via the jasmonic acid (JA) signaling pathway. We speculate that ADK may play a role as an inhibitor in the regulation of SL-related physiologic responses in rice.

Actin 7 (ACT7) is an important cytoskeleton protein in plants. Its differential expression has been found in the root proteome in rice treated with salt [50]. The down-regulated expression of ACT7 protein (spot G6, Figure 2) might be related to the inhibitory effect of SLs on cell growth.

Vitamin B12-independent methionine synthase (MetE) is required for the biosynthesis of methionine and tetrahydrofolate from homocysteine and 5-methyl tetrahydrofolate [51,52]. Methionine is an essential amino acid for protein biosynthesis and is also a precursor for S-adenosyl methionine (SAM), a methyl donor of various methylation reactions in cells. Under oxidative and temperature stress conditions, MetE is inactivated in bacteria and algae [53]. Our data showed that MetE expression level was down-regulated by GR24 (Figure 2, Table 1). It appears that MetE may represent a target of stress and phytohormones.

Figure 3. Differential expression of proteins in etiolated d10 seedlings treated with GR24. (a–b) 2-DE maps of total proteins extracted from XJC seedlings grown in the absence of GR24 (CK, control) or 1.0 μM GR24 (TR). (c) Magnified images containing differentially expressed protein spots from G1 to G10. (d) Abundance analysis of the differentially expressed protein spots determined using the PDQuest 2D analysis software (BioRad). Normalized quantitative volumes of protein spots were exported from PDQuest and graphical presentations of protein spots were generated using Excel software. Normalized volumes (N. volume x1000) of protein spots are shown in y-axes. The empty histograms represent spot volume means of the control (CK) and the solid histograms represent the treatments (TR).

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Table 1. Identification of differentially expressed proteins in response to SL treatment.

<table>
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<th>Spot</th>
<th>GenBank Accession</th>
<th>Protein name (abbreviation)</th>
<th>Mr(Da)/pI</th>
<th>MP</th>
<th>SC (%)</th>
<th>Mowse score</th>
<th>Exp. score</th>
<th>Protein change</th>
<th>FC ± SE</th>
<th>p-value</th>
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<td>25</td>
<td>83</td>
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<td>Up</td>
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<td>Vitamin B12-independent methionine synthase (MetE)</td>
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<td>44</td>
<td>214</td>
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<td>1.7±0.1</td>
<td>0.0027</td>
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<td>26S proteasome regulatory subunit 6A</td>
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<td>36</td>
<td>71</td>
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<td>59</td>
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<td>G8</td>
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*Rice d10 mutant XJC seedlings were germinated and grown on medium in the presence of 1.0 μM GR24 for 6 days in darkness. Seedlings grown without exogenous application of GR24 served as a control. Total proteins were extracted from seedlings and resolved by 2-DE. Protein spots that exhibited differences in intensity were excised for identification by mass spectrometry.

1Relative molecular mass (Mr) in Da and isoelectric point (pI) of proteins are theoretical.

2Number of matched peptides (MP).

3Sequence coverage (SC, %) of the identified peptides.

4Proteins with a Mowse score of more than 61 were considered to be identified at a statistically significant level. Significance threshold p<0.05.

5Expectation scores are the probability-based values obtained from database search by MASCOT. Values less than their threshold (p<0.05) are statistically confident with more than 95% certainty.

6Fold changes (FC) with standard errors (±SE) from three biological replicates were calculated as the ratio of normalized spot intensities between the treatment and control.

7Fold changes (FC) in protein spot intensity with a p-value of less than 0.05 were considered to be statistically significant.

doi:10.1371/journal.pone.0093947.t001
The G3 spot that was undetectable after GR24 treatment was identified as the hypothetical protein OsI_09330, which is homologous to maize 26S proteasome regulatory subunit 6A, a highly conserved protein in plants. It belongs to the P-loop nucleoside triphosphatase (NTPase) superfamily. By turning off the expression of the proteasome subunit, SLs may target the ubiquitin-mediated protein degradation machinery.

Late embryogenesis abundant (LEA) protein may protect plant cells from damages caused by stress [54]. The expression of LEA is known to be regulated by ABA, drought and wound [55]. L-Ascorbate peroxidase 2 (APX2) is an important antioxidant enzyme in protecting cells from oxidative stress. A previous report identifies APX2 as a differentially expressed protein in etiolated shoots of rice [56]. This is consistent with our observation that APX2 was found to be high in dark-grown rice seedlings, and was down-regulated in d10 mutant plants in response to GR24.

Because both LEA (spot 4) and APX2 (spot G8, Figure 2) were down-regulated by GR24, it appears that SLs may regulate plant growth and development through regulation of enzymes that function in defense against oxidative stress.

Mitochondrial phosphate translocator (MPT, spot G10) was found to be highly expressed in the dark-grown d10 mutant seedlings, but became undetectable upon GR24 treatment (Figure 3, Table 1). This differential expression might be related to the inhibitory effect of GR24 on cell division. MPT is present as a homodimer in the inner mitochondrial membrane and transports inorganic phosphate (Pi) to the mitochondrial matrix. It is a symporter of phosphate (H2PO4−) and proton (H+) and

**Figure 4. Phosphoproteins of d10 seedlings in response to GR24 treatment.** Total proteins were extracted from 6 day-old dark-grown d10 mutant (XJC) seedlings in the absence (CK, the control) or presence of 1.0 μM GR24 (TR), and probed with anti-phosphoserine (a–b) or anti-phosphothreonine (c–d) antibodies. Phosphoproteins that changed phosphorylation intensities in response to GR24 treatment are indicated by S1–S9 and T1–T2. Intensity changes of phosphorylation status were quantified with the PDQuest 2D analysis software (e). Normalized volumes (N, volume x 1000) of phosphoprotein spots in the control (CK) or GR24 treated samples (TR) were presented.

doi:10.1371/journal.pone.0093947.g004
Table 2. Identification of proteins that changed phosphorylation intensities in response to SL treatment.  

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5**Proteins with a Mowse score of more than 61 were considered to be identified at a statistically significant level. Significance threshold p<0.05.

6**Expectation scores are the probability-based values obtained from database search by MASCOT. Values less than their threshold (p<0.05) are statistically confident with more than 95% certainty.

7**Protein phosphorylation intensity changes were distinguished as induced phosphorylation (+), induced de-phosphorylation (-), and reduced phosphorylation (↓) in response to GR24 treatment.

8**Fold changes (FC) with standard errors (±SE) from three biological replicates were calculated as the ratio of normalized spot intensities between the treatment and control.

9**Fold changes (FC) in spot intensity with a p-value of less than 0.05 were considered to be statistically significant.

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plays an essential role in the oxidative phosphorylation from ADP to ATP in mitochondria [57–59]. A recent report has suggested that elevated SL levels by phosphate starvation contribute to the inhibition of tiller bud outgrowth in rice seedlings [60]. It is speculated that during phosphate deficiency, SLs may function as a rhizosphere signal to maximize AM fungi symbiosis for improved phosphate acquisition and may also serve as an endogenous hormone to optimize shoot branching for efficient \( \Pi \) utilization. Our data showed that MTP disappeared in \( d10 \) seedlings after SL treatment (Figure 3). This suggests that SLs may have a more complex regulation role in the \( \Pi \)-related metabolism and cell division control in rice.

Potential targets of the SL-signaling pathway

Protein phosphorylation has been recognized as an important mechanism for cell signaling and regulation of cell division and plant growth [61]. A great number of phosphoproteins have been implicated in the signaling pathways of phytohormones including GA, ABA and BR [42–44,47,62]. SLs represent a newly identified phytohormone. Whether phosphoproteins are involved in the signaling pathway of SLs remain to be explored. A total of 11 phosphoprotein spots were found to be responsive to GR24 treatment (Figure 4, Table 2). Eight proteins were identified by MALDI-TOF/MS, including six phosphoserine-containing proteins and two phosphothreonine-containing proteins. Unfortunately, four phosphoprotein spots (S2, S7, S9 and T2) were not identified successfully by mass spectrometry probably due to their low abundance. The signal intensity on X-ray films of Western blotting analysis using anti-phosphoserine/threonine antibodies was proportional to the phosphorylation levels of a protein, but not to the protein content. Despite of the high intensity on Western blots, these four phosphoprotein spots were faint on 2-DE gels (Figure 4). As the mass spectrometry technology continues to improve, these relatively low abundant proteins may be identified in future work.

Phosphoglycerate kinase (PGK) and triosephosphate isomerase (TPI) both are key enzymes in glycolysis. PGK catalyzes the conversion of 1,3-diphosphoglycerate to 3-phosphoglycerate, coupling with substrate level phosphorylation of ADP to ATP. TPI catalyzes the inter-conversion between dihydroxyacetone phosphate (DHAP) and 3-phosphoglyceraldehyde (PGA) in the glycolytic pathway. Phosphorylation of TPI has been found in the proteomes of rice leaves treated with exogenous ABA [47]. This suggests that TPI may be a target of phytohormone signaling. Our result is also consistent with a recent report on phosphoproteins in rice responsive to different hormones/stresses, which has concluded that the glycolytic pathway enzymes are the main targets of phosphorylation events [63].

Phosphomannose mutase (PMY), cytoplasmic aconitate hydratase (ACO), UDP-glucose pyrophosphorylase (UGP), and mitochondrial NADH:ubiquinone oxidoreductases (Complex I) are all important enzymes involved in the carbohydrate metabolism and ATP biosynthesis. PMY is the key enzyme in mannose metabolism and contributes to the biosynthesis of ascorbic acid,GDP mannose and mannosyl monophosphopolyrenol [64,65]. ACO catalyzes the reversible isomerization between cis-aconitic acid and isocitric acid, and the cytoplasmic ACO has been implicated in multiple physiological processes including cytosolic citrate metabolism, the glyoxylate cycle, iron homeostasis and oxidative stress resistance. Complex I of the mitochondrial inner membrane consists of multiple subunits and provides a link of the tricarboxylic acid cycle with the oxidative phosphorylation. It uses the energy from the oxidation of NADH to produce reduced form of ubiquinone and generate a proton gradient that drives ATP biosynthesis.

UDP-glucose pyrophosphorylase (UGP) plays an important role in the synthesis of uridine diphosphoglucose (UDPG), which serves as the glucose donor for the biosynthesis of sucrose, cellulose, callose and cell wall hemicelluloses. Although the protein expression level of UGP did not change between control and GR24 treatment, the degree of de-phosphorylation of UGP was found to be responsive to GR24 (Figure 4, Table 2).

Aminotransferases transfer the amino group from an \( \alpha \)-amino acid to an \( \alpha \)-ketone acid, generating a new amino acid. A previous study has showed that the \( \alpha \)-amino acid substrate(s) has yet to be defined [67]. The AGD2 aminotransferase appears to be involved in the synthesis of an essential amino acid-derived molecule affecting development and defense [67]. The roles of AGD2 phosphorylation changes in plant growth in response to SLs need further investigation.

Methylmalonate semialdehyde dehydrogenases (MSDH) belong to the CoA-dependent aldehyde dehydrogenase superfamily and catalyze the irreversible oxidative decarboxylation reaction of methylmalonate semialdehyde, generating NADH and propanoyl-CoA. MSDH may participate in the pathways of inositol, amino acid and propanoate metabolism. How the phosphorylation changes of MMDSH is related to branching and mesocotyl elongation in response to SL treatment remains to be investigated.

Understanding protein phosphorylation and its regulation is a crucial part of functional biology of plants. To investigate whether the eight SL-regulated phosphoproteins contain a conserved phosphorylation-site, we analyzed these proteins using Phospho-protein BLAST in P3DB [http://www.p3db.org/] [68]. Except for S4 (Q7XPW5, phosphomannose mutase), other seven rice proteins have at least one orthologous phosphoprotein in Arabidopsis thaliana, Medicago truncatula and Vitis vinifera identified by different research groups [69–76]. Among them, S1 (Q6H6C7, phosphogllycerate kinase) and T1(1) (Q6Z4E4, methylmalonate semialdehyde dehydrogenase) have been identified as phosphoproteins by large-scale comparative phosphoproteomics in rice [69]. Eight orthologous accessions of S1 in the database include four from Oryza sativa, one from Arabidopsis thaliana and three from Medicago truncatula. The phosphorylation site of S1 (Q6H6C7) has been identified as serine-201 in rice [69].

It has recently been shown that depletion of the auxin efflux carrier PIN1 from the plasma membrane occurs within 10 minutes after SL treatment and does not depend upon protein synthesis [77]. This suggests that a post-translational regulatory mechanism, rather than a transcriptional cascade, may carry out the initial responses to SL perception. As an important post-translational modification, protein phosphorylation and de-phosphorylation might have rapid effects on enzyme activity, structural changes and subcellular localization. Thus protein phosphorylation status might be an important target of SL action.

Despite of the successful identification of phosphorylation changes of several proteins, some known SL-related signaling components such as D3 (an F-box protein) and D14 (an \( \alpha \)-/\( \beta \)-fold hydrolase) were not detected in our study. It is possible that they are present in low abundance or the limited pHi range of IPG strips are not suitable for their identification.

D3/MAX2 encodes an F-box protein that is highly conserved among land plants [78,79]. F-box proteins serve as the adapter components of SCF E3 ubiquitin ligase complexes, and confer substrate specificity for ubiquitination. Target proteins recognized...
by the F-box protein are typically polyubiquitinated and subsequently degraded by the 26S proteasome [80]. There are remarkable similarities in protein components between the SL and gibberellin signaling pathways. Both the gibberellin receptor and the putative SL receptor D14 are members of the \( \alpha/\beta \) hydrolase protein family, and in both cases, the hormone signaling requires the function of an F-box protein [81–83]. Two recent reports on rice D53, a member of the double CIP-N motif-containing P-loop nucleoside triphosphate hydrolase superfamily, further reveal similarities between the signaling mechanisms between SL and gibberellins [84,85]. D53 functions as a repressor of SL signaling and a target of D3/MAX2. Perception of SL by D14 and the SCF\(^{D3} \) complex leads to ubiquitination of D53 and its subsequent degradation by the ubiquitin proteasome system, which in turn releases the repression of downstream target genes [84,85]. As an important component of the ubiquitin proteasome system, 26S proteasome regulatory subunit 6A (spot G3) was identified in this study as a differentially expressed protein in response to GR24 treatment. We speculate that it might play a role in the ubiquitin-mediated protein degradation machinery in the SL signaling pathway. It remains to be determined whether this 26S proteasome regulatory subunit interacts with the D3-D14-D53 complex, and if it plays a role in the ubiquitin-mediated protein degradation machinery in the SL signaling pathway.

In summary, the 8 differentially expressed proteins responded to SLs did not overlap with the 8 proteins that exhibited phosphorylation status changes. Thus, different approaches have apparent different sensitivities and specificities in the identification of proteins in response to SL treatment. An integrative approach to combine the use of multiple techniques may complement with each other and lead to better identification of molecular changes in rice plants. It will be of interest to determine the exact functions of the SL-regulated proteins identified in this work. Due to the inherent bias of the 2-DE approach toward the identification of more abundant proteins, components of signaling pathways with lower abundance would be more difficult to detect. Enrichment of these low abundant proteins using pre-fractionation methods and application of more sensitive technologies will be helpful. This work represented an attempt to use proteomic approach in identification of proteins that responded to SL treatment in rice, and may pave an avenue for future investigations aimed at exploration of new signaling pathways in plants.

Materials and Methods

Plant materials and growth conditions

Rice (Oryza sativa L. ssp. indica) cultivar GC13 (the wild-type) and its \( d10 \) mutant line XJC (\( d10 \)) were used in this research. XJC was originally isolated from a mutation population generated by \( \gamma \)-ray irradiation. After inbreeding for several generations, XJC attained genetic stability and exhibited semi-dwarf plant height and a high-tillering phenotype. More than 120 tillers could be observed in XJC at the maturing stage [46].

For the phenotype rescuing experiment, pregerminated rice seeds were grown hydroponically in growth chambers and cultured at 30°C under fluorescence white light with a 16 h light/8 h dark photoperiod for two weeks. The seedlings were maintained in the Kimura B hydroponic solution [86].

For the mesocotyl elongation experiment, rice seedlings were grown under conditions as described elsewhere [38] with minor modifications. Dehusked rice caryopses were sterilized in a 10% \( (v/v) \) sodium hypochlorite solution for 1 h. After washing with deionized water, seeds were kept in water for 12 h at room temperature in darkness. Each agar plate (0.7%, w/v) was planted with 6 seeds, and placed in a box covered by black cloths. The box was kept in a growth chamber at 30°C for 6 days.

GR24 treatment and analysis

GR24, kindly provided by Prof. S. Yamaguchi, was dissolved in 100% acetone as a 100 mM stock solution, and was added to the hydroponic culture medium in a final concentration of 1.0 \( \mu \)M. Hydroponic culture medium with 0.001% acetone served as a negative control. For the mesocotyl elongation experiment, GR24 (100 mM stock solution in acetone) was added to melted 0.7% (w/v) agar medium (about 30°C) in a final concentration of 1.0 \( \mu \)M GR24. Acetone at a final concentration of 0.001% was present in each agar plate including the mock treatment. Sterilized seeds were placed on agar plates and seedlings were grown at 30°C for 6 days under complete darkness. The mesocotyl length of wild type GC13 and \( d10 \) mutant XJC seedlings were recorded. Total proteins were extracted from XJC seedlings with or without GR24 treatment, and analyzed by the proteomic approach.

Preparation of total proteins

Etiolated XJC seedlings, cleared off residual seeds, were quickly frozen in liquid nitrogen and stored at \(-80^\circ\)C. Total proteins were extracted in extraction buffer and precipitated by trichloroacetic acid as described previously [87]. Protein pellets were dissolved in a final concentration of approximately 25 mg/ml protein in the O’Farrells UKS buffer (9.5 M urea, 3 mM K$_2$CO$_3$, 1.25% sodium dodecyl sulfate [SDS], 0.5% DTT, pH 3.5 to 10.0 2% pharmalyte, 6% Triton X-100) [39]. To facilitate protein dissolution, the pellet-buffer mixture was shaken gently at room temperature for 1 h, followed by centrifugation at 15,000 \( g \) at 15°C for 30 min. Protein content of the soluble fraction was determined using the BCA Protein Assay (Pierce). The soluble fraction was used for two-dimensional polyacrylamide gel electrophoresis (2-DE) or stored at \(-80^\circ\)C. Because most of the proteins were found within pl 4–7 on 2D gels (Supplemental Figure S1), precast immobilized \( pH \) gradient [IPG] strips (pH 4–7, 7 cm, Bio-Rad) were used in the first dimension of electrophoresis. Proteins (approximately 100 \( \mu \)g) were loaded on each strip. After isoelectric focusing electrophoresis, the IPG strips were soaked in equilibration buffer and placed directly onto 12% polyacrylamide-SDS slab gels. After electrophoresis, the gels were immuno-blotted using a phospho-ami no-acid-specific antibody. Identical gels were also stained by Coomassie blue or silver staining solution.

Immunoblot analysis

Proteins resolved on 2-DE gels were transferred onto PVDF membranes (Millipore). The membranes were blocked with 5% \( (w/v) \) bovine serum albumin (BSA) in TBST (20 mM Tris–HCl, pH 7.5, 150 mM NaCl, 0.05% \( (v/v) \) Tween-20) for overnight at 4°C, and then incubated with mouse anti-phosphoserine (Millipore) or rabbit anti-phosphothreonine antibodies (Cell Signaling) at 1 \( \mu \)g/ml in TBST for 1.5 hr. After washing with TBST, membranes were incubated with goat anti-mouse IgG or donkey anti-rabbit IgG (Protein Tech Group) conjugated with horseradish peroxidase at a 1:4000 dilution for 1 h at room temperature. Immunoreactive spots were detected by enhanced chemiluminescence (GE) followed by exposure to X-ray film according to the manufacturer’s instructions.

Image acquisition and data analysis

Stained gels and developed films were scanned with the Molecular Imager ChemiDoc XRS System (Bio-Rad, USA) in transmission mode. All 2-DE gel images were captured,
digitalized, and analyzed with PDQuest software (Bio-Rad) using tenfold over background as a minimum criterion for presence/absence. The analysis was re-evaluated by visual inspection. The spots present in all three biological replicates showing statistically significant, qualitative or quantitative differences between treatment (TR) and control (CK) were drawn after comparisons. On the basis of total quantity and intensity of valid spots in each 2D-gel, a normalization factor was generated. A normalized spot volume was determined for each protein spot. Data analysis was performed with Excel software (Microsoft). Error bars shown in figures represent the standard deviation values.

Mass spectrometry analysis and database search

Selected protein spots were cut from the gel and digested with sequencing-grade trypsin (Promega). The peptides were dissolved in 2 μL of 0.5% trifluoroacetic acid (TFA) solution and mixed with an equal volume of a saturated matrix solution consisting of α-cyanо-4-hydroxy-cinnamic acid in 70% acetonitrile with 0.1% TFA. The resulting mixture was loaded onto the point template. After drying at room temperature, the template was introduced into a Bruker Reflex™ matrix-assisted laser-desorption ionization-time of flight mass spectrometry (MALDI-TOF MS) system (Bruker, Germany). The peak lists of peptide mass fingerprints were created using the FlexAnalysis 2.0 software (Bruker Daltonics), and interpreted with Mascot (Matrix Science, London, UK. http://www.matrixscience.com/) against the MSDB and NCBI databases.

Supporting Information

Figure S1 Comparison of protein distribution patterns separated by different precast immobilized pH gradient (IPG) strips. Total proteins isolated from rice seedlings were separated using IPG strips with a pH range of 3–10 (left) and pH 4–7 (right). Note that the majority of proteins were present within the pH range of 4–7. (PDF)

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Author Contributions

Conceived and designed the experiments: FC HW ZH YH. Performed the experiments: FC IJ JZ RH. Analyzed the data: FC. Contributed reagents/materials/analysis tools: FC. Wrote the paper: FC HW ZH YH.

References


