Cytological and proteomic analyses of horsetail (Equisetum arvense L.) spore germination

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Spermatophyte pollen tubes and root hairs have been used as single-cell-type model systems to understand the molecular processes underlying polar growth of plant cells. Horsetail (Equisetum arvense L.) is a perennial herb species in Equisetopsida, which creates separately growing spring and summer stems in its life cycle. The mature chlorophyllous spores produced from spring stems can germinate without dormancy. Here we report the cellular features and protein expression patterns in five stages of horsetail spore germination (mature spores, rehydrated spores, double-celled spores, germinated spores, and spores with protonemal cells). Using 2-DE combined with mass spectrometry, 80 proteins were found to be abundance changed upon spore germination. Among them, proteins involved in photosynthesis, protein turnover, and energy supply were over-represented. Thirteen proteins appeared as proteoforms on the gels, indicating the potential importance of post-translational modification. In addition, the dynamic changes of ascorbate peroxidase, peroxiredoxin, and dehydroascorbate reductase implied that reactive oxygen species homeostasis is critical in regulating cell division and tip-growth. The time course of germination and diverse expression patterns of proteins in photosynthesis, energy supply, lipid and amino acid metabolism indicated that heterotrophic and autotrophic metabolism were necessary in light-dependent germination of the spores. Twenty-six proteins were involved in protein synthesis, folding, and degradation, indicating that protein turnover is vital to spore germination and rhizoid tip-growth. Furthermore, the altered abundance of 14-3-3 protein, small G protein Ran, actin, and caffeoyl-CoA O-methyltransferase revealed that signaling transduction, vesicle trafficking, cytoskeleton dynamics, and cell wall modulation were critical to cell division and polar growth. These findings lay a foundation toward understanding the molecular mechanisms underlying fern spore asymmetric division and rhizoid polar growth.

Keywords: spore germination, Equisetum arvense L., fern, proteomics, single cell, polar growth

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Introduction

Sexual reproduction is crucial in plant life cycle. Spermatophyte seeds, pollen grains, and fern spores play central roles in sexual reproduction with common capability of surviving under harsh conditions upon emergence from dormancy. They have physiological resemblance with each other as to features of resurrection and polar growth during the emergence of roots, pollen tubes, and rhizoids, respectively. Previous reports have addressed the physiological and molecular characteristics of pollen and seed germination using genomics and proteomics approaches (Dai et al., 2007a,b; Tan et al., 2013), but comparable information is lacking for fern spore germination. The similarity of germination among spermatophyte pollen grains, seeds, and fern spores includes mobilization and organization of limited reserves for polar growth within a short time frame. Previously, 33 and 123 genes were found to be commonly expressed when Ceratopteris richardii spores were compared with Arabidopsis thaliana pollen and seeds, respectively (Bushart and Roux, 2007; Salmi et al., 2007). Some proteins encoded by these genes (e.g., Rop GTPase, Mago nashi, calmodulin 2, No pollen germination 1, phospholipase D, synaptobrevin, and constitutive photomorphogenic 9) are mainly involved in calcium signaling, vesicle trafficking, and ubiquitin-mediated protein degradation. Although the protein functions encoded by some of the important genes (e.g., Rop GTPase, calmodulin, and phospholipase D) in pollen grains have been well-studied (Malhó et al., 2006), their roles in fern spore germination are not clear. In spite of polar growth similarities, pollen and fern spores are evolutionarily distinct. The pollen grains are reduced male gametophytes with two/three cells to reach ovary by tip-growing pollen tube after recognition on appropriate stigma, while the mononuclear fern spores can generate live-independent gametophyte through germination (Dai et al., 2008). Fern spore germination is somewhat more complex in that they undergo extensive cell division and differentiation during the emergence of rhizoid and prothallus under various environmental conditions (Bushart and Roux, 2007). Fern spores represent a new single-celled model for investigating asymmetric cell division, differentiation, and polar growth (Chatterjee et al., 2000; Bushart and Roux, 2007).

Previous physiological studies have reported that spore germination of more than 200 fern species was modulated by various environmental factors, such as light, gravity, calcium, phytohormones, and temperature (Suo et al., 2015). Recently, gene function analyses revealed that several signaling pathways are crucial for fern spore germination. Phytochrome and cryptochrome signaling is important for the first spore cell mitosis (Suetsugu and Wada, 2003; Kamachi et al., 2004; Tsuibo et al., 2012). Besides, gravity and calcium signaling determines polarity establishment, cell asymmetry division, and the direction of rhizoid elongation (Chatterjee et al., 2000; Bushart and Roux, 2007; Bushart et al., 2014), and nitric oxide functions as a signal molecule in the regulation of gravity-directed fern spore polarity (Salmi et al., 2007). Moreover, gibberellin and antheridiogen can initiate and promote spore germination in many species, but abscisic acid, jasmonic acid, and ethylene have only minor promoting effects (Suo et al., 2015). In addition, the enzymes involved in glyoxylate cycle (e.g., isocitrate lyase and malate synthase) and genes encoding aconitase in the conversion from citrate to isocitrate were induced in germinating spores from Onoclea sensibilis (DeMaggio and Stetler, 1980), Anemia phyllitidis (Gemmrich, 1979), and C. richardii (Banks, 1999), indicating the importance of lipid reserve degradation and mobilization during spore germination. However, the molecular regulatory mechanisms in these processes are still unknown.

Germination of fern spores, especially the chlorophyllous spores (green spores), is a sophisticated signaling and metabolic process. Spores from Equisetum species are chlorophyll-bearing and of short viability (only a few weeks) (Ballesteros et al., 2011). Equisetum spores can germinate immediately under appropriate conditions with high humidity (Lebkuecher, 1997), therefore, they are good materials for studying the chlorophyllous spore germination. Equisetum is the oldest living genus of vascular plants, containing 15–25 extant hollow-stemmed taxa (Guillon, 2007). Most Equisetum species are regarded as persistent weeds in wetlands (Large et al., 2006). The reproduction of Equisetum species is mainly dependent on the growth of the rhizomes underground, but not the spore germination. Large-scale comparative proteomics of developing rhizomes of Equisetum hyemale have revealed that 1911 and 1860 proteins in rhizomes apical tip and elongation zone, respectively (Balbuena et al., 2012). However, the cellular and proteomic features of spore germination are still lacking. In the present study, we carried out cellular and 2-DE based proteomics analysis of horsetail (Equisetum arvense L.) spore germination to reveal the signaling and metabolic characteristics.

Materials and Methods

Collection and Germination of Mature Horsetail Spores

The mature fertile sporophylls (the separate stalks in the spring) of horsetail (E. arvense) were collected in suburb of Harbin (45° 27′ N, 127° 52′ E), Heilongjiang province, China. The mature spores (MS) were released by its elaters moving from the sporangia during dehydration at room temperature (25°C). To synchronize spore germination, the collected fresh MS were soaked in deionized H2O overnight in the dark. Then, the rehydrated spores (RS) were transferred into a liquid germination medium (4.6 mM Ca(NO3)2·4H2O, 2 mM KNO3, 1.5 mM KH2PO4, 0.8 mM MgSO4·7H2O) and cultured in an...
environmentally controlled chamber at 25 ± 2°C, 24 h light at 60 µmol·m⁻²·s⁻¹. The amount of germinating spores at various stages of RS, double-celled spores (DCS), germinated spores (GS), and spores with protonemal cells (SPC) was examined under a microscope. All these spores were collected by centrifugation at 500 × g for 5 min, and used immediately after collection for protein extraction or storage at −80°C.

Observation of Spore Morphology Upon Germination
Morphological characteristics of spores upon germination were examined under a Axioskop 40 fluorescence microscope (Zeiss, Oberkochen, Germany) without or with staining using 0.2 µg·mL⁻¹ 4,6-diamidino-2-phenylindole (Molecular Probes, Carlsbad, USA). Observation of MS was performed under HITACHI S-520 scanning electron microscope (Hitachi, Tokyo, Japan) (Dai et al., 2002). Living MS were prepared by standard techniques for scanning electron microscope observation. Spores were fixed, gradually dehydrated, and critically point dried by Balt-Tec CPD-030 critical point dryer (Balt-Tec AG, Balzers, Liechtenstein) according to Dai et al. (2002). Dry spores were mounted on aluminum stubs using double-sided tape and coated with gold-palladium using a Denton Vacuum Desk II sputter coater (Denton Vacuum Inc., Cherry Hill, USA).

Protein Extraction and Quantification
For protein preparation, 0.5 g spores in the five stages of germination (MS, RS, DCS, GS, and SPC) were ground to powder in liquid nitrogen using chilled mortar and pestle. Total protein of spores was extracted according to the method of Wang et al. (2010). Protein samples were prepared independently from three different batches of plants, considered as three biological replicates. Protein concentration was determined using a Quant kit according to manufacturer’s instructions (GE Healthcare, Salt Lake City, USA).

2-DE, Gel Staining and Protein Abundance Analysis
The protein samples were separated and visualized using 2-DE according to Dai et al. (2006). An aliquot of 1.3 mg total protein was diluted with rehydration buffer (7 M urea, 2 M thiourea, 0.5% CHAPS, 20 mM DTT, 0.5% immobilized pH gradient IPG buffer 4–7, and 0.002% bromphenol blue) to a final volume of 450 µL and loaded onto an IPG strip holder containing a 24 cm, pH 4–7 linear gradient IPG strip (GE Healthcare, Salt Lake City, USA). Isoelectric focusing was performed in the Ettan IPGphor isoelectric focusing system (GE Healthcare, Salt Lake City, USA) following the protocol of the manufacturer. For SDS-PAGE, the equilibrated IPG strips were transferred onto 12.5% acrylamide gels by using an Ettan DALT Six Electrophoresis Unit (GE Healthcare, Salt Lake City, USA). The gels were stained by Coomassie Brilliant Blue. Gel image acquisition and analysis were conducted as previously described (Wang et al., 2010). Images were acquired by scanning each stained gel using an ImageScanner (GE Healthcare, Salt Lake City, USA) at a resolution of 300 dpi and 16-bit grayscale pixel depth, and then analyzed using ImageMaster 2D version 5.0 (GE Healthcare, Salt Lake City, USA). The experimental molecular weight of each protein was estimated by comparison with the coseparated molecular weight markers. The experimental pI of each protein was determined by its migration on IPG linear strips. For quantitative analysis, the average vol% values were calculated from three biological replicates. Protein spots with reproducible and statistically significant changes in intensity (greater than 1.5-fold and p < 0.05) were considered to be differentially abundant protein (DAP) spots.

Protein Identification Using Mass Spectrometry and Database Searching
The DAP spots were manually excised from the 2-DE gels, and the in-gel digestion was performed as described previously (Dai et al., 2006). Tandem mass spectrometry spectra were acquired on a ESI-Q-TOF mass spectrometry (QSTAR XL) and a ESI-Q-Trap mass spectrometry (Applied Biosystems, Foster City, USA) (Wang et al., 2010; Yu et al., 2011). The tandem mass spectrometry spectra were searched against the NCBI non-redundant protein database (http://www.ncbi.nlm.nih.gov/) using Mascot software (Matrix Science, London, UK), according to the searching criteria described previously (Yu et al., 2011). The taxonomic category was green plants (3,019,757 sequence entries), mass accuracy was 0.3 Da, and the maximum number of missed cleavages was set to one. To obtain highly confident identification, proteins had to meet the following criteria: (1) the top hits on the database searching report, (2) a probability-based MOWSE score greater than 43 (p < 0.05), and (3) more than two peptides matched with nearly complete y-ion series and complementary b-ion series. The mass spectrometry proteomics data have been deposited to the ProteomeXchange Consortium (Vizcaino et al., 2014) via the PRIDE partner repository with the dataset identifier PXD002218.

Protein Classification and Hierarchical Clustering Analysis
For function classification of DAPs, the peptide sequences of each DAP were aligned against the Gene Ontology (GO) protein database (http://geneontology.org/) following the policies and procedures provided by the GO Consortium (http://geneontology.org/). The proteins were classified according to their cellular component, molecular function, and biological process. Also, PSI and PHI-BLAST programs (http://www.ncbi.nlm.nih.gov/BLAST/) were used to search against the NCBI non-redundant protein database for protein functional domain annotation. Besides, the biological function of protein was obtained from the Kyoto Encyclopedia of Genes and Genomes pathway database (http://www.kegg.jp/kegg/). In addition, the conservative protein function during spore germination was predicted from previous publications on the germinating seeds and pollen. Finally, by integrative analysis of all the information collected from aforementioned processes, each DAP was classified into certain functional category defined by us. The definition of functional category is referred from literatures on pollen and seed germination (Supplementary Figure S1). Log (base 2) transformed ratios were used for hierarchical clustering analysis using Cluster 3.0 available on
the Internet (http://bonsai.hgc.jp/~mdehoon/software/cluster/software.htm). Protein abundance ratio was calculated as protein abundance at MS stage divided by abundance at each stage. Using a tree algorithm, these DAPs were organized based on similarities in the expression profile. These proteins can be joined by very short branches if they are very similar to each other, and by increasingly longer branches as their similarity decreases. Java TreeView (http://jtreeview.sourceforge.net/) was used for data visualization.

**Protein Subcellular Location and Protein-protein Interaction Analysis**

The subcellular location of the identified proteins was predicted using five internet tools: (1) YLoc (http://abi.inf.uni-tuebingen.de/Services/YLoc/webloc.cgi), confidence score ≥ 0.4; (2) LocTree3 (https://rostlab.org/services/loctree3/), expected accuracy ≥ 80%; (3) ngLOC (http://genome.unmc.edu/ngLOC/index.html), probability ≥ 80%; (4) TargetP (http://www.cbs.dtu.dk/services/TargetP/), reliability class ≤ 3; (5) Plant-mPLoc (http://www.csbio.sjtu.edu.cn/bioinf/plant-multi/), no threshold value in Plant-mPLoc. Only the consistent predictions from at least two tools were accepted as a confident result. For the inconsistent prediction results among five tools, subcellular localizations for corresponding proteins were predicted based on literatures.

The protein-protein interactions were predicted using the web-tool STRING 9.1 (http://string-db.org). The DAPs homologs in Arabidopsis were found by sequence BLASTing in TAIR database (http://www.arabidopsis.org/Blast/index.jsp). The homologs were subjected to the molecular interaction tool of STRING 9.1 for creating the proteome-scale interaction network.

**Statistical Analysis**

All the results were presented as means ± standard deviation of three biological replicates. Data were analyzed by One-Way ANOVA using the statistical software SPSS 17.0 (SPSS Inc., Chicago, USA). The mean values from different stages of spore germination were compared by least significant difference post hoc test. A p-value less than 0.05 was considered statistically significant.

**Results and Discussion**

**Horsetail Chlorophyllous Spore Germination Process**

Chlorophyllous spores from fern species in a few unrelated taxa in Pteridophyta can germinate in less than 3 days and their viability lasts about 1 year or less (Lloyd and Klekowski, 1970). *Equisetum* spores are typical chlorophyllous spores with short viability, which can germinate immediately under humid conditions and remain viable for about 2 weeks (Lebkuecher, 1997). In this study, the spores were sown and cultured on Knop's medium under two different illumination levels (i.e., 30 µmol m⁻²·s⁻¹ and 60 µmol m⁻²·s⁻¹) after 12 h dark imbibition in deionized H₂O at room temperature. We found that the light illumination level has obvious effects on horsetail spore germination rate. At 16 h after illumination (HAI), the germination rate of spores under higher illumination level (60 µmol m⁻²·s⁻¹) was over 50%, but the germination rate of spores under lower illumination level (30 µmol m⁻²·s⁻¹) was only 20%. The spores under the higher light intensity reached to the maximum germination rate of 95% at 32 HAI, but the maximum germination rate of spores under 30 µmol m⁻²·s⁻¹ was only 35% (Supplementary Figure S2). Thus, the 60 µmol m⁻²·s⁻¹ light was used for horsetail spore germination. Actually, the horsetail MS have initiated germination during the process of dark imbibition. Eighty-seven percent of the RS finished their cell nucleus polar migration toward the gravity, and the nucleus of spore started their division at 12 HAI for the preparation of asymmetric cell mitosis (Figure 1). At 8 HAI, 82% spores finished the first mitosis to generate DCS containing a larger cell and a smaller cell, and the smaller cell of 4% spores elongated to form an obvious rhizoid. At this time point, we started to calculate the spore germination rate that is defined as the ratio of GS number to total spore number. At 18 HAI, more than 70% of spores generated the rhizoids, being defined as GS. At 32 HAI, spores obtained the maximum germination rate of over 95%, and the larger cell (protonemal cell) of over 87% spores have finished the second mitosis to give rise to the photosynthetic prothallus. The spores at this stage were defined as SPC (Figure 1).

**Cytological Characteristics of Horsetail Germinating Spores**

The horsetail chlorophyllous spores were generated in sporangium from strobili. The MS were released from partially dried strobili. Horsetail spores are unusual in the morphological and physiological aspects. The spores are typically about 25–35 µm in diameter with four flexible ribbon-like elaters about 100 µm each (Figures 2A,E,I,Q). The elaters initially wrapping around the spore body can deploy upon dehydration and fold back in humid air. The elater movement driven by humidity variations led to the spore exiting from sporangium, and especially can catch the wind again when they were jumping from the ground, which is believed to be a novel type of efficient self-propelled dispersal mechanism (Marmottant et al., 2013). The single nuclear was visible clearly in the center of the spores (Figures 2E,I). The elaters were lost when spores were cultured in liquid medium and cell nucleus has migrated toward gravity (Figures 2N,P). After 12 h dark imbibition, the RS were swelled to the diameter of about 50 µm. In RS, the nucleus has completed the first mitosis and two nuclei in the bottom side of the spore can be clearly observed (Figures 2B,F,J). Subsequently, at 8 HAI, the novel cell wall was formed between the two nuclei to generate an asymmetric DCS, containing a larger cell and a smaller cell (Figures 2C,G,K). The smaller cell started to elongate out of the spore and an approximate 30 µm long rhizoid emerged at 18 HAI to form GS (Figures 2D,H,L). In the GS, the cell nucleus of rhizoid still left inside the spores (Figures 2H,I). As rhizoid elongated, its nucleus was moving outside from the spore to the center of the rhizoid at 28 HAI (Figures 2O,R). At the same time, the larger cell inside the spore can complete the second mitosis to form a new protonemal cell (Figures 2O,R). Thus, the spore finish its germination by the formation of...
FIGURE 1 | Germination time course of spores from E. arvense.
Mature spores (MS) (open rhombuses) were cultured on Knop’s medium. After 12 h dark imbibition, 87% of the rehydrated spores (RS) (open squares) completed rehydration, which were enlarged with an open trilet aperture. Then, spores were cultured under continuous illumination. At 8 h after illumination (HAI), 82% spores completed first mitosis to form double-celled spores (DCS) (open triangles). At 18 HAI, over 70% spores generated rhizoids, which were defined as germinated spores (GS) (crosses). At 32 HAI, more than 87% spores finished the second cell division, which were called spores with protonemal cells (SPC) (open circles). The spore germination rate (filled rhombuses) was increased gradually in this time course, and the maximum germination rate was over 95% at the stage of SPC. Error bar indicates ± standard deviation.

DAPs Upon Spore Germination
To determine DAPs in mature and germinating spores, protein samples collected at the five stages (MS, RS, DCS, GS and SPC) were subjected to 2-DE analysis. On the Coomassie Brilliant Blue-stained gels (pH 4–7, 24 cm IPG strip), 1243 ± 47, 1247 ± 34, 1254 ± 31, 1229 ± 57, and 1234 ± 40 spots from MS, RS, DCS, GS, and SPC were detected, respectively (Figure 3, Supplementary Figure S3). Among them, 139 protein spots showed differential abundances in five distinct stages of spore germination (>1.5-fold, p < 0.05). A total of 131 spots were identified using tandem mass spectrometry and Mascot database searching, and eight spots were not matched in database. Among the proteins, 28 spots contained only single peptide match, which were considered as un-identified according to our criteria. In the remaining 103 spots, 80 spots contained a single protein each (Table 1, Supplementary Table S1) and 23 spots contained more than one protein each (Supplementary Table S2). Thus, the 80 proteins were DAPs during spore germination.

The 80 DAPs were classified into eleven groups, including photosynthesis (17), carbohydrate and energy metabolism (9), other metabolisms (5), signaling and vesicle trafficking (8), cell structure (5), cell cycle (2), transcription related (1), protein synthesis (10), protein folding and processing (8), protein degradation (8), stress and defense (7) (Table 1). Among them, proteins involved in photosynthesis (21%) and protein synthesis (13%) were over-represented (Figure 4A), indicating active photosynthesis and de novo protein synthesis are pivotal for the germinating chloropyllous spores.

Interestingly, the 80 DAPs represented 48 unique proteins. Thirteen proteins (16.3%) had multi-proteoforms, which were mainly involved in photosynthesis, tricarboxylic acid (TCA) cycle, vesicle trafficking, protein synthesis, folding and turnover, as well as reactive oxygen species (ROS) scavenging (Table 1, Supplementary Table S3). These proteoforms might be generated from alternative splicing and various post-translational modifications.

The subcellular localization of the 80 DAPs was predicted based on five internet tools (i.e., YLoc, LocTree3, ngLOC, TargetP, and Plant-mPLoc) and literature. In total, 32 DAPs were predicted to be localized in chloroplast, 27 in cytoplasm, one in Golgi apparatus, two in mitochondria, and six in nucleus. Besides, seven proteins were predicted to be localized in two organelles, and five proteins in four organelles (Figure 4B, Supplementary Table S4).

To better understand the protein expression characteristics during the five stages, hierarchical clustering analysis was
applied to the 80 proteins, which revealed three main clusters. Cluster I contained 21 proteins, which were not expressed in MS, but induced in other stages of germination. They were involved in photosynthesis, glycolysis, protein folding and turnover, signal transduction, and ROS scavenging, implying these pathways were specially induced in certain germination stages (Figure 4C). Cluster II included 35 proteins that were mainly decreased during spore germination. These proteins were divided into two subclusters. Subcluster II-1 contained the proteins mainly decreased during germination, and subcluster II-2 included proteins induced at the stage of GS but obviously reduced in SPC. The proteins in cluster II covered ten function categories, in addition to category of transcription (Figure 4D). The rest 24 proteins were grouped into cluster III, representing increased proteins during spore germination. Cluster III contained two subclusters. Subcluster III-1 contained five proteins induced in stages of RS and DCS, but reduced in GS and SPC, and subcluster III-2 included 19 proteins significantly increased at all stages of germination (Figure 4D).
Photosynthesis and Reserve Mobilization are Active in the Germinating Spores

Mature chlorophyllous spores of *Equisetum* species contain chloroplasts and water, making them metabolically active and short-lived. After released from sporangium, the green spores can survive desiccation for less than 2 weeks (Lebkuecher, 1997). It has been found that the *E. hyemale* spores can tolerate desiccation of 2% relative humidity for 24 h. Upon rehydration, they can rapidly regain photosynthetic competence (Lebkuecher, 1997). In this study, we found the viability of
### TABLE 1 | Differentially abundant proteins during *E. arvense* spore germination.

| Spot No. | Protein name | Subcellular location | Plant species | Gi No. (a) | Thr. MW(Da) /pI(l) | Exp. MW(Da) /pI(a) | Cov (%) | Sco (l) | QM(l) | V% ± SD(l) | MS RS DC SPC |
|----------|--------------|----------------------|---------------|------------|---------------------|---------------------|---------|---------|-------|-----------|-------------|
| 121      | Chlorophyll a/b binding protein (CAB) | Chl | Hedera helix | 12,582 | 20,759/ 4.83 | 25,376/ 4.71 | 10 | 57 | 3 |
| 346      | Ribulose-1,5-bisphosphate carboxylase/oxygenase large subunit (RBCL) | Chl | Eryngium bourgatii | 1,292,976 | 53,093/ 5.56 | 28,686/ 6.28 | 6 | 59 | 3 |
| 763      | Ribulose-1,5-bisphosphate carboxylase/oxygenase large subunit (RBCL) | Chl | Donatia fascicularis | 1,304,292 | 49,896/ 5.56 | 134,736/ 5.48 | 8 | 55 | 5 |
| 902      | Ribulose-1,5-bisphosphate carboxylase/oxygenase large subunit (RBCL) | Chl | Grammitis diminuta | 340,031,166 | 46,908/ 6.26 | 50,502/ 6.31 | 5 | 61 | 2 |
| 117      | Ribulose-1,5-bisphosphate carboxylase/oxygenase large subunit (RBCL) | Chl | Equisetum telmateia | 16,565,336 | 48,750/ 6.26 | 52,974/ 6.47 | 15 | 57 | 7 |
| 196      | Ribulose-1,5-bisphosphate carboxylase/oxygenase large subunit (RBCL) | Chl | E. telmateia | 16,565,336 | 48,750/ 6.26 | 51,502/ 6.47 | 16 | 52 | 6 |
| 194      | Ribulose-1,5-bisphosphate carboxylase/oxygenase large subunit (RBCL) | Chl | E. telmateia | 16,565,336 | 48,750/ 6.26 | 51,418/ 6.33 | 13 | 53 | 6 |
| 400      | Ribulose-1,5-bisphosphate carboxylase/oxygenase large subunit (RBCL) | Chl | E. telmateia | 16,565,336 | 48,750/ 6.26 | 49,452/ 6.31 | 11 | 53 | 5 |
| 23       | Ribulose-1,5-bisphosphate carboxylase/oxygenase large subunit (RBCL) | Chl | E. bourgatii | 1,292,976 | 53,093/ 5.56 | 50,872/ 5.02 | 4 | 78 | 2 |
| 528      | Ribulose-1,5-bisphosphate carboxylase/oxygenase large subunit (RBCL) | Chl | Equisetum arvense | 1,352,773 | 52,493/ 5.86 | 67,620/ 4.64 | 7 | 60 | 3 |
| 100      | Ribulose-1,5-bisphosphate carboxylase/oxygenase large subunit (RBCL) | Chl | Isoetes capensis | 83,032,384 | 47,561/ 6.30 | 42,425/ 5.47 | 6 | 55 | 3 |
| 496      | Ribulose-1,5-bisphosphate carboxylase/oxygenase large subunit-binding protein subunit beta (RBP) | Chl | Solanum lycopersicum | 460,379,814 | 64,527/ 5.46 | 71,249/ 5.21 | 4 | 61 | 2 |
| 34       | Ribulose-1,5-bisphosphate carboxylase/oxygenase activase (RCA) | Chl | Gossypium hirsutum | 12,620,883 | 48,609/ 5.06 | 48,999/ 5.19 | 11 | 98 | 4 |
| 184      | Ribulose-1,5-bisphosphate carboxylase/oxygenase activase (RCA) | Chl | Hordeum vulgare | 100,614 | 47,496/ 5.64 | 47,356/ 6.31 | 11 | 94 | 4 |

(Continued)
| Spot No. | Protein name[a] | Subcellular location[c] | Plant species[d] | Gi No. | Thr. MW(Da)/pI[f] | Exp. MW(Da)/pI[g] | Cov (%)[h] | Sco[10] | QM[j] | V% ± SD[k] | MS RS DCS GS SPC |
|---------|----------------|-------------------------|------------------|--------|------------------|------------------|------------|--------|------|------------|-----------------|
| 125     | Ribulose-1,5-bisphosphate carboxylase/oxygenase activase (RCA) | Chl | Musa acuminata subsp. malaccensis | 695,062,479 | 47,898/6.18 | 42,859/5.67 | 8 | 75 | 3 | a b d a | |
| 308     | Transketolase (TK) | Chl | Spinacia oleracea | 2,529,342 | 80,744/6.20 | 95,817/6.20 | 8 | 156 | 5 | b a a a | |
| 84      | Transketolase (TK) | Chl | S. lycopersicum | 460,388,792 | 80,615/6.26 | 85,056/6.11 | 8 | 90 | 7 | a b c b | |

**CARBOHYDRATE AND ENERGY METABOLISM (9)**

| 685     | Unknown, pyruvate dehydrogenase E1 component subunit beta* (PDH) | Chl | Glycine max | 255,647,166 | 44,596/6.28 | 27,410/6.46 | 4 | 60 | 2 | a b c b | |
| 160     | Malate dehydrogenase (MDH) # | Chl, Cyt, Mit, Pox | Arabidopsis thaliana | 15,219,721 | 35,890/6.11 | 46,411/4.98 | 12 | 74 | 3 | a b c a | |
| 503     | Malate dehydrogenase (MDH) # | Chl, Cyt, Mit, Pox | A. thaliana | 11,133,509 | 35,548/6.11 | 29,729/6.41 | 6 | 59 | 3 | a b c a | |
| 260     | Malate dehydrogenase (MDH) # | Chl, Cyt, Mit, Pox | A. thaliana | 731,361,010 | 41,677/8.81 | 32,651/6.16 | 5 | 21 | 2 | b a b a | |
| 164     | Malate dehydrogenase (MDH) # | Chl, Cyt, Mit, Pox | Beta vulgaris subsp. vulgaris | 731,361,010 | 41,677/5.74 | 36,353/5.84 | 12 | 193 | 4 | b a b a | |
| 163     | Malate dehydrogenase (MDH) | Chl | Brachypodium distachyon | 357,147,942 | 41,864/6.97 | 38,560/5.94 | 10 | 60 | 4 | b a c a | |
| 486     | Enolase | Cyt | Tarenaya hassleriana | 729,317,446 | 51,639/5.91 | 48,823/5.62 | 3 | 54 | 2 | b a b b | |
| 489     | 6-phosphogluconate dehydrogenase (6-PGDH) # | Chl, Cyt | G. max | 356,513,305 | 54,116/6.25 | 24,817/5.21 | 12 | 53 | 5 | b a b a | |
| 192     | Fructokinase (FK) | Chl | Lycopersicon esculentum | 23,476,263 | 40,620/5.41 | 37,710/4.91 | 8 | 129 | 3 | b a b c | |

**OTHER METABOLISMS (5)**

| 135     | Enoyl-acyl carrier protein reductase (EAR) | Chl | Oryza sativa subsp. japonica | 75,225,229 | 39,277/8.81 | 32,651/6.16 | 5 | 21 | 2 | b a b a | |
| 197     | Hypothetical protein PHAVU_001G035500g, glycine decarboxylase* (GDC) | Mit | Phaseolus vulgaris | 593,795,946 | 116,167/6.65 | 117,088/6.30 | 11 | 157 | 12 | b a b a | |
| 362     | 3-isopropylmalate dehydrogenase (IPMDH) | Chl | A. thaliana | 15,241,338 | 44,305/5.75 | 28,512/4.87 | 23 | 67 | 7 | b a b c | |

(Continued)
### TABLE 1 | Continued

| Spot No. | Protein name | Subcellular location | Plant species | Gi No. | Thr. MW(Da)/pI | Exp. MW(Da)/pI | Cov (%) | Sco | QM | V% ± SD | MS RS DCS GS SPC |
|----------|--------------|----------------------|---------------|--------|----------------|----------------|--------|-----|-----|---------|------------------|
| 445      | Hypothetical protein SELMDRAFT_406755, containing cd00517 ATP-sulfurylase domain* (ATPS) | Chl | Selaginella moellendorffii | 302,763,978 | 56,179/6.69 | 26,370/5.59 | 5 | 60 | 3 | a b c d | |
| 190      | Predicted protein, pyridoxal biosynthesis protein PDX1* | Cyt | Physcomitrella patens | 168,019,502 | 33,769/6.03 | 29,893/5.82 | 15 | 324 | 8 | a b c d | |
| 1063     | Hypothetical protein SELMDRAFT_151778, containing cd00200 WD40 domain* (WD40) | Nuc | S. moellendorffii | 302,791,020 | 37,267/5.65 | 36,189/5.74 | 6 | 84 | 2 | b c d | |
| 207      | 14-3-3 protein | Cyt | S. oleracea | 440,573,600 | 30,029/4.84 | 27,277/4.79 | 16 | 79 | 6 | a b c d | |
| 276      | Unknown, containing cd00877 Ran GTPase domain* (RAN) | Nuc | P. sitchensis | 116,794,384 | 25,374/6.30 | 27,727/6.66 | 20 | 52 | 4 | a b c d | |
| 317      | GTP-binding nuclear protein Ran/TC4 (RAN) | Nuc | Vicia faba | 586,783 | 25,274/6.39 | 27,750/6.73 | 27 | 187 | 6 | a b c d | |
| 340      | Unknown, containing cd00877 Ran GTPase domain* (RAN) | Nuc | P. sitchensis | 116,794,384 | 25,374/6.30 | 30,115/5.59 | 24 | 53 | 5 | a b c d | |
| 764      | Unknown, containing cd00877 Ran GTPase domain* (RAN) | Nuc | P. sitchensis | 116,794,384 | 25,374/6.30 | 44,900/6.58 | 30 | 51 | 6 | a b c d | |
| 123      | ADP-ribosylation factor (ARF) | Gol | Chlamydomonas reinhardtii | 1,703,374 | 20,747/6.92 | 19,121/6.71 | 29 | 92 | 4 | a b c d | |
| 397      | Os03g0718100, actin* | Cyt | O. sativa subsp. japonica | 115,454,971 | 42,014/5.30 | 32,585/4.95 | 9 | 82 | 3 | a b c d | |
| 554      | Reversibly glycosylated polypeptide (RGP) | Cyt | Ricinus communis | 223,548,230 | 41,557/5.82 | 39,973/5.41 | 14 | 105 | 5 | a b c d | |
| 118      | Reversibly glycosylated polypeptide (RGP) | Cyt | Solanum tuberosum | 34,582,499 | 42,146/5.71 | 35,827/5.36 | 24 | 74 | 8 | a b c d | |
| 299      | Unknown protein, rhamnose biosynthetic enzyme 1* (RBE) | #Cyt | A. thaliana | 8,493,590 | 33,861/5.73 | 29,119/6.48 | 7 | 60 | 3 | a b c d | |
| 482      | Caffeoyl-CoA O-methyltransferase (CCoAOMT) | #Cyt | Eucalyptus gunni | 3,023,419 | 28,010/5.02 | 29,093/4.81 | 11 | 73 | 2 | a b c d | |

(Continued)
### TABLE 1 | Continued

| Spot No. | Protein name | Subcellular location | Plant species | Gi No. | Thr. MW(Da)/pI | Exp. MW(Da)/pI | Cov (%) | Sco | QM | V% ± SD | MS | RS | DCS | GS | SPC |
|----------|--------------|----------------------|---------------|--------|----------------|----------------|---------|-----|----|--------|-----|-----|------|----|-----|
| 104      | Cell division cycle protein 48 homolog (CDC48) | Cyt | *T. hassleriana* | 729,396,339 | 89,888/5.09 | 103,252/5.22 | 21 119 14 |   |   |   | a | b | c | c |   |
| 139      | Proliferation-associated protein 2G4-like (PA2G4) | Nuc | *Nicotiana tomentosiformis* | 697,180,533 | 43,837/5.96 | 55,758/6.73 | 5 52 2 |   |   |   | d | d |   | a | ab c |
| 938      | Predicted protein, containing cd00771 threonyl-tRNA synthetase class II core catalytic domain* (ThrRS) | Cyt | *P. patens* | 168,012,416 | 71,002/5.96 | 80,637/6.37 | 3 54 2 |   |   |   | b | c | c | c |   |
| 83       | Predicted protein, 40S ribosomal protein SA (RPSA) | Cyt | *P. patens* | 168,017,628 | 31,539/4.72 | 43,804/4.58 | 7 101 2 |   |   |   | a | b | c | d |   |
| 454      | Hypothetical protein AMTR_s00087p00135370, containing cd08065 eukaryotic translation initiation factor 3* (eIF3) | Chl, Cyt, Mit, Nuc | *Amborella trichopoda* | 548,842,775 | 44,710/5.17 | 39,587/4.64 | 5 82 2 |   |   |   | b | c | d | a |   |
| 66       | Hypothetical protein, eukaryotic initiation factor 4A* (eIF4A) | Cyt | *P. patens* | 168,026,095 | 47,119/5.46 | 48,999/5.32 | 25 162 11 |   |   |   | a | b | c | c |   |
| 878      | Eukaryotic initiation factor 4A (eIF4A) | Cyt | *Nicotiana tabacum* | 1,170,511 | 47,098/5.37 | 49,134/5.23 | 14 118 6 |   |   |   | a | b | c | c |   |
| 94       | OSJNBA0020P07.3, elongation factor 2* (EF2) | Cyt | *O. sativa* subsp. *japonica* | 38,344,860 | 94,939/5.85 | 115,425/6.42 | 9 111 7 |   |   |   | b | c | c | c |   |
| 538      | Hypothetical protein SELMODRAFT_411087, elongation factor 2* (EF2) | Cyt | *S. moellendorfii* | 302,773,640 | 94,568/6.00 | 114,195/6.33 | 5 167 5 |   |   |   | a | b | c | d |   |
| 510      | Hypothetical protein SELMODRAFT_143627, elongation factor G* (EF-G) | Chl, Mit | *S. moellendorfii* | 302,765,284 | 75,483/5.31 | 119,422/6.41 | 6 75 4 |   |   |   | a | b | c | c |   |
| 942      | Elongation factor Tu (EF-Tu) | Mit | *Setaria italica* | 514,812,465 | 48,530/5.99 | 44,724/6.18 | 16 63 8 |   |   |   | d | d | b | c |   |
| 205      | Uncharacterized protein LOC105056625, containing cd14275 elongation factor Ts domain* (EF-Ts) | Chl | *Elaeis guineensis* | 743,840,139 | 125,269/4.92 | 27,102/5.82 | 2 130 5 |   |   |   | b | c | b | c |   |
| 625      | Tyrosine phosphorylated protein A (TypA) | Chl | *Suaeda salsa* | 162,424,768 | 75,474/6.71 | 85,342/5.71 | 6 205 4 |   |   |   | a | b | b | c |   |

(Continued)
### TABLE 1 | Continued

| Spot No. (a) | Protein name(b) | Subcellular location(c) | Plant species(d) | Gi No. (e) | Thr MW/DA(pI)(f) | Exp MW/DA(pI)(g) | Cov (%) (h) | Sco (i) | QM (j) | V% ± SD (k) | MS RS DCS GS SPC |
|-------------|-----------------|------------------------|-----------------|---------|----------------|----------------|------------|--------|--------|-------------|-----------------|

#### PROTEIN FOLDING AND PROCESSING (8)

542  Hypothetical protein SELMODRAFT_440382, containing pfam00012 heat shock protein 70 domain* (HSP70)
  Cyt  S. moellendorffii  302,770,212  71,931/ 5.17  80,237/ 5.18  13  295  7  

1002  Heat shock protein 70 (HSP70)
  Cyt  Populus trichocarpa  224,098,390  71,620/ 5.14  80,105/ 5.21  16  135  8  

1566  Heat shock protein 70 (HSP70)
  Cyt  Petunia × hybrida  20,559  71,137/ 5.07  106,146/ 6.34  4  92  2  

314  Heat shock protein 70 (HSP70)
  Cyt  Dactylis glomerata  188,011,548  72,002/ 5.03  86,310/ 5.38  21  338  13  

928  Heat shock protein 70 like protein (HSP70)
  Cyt  S. lycopersicum  460,394,037  72,308/ 5.16  91,122/ 5.25  7  122  5  

598  Predicted protein, containing pfam00183 heat shock protein 90 domain* (HSP90)
  Cyt  P. patens  168,034,606  79,652/ 4.93  87,379/ 5.25  5  89  3  

259  T-complex protein 1 subunit alpha (TCP1α)
  Cyt  A. thaliana  135,535  59,477/ 5.93  66,881/ 6.22  8  246  4  

106  Hypothetical protein VITISV_000290, T-complex protein 1 subunit gamma* (TCP1γ)
  Cyt  V. vinifera  147,784,740  61,064/ 6.06  72,989/ 6.14  5  94  3  

#### PROTEIN DEGRADATION (8)

284  Alpha7 proteasomal subunit (PSA7)
  Cyt, Nuc  N. tabacum  14,894,925  27,466/ 6.11  25,449/ 5.20  8  79  2  

199  Zinc dependent protease (ZDP)
  Chl  Trifolium pratense  84,468,286  74,746/ 5.82  73,106/ 5.66  12  106  7  

227  Zinc dependent protease (ZDP)
  Chl  T. pratense  84,468,286  74,746/ 5.82  72,520/ 5.57  5  93  3  

37  Zinc dependent protease (ZDP)
  Chl  T. pratense  84,468,286  74,746/ 5.82  72,520/ 5.57  5  93  3  

869  Zinc metalloprotease (ZMP)
  Chl, Mt  A. thaliana  10,120,424  121,539/ 5.39  131,816/ 5.10  1  44  2  

354  Predicted protein, containing pfam06480 FtsH extracellular domain* (FtsH)
  Chl  Micromonas pusilla CCMP1545  303,275,720  77,421/ 5.29  75,882/ 5.26  6  113  4  

(Continued)
TABLE 1 | Continued

| Spot No. (a) | Protein name (b) | Subcellular location (c) | Plant species (d) | Gi No. (e) | Thr. MW(Da) /pI (f) | Exp. MW(Da) /pI (g) | Cov (%) (h) | Sco (i) | QM (j) | V% ± SD (k) | MS RS DCS GS SPC |
|-------------|-----------------|-------------------------|------------------|------------|---------------------|---------------------|------------|-------|--------|------------|----------------|
| 508         | Predicted protein, ATP-dependent zinc metalloprotease FtsH* (FtsH) | Chl | *P. patens | 188,001,910 | 66,993/5.23 | 108,174/5.50 | 3 | 75 | 2 |
| 59          | ATP-dependent Clp protease ATP-binding subunit ClpC (CLPC) | Chl | *A. thaliana | 9,758,239 | 103,616/6.36 | 94,506/5.45 | 9 | 415 | 9 |
| STRESS AND DEFENSE (7) |
| 414 | Hypothetical protein MIMGU_mgv1a021611mg, containing cd03015 2-cys peroxiredoxin domain* (Prx) | Chl, Cyt | Erythranthe guttata | 604,334,612 | 21,153/4.93 | 21,812/4.68 | 17 | 132 | 4 |
| 92          | 2-cys peroxiredoxin-like protein (Prx) | Chl, Cyt | *Hyacinthus orientalis | 47,027,073 | 21,966/4.93 | 22,930/5.24 | 12 | 58 | 3 |
| 166         | Ascorbate peroxidase (APX) | Cyt | Eucalyptus grandis | 702,241,628 | 27,613/6.07 | 25,671/5.66 | 7 | 104 | 2 |
| 244         | Dehydroascorbate reductase-like protein (DHAR) | Cyt | *S. tuberosum | 76,573,291 | 23,610/6.32 | 24,534/5.60 | 11 | 147 | 3 |
| 361         | Dehydroascorbate reductase-like protein (DHAR) | Cyt | *S. tuberosum | 76,573,291 | 23,610/6.32 | 24,783/5.64 | 11 | 105 | 3 |
| 899         | Dehydroascorbate reductase-like protein (DHAR) | Cyt | *S. tuberosum | 76,160,951 | 23,596/6.09 | 44,783/6.60 | 11 | 92 | 3 |
| 294         | Chloroplast drought-induced stress protein of 32 kDa (CDSP32) | Chl, Cyt | *S. tuberosum | 2,582,822 | 33,779/8.07 | 31,436/6.26 | 8 | 87 | 3 |

*Assigned spot number as indicated in Figure 3.
1 The name and functional category of the proteins identified by ESI-Q-TOF and ESI-Q-Trap tandem mass spectrometry. Protein names marked with an asterisk (*) have been edited by us depending on searching against NCBI non-redundant protein database for functional domain. The abbreviations for the protein names are indicated in the bracket after protein names.
2 Subcellular localization predicted by softwares (YLoc, LocTree3, Plant-mPLoc, ngLOC, and TargetP). Only the consistent predictions from at least two tools were accepted as a confident result. Pounds (#) indicate prediction results were inconsistent among five tools. The subcellular localizations were predicted based on literature listed in Supplementary Table S4. Chl, chloroplast; Cyt, cytoplasm; Gol, Golgi apparatus; Mit, mitochondria; Nuc, nucleus; Pox, peroxisome.
3 The plant species that the peptides matched from.
4 Database accession number from NCBI non-redundant protein database.
5 Theoretical (f) and experimental (g) molecular weight (Da) and pI of identified proteins. Theoretical values were retrieved from the protein database. Experimental values were calculated using ImageMaster 2D version 5.0.
6 The amino acid sequence coverage for the identified proteins.
7 The Mascot score obtained after searching against the NCBI non-redundant protein database.
8 The number of matched peptides for each protein.
9 The mean values of protein spot volumes relative to total volume of all the spots. Five spore germination stages, MS, mature spores; RS, rehydrated spores, DCS, double-celled spores, GS, germinated spores; and SPC, spores with protonemal cells were performed. Error bar indicates ± standard deviation (SD). Letters indicate statistically significant differences (p < 0.05) among five stages of spore germination as determined by One-Way ANOVA.

spores from *E. arvense* lasted less than 3 weeks, and the fresh spores germinated in 32 h (Figure 1). The extremely short viability and rapid germination were mainly due to active photosynthesis and respiration in the chlorophyllous spores. This is different from non-green spores with long dormancy and viability (Lloyd and Klekowski, 1970). Upon germination, chlorophyllous spores also exhibited features distinct from the non-green spores. In our proteomic results, we found that 17 DAPs were photosynthesis-related proteins, including a chlorophyll a/b-binding protein, 10 proteoforms of...
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FIGURE 4 | Functional categorization, subcellular localization, and hierarchical clustering of differentially abundant proteins upon E. arvense spore germination. (A) A total of 80 proteins were classified into 11 functional categories. The percentage of proteins in different functional categories are shown in the pie. (B) Subcellular localization categories of proteins predicted based on softwares and literature. The numbers of proteins with different locations are shown in the pie. Chl, chloroplast; Cyt, cytoplasm; Gol, Golgi apparatus; Mit, mitochondria; Nuc, nucleus; Pox, peroxisome. (C,D) Dendrogram of 80 differentially abundant proteins obtained by hierarchical clustering analysis. The columns represent different spore germination stages, including MS, mature spores; RS, rehydrated spores; DCS, double-celled spores; GS, germinated spores; SPC, spores with protonemal cells. The rows represent individual proteins. Three main clusters (I–III) and subclusters of II and III (II-1, II-2, III-1, and III-2) are shown on the left side. Functional categories indicated by capital letters, spot numbers, and protein name abbreviations are listed on the right side. The scale bar indicates log (base 2) transformed protein abundance ratios ranging from -13.0 to 3.0. The ratio was calculated as protein abundance at MS stage divided by abundance each stage, respectively. The increased and decreased proteins are represented in red and green, respectively. The color intensity increases with increasing abundant differences. Undetected proteins are indicated in gray. Abbreviations for functional categories: A, Photosynthesis; B, Carbohydrate and energy metabolism; C, Other metabolisms; D, Signaling and vesicle trafficking; E, Cell structure; F, Cell cycle; G, Transcription related; H, Protein synthesis; I, Protein folding and processing; J, Protein degradation; K, Stress and defense. Detailed information on protein names and abbreviations can be found in Table 1.

Ribulose-1,5-bisphosphate carboxylase/oxygenase (RuBisCO) large subunit, a RuBisCO large subunit binding protein, three proteoforms of RuBisCO activase, and two proteoforms of chloroplast transketolase (Table 1). The increased chlorophyll a/b-binding protein indicated that the photosystem II was enhanced upon spore germination. The multiple proteoforms of RuBisCO large subunit, RuBisCO activase, and transketolase changed in abundance possibly due to protein phosphorylation (Guitton and Mache, 1987). This implies that the carbon assimilation is active in germinating spores. Importantly, we also found nine enzymes involved in carbohydrate and energy metabolism. They were six TCA cycle enzymes (i.e., a pyruvate dehydrogenase and five malate dehydrogenases), an enolase involved in glycolysis, a cytosolic 6-phosphogluconate dehydrogenase (an enzyme in pentose phosphate pathway), and a fructokinase in charge of sucrose and fructose metabolism. All these enzymes, taking up 32% of DAPs, were considered as key enzymes for carbon and energy supplies during spore germination. Their variations also indicate that the heterotrophic metabolism is crucial for chlorophyllous spore germination.

We also found several proteins involved in fatty acid synthesis, amino acid metabolism, sulfur assimilation, and secondary metabolism (Table 1). Among them, enoyl-acyl carrier protein reductase is a key enzyme of the type II fatty acid synthesis system
in plasts. Enoyl-acyl carrier protein reductase was proved to be important for fatty acid deposition in developing seeds (de Boer et al., 1999) and pollen grains (Poghosyan et al., 2005). In our results, enoyl-acyl carrier protein reductase was decreased at the stages of DCS, GS and SPC during germination. This indicates that fatty acid synthesis is reduced upon spore germination, which is consistent with the notion that mobilization of storage lipids has been triggered for energy supply in germinating spores (DeMaggio and Stetler, 1985). Besides, we found three increased enzymes at certain stages involved in amino acid metabolism, including glycine decarboxylase, 3-isopropylmalate dehydrogenase (IPMDH), and ATP-sulfurylase. Glycine decarboxylase participates in glycine, serine and threonine metabolism, and was found to be abundant in mitochondria of C3 leaves that functions in photosynthetic carbon recovery (Timm et al., 2012). Whether glycine decarboxylase is involved in photosynthetic in germinating fern spores needs to be further investigated. IPMDH catalyzes the oxidative decarboxylation of 3-isopropylmalate in leucine biosynthesis, and was proved to be essential for Arabidopsis pollen development (He et al., 2011). In our results, IPMDH was obviously induced in the stage of GS, suggesting that IPMDH is also pivotal for spore germination. Interestingly, ATP-sulfurylase acts as the metabolic entry point into the sulfur assimilation pathway. It was found that the increase of ATP-sulfurylase level during soybean seed development could lead to an increase in the availability of sulfur amino acids (Phartiyal et al., 2006). The induced ATP-sulfurylase in GS and SPC may facilitate the synthesis of sulfur rich amino acids for spore germination. Additionally, pyridoxal biosynthesis protein (PDX) was specially expressed in germinating spores, but not found in MS. PDX family in plants is primarily known for its role in vitamin B6 biosynthesis. Recently, PDX1.2 was proved to be critically required for hypocotyl elongation and primary root growth (Leuendorf et al., 2014). The increase of PDX in germinating spores implies that it probably functions in rhizoid elongation.

Signaling and Vesicular Trafficking are Important for Spore Germination

The critical roles of hormone signaling and vesicular trafficking in germinating seeds and pollen grains have been well-studied (Ellis and Turner, 2002; Dai et al., 2007a), but little information is available for fern spore germination. In this study, we found some novel factors in signal transduction and vesicular trafficking upon fern spore germination, including chalcone isomerase, WD40, 14-3-3 protein, GTP-binding nuclear protein Ran (RAN), and ADP-ribosylation factor (ARF) (Table 1). Among them, chalcone isomerase involved in flavonoid biosynthesis was increased at the stage of RS. Flavonoid participates in auxin signaling and facilitates pollen-tube growth (Falcone Ferreyra et al., 2012). In maize (Zea mays) and Petunia hybrida mutants, the flavonoid-deficient pollen failed to produce a functional pollen tube (Mo et al., 1992). The increase of chalcone isomerase in spores implies that flavonoids may function in fern spore germination as well. Interestingly, flavonoid biosynthesis is regulated by conserved WD40 domain–contained transcription factors (Falcone Ferreyra et al., 2012). Members of WD40 protein superfamily are known as key regulators of multi-cellular processes (e.g., cell division, light signaling, protein trafficking, cytoskeleton dynamics, nuclear export, RNA processing, chromatin modification, and transcriptional mechanism), acting as scaffolding molecules assisting proper activity of other proteins (Stürnemann et al., 2010). Moreover, some WD40 proteins have been found to be crucial for Arabidopsis seed germination (Gachomo et al., 2014) and pollen viability in flax (Linum usitatissimum L.) (Kumar et al., 2013). In our results, the nucleus-localized WD40 was induced in horsetail germinating spores, indicating its regulatory function for spore germination and rhizoid tip-growth. Similarly, 14-3-3 protein family is also a highly conserved eukaryotic proteins with multiple molecular and cellular functions by binding to phosphorylated client proteins to modulate their function (Denison et al., 2011). In lily (Lilium longiflorum) pollen, 14-3-3 protein was involved in the regulation of plasma membrane H+-ATPase via modulation of its activity, which is essential for germination and tube elongation (Pertl et al., 2010). In this study, we found a 14-3-3 protein was induced in the stages of RS and SPC, indicating it would play roles in cell nuclear migration and rhizoid tip-growth. Importantly, we also found four proteoforms of RAN were significantly increased in germinating spores. RAN, a primarily nucleus-localized small GTPase, is essential for nuclear transport and assembly, mRNA processing, and cell cycle (Ciciarello et al., 2007). It was reported that overexpression of RANI in rice (Oryza sativa) and Arabidopsis altered the mitotic progress and sensitivity to auxin (Wang et al., 2006). The multi-proteoforms of RAN induced in germinating horsetail spores were probably due to the different phosphorylation levels, implying their probable key roles in cell division and polar growth during spore germination. Besides of RAN, we found another important small GTPase, ARF, was induced at the stages of RS and DCS. ARF contributes to the regulation of multiple trafficking routes with respect to Golgi organization, endocytic cycling, cell polarity and cytokinesis (Yorimitsu et al., 2014). It was found that ARF played essential roles for endosomal recycling during Arabidopsis pollen and root hair polarized tip growth (Richter et al., 2011). The induced ARF in horsetail spores meets the specific requirement of early-secretory and polar vesical recycling to facilitate rhizoids tip growth.

Cytoskeleton and Cell Wall Dynamics are Necessary for Spore Germination

Actin cytoskeleton directs the flow of vesicles to the apical domain, where they fuse with the plasma membrane and contribute their contents to the expanding cell wall (Hepler et al., 2013). The local changes in contents and viscosity of the apical wall control the local expansion rate and cell elongation. Precise mechanisms in the organization of actin cytoskeleton and cell wall dynamics have been well-studied in growing pollen tubes (Hepler et al., 2013; Qu et al., 2015), but the mechanisms remain to be further elucidated in fern spores. Here we found the levels of actin, reversibly glycosylated polypeptide, rhamnose biosynthetic enzyme 1, and caffeoyl-CoA O-methyltransferase (CCoAOMT) were altered during horsetail spore germination. Actin is known to be the main content of actin cytoskeleton, and the other three enzymes are essential
for cell wall modulation. Reversibly glycosylated polypeptide is implicated in polysaccharide biosynthesis (Langeveld et al., 2002), and may function in cell wall construction in pollen from rice and *Picea meyeri* (Dai et al., 2006; Chen et al., 2009). Besides, rhamnose biosynthetic enzyme catalyzes the synthesis of L-rhamnose, an important constituent of pectic polysaccharides in cell wall of pollen tube (Yue et al., 2014). In addition, CCoAOMT has a proven role in lignin monomer biosynthesis, which is crucial for pollen wall development (Arnaud et al., 2012). In our results, rhamnose biosynthetic enzyme was induced in GS and decreased in SPC, CCoAOMT was induced in RS and reduced in the stages of DCS and SPC, and two proteoforms reversibly glycosylated polypeptide displayed different changes. All these alterations would modulate the biosynthesis of cell wall components (e.g., polysaccharide, rhamnose, and lignin), leading to the dynamics of rhizoid cell wall rigidity for sustained polarized growth.

**Rapid Protein Synthesis, Processing, and Turnover are Essential for Fern Spore Germination**

The germinating pollen and fern spores exhibit quick switches from metabolic quiescent state to active state. The substance and energy supply for rapid cell division and polar tip-growth need to be triggered in a short time period. Although it has been found that mature pollen grains have pre-synthesized mRNA and proteins for germination and tube growth (Taylor and Hepler, 1997; Dai et al., 2006), *de novo* protein synthesis is necessary for pollen tube elongation (Dai et al., 2007a,b). For fern species, the germination of green spores from *O. sensibilis* and non-green spores from *A. phyllitidis, Marsilea vestita, Pteridium aquilinum,* and *Pteris vittata* were not inhibited by actinomycin D (Raghavan, 1977, 1991, 1992; Kuligowski et al., 1991). In horsetail spores, similar metabolic features were discovered from our proteomics data. We found eleven DAPs involved in transcription and protein synthesis, including a threonyl-tRNA synthetase, a 40S ribosomal protein SA, three eukaryotic translation initiation factors (eIFs) (a eIF3 and two eIF4A), five elongation factors (EFs) (two proteoforms of EF2, a EF-G, a EF-Tu, and a EF-Ts), and a protein translation-related GTP-binding protein tyrosine phosphorylated protein A (TypA) (*Table 1*). The increases of threonyl-tRNA synthetase in GS and SPC and increases of 40S ribosomal protein SA in RS and DCS indicate that the protein synthesis machinery is enhanced during spore germination, while the changes in multi-proteoforms of eIF and EF imply that certain specific protein synthesis is regulated in diverse modes in different stages of germinating spores. It is interesting to note that these proteins are localized in cytoplasm, mitochondria, and chloroplast, respectively (*Table 1*). This indicates that not only the synthesis of nuclear gene-encoding proteins are necessary, but also the protein synthesis machineries in chloroplast and mitochondria are all triggered for the active metabolism in germinating spores.

Molecular chaperones not only control house-keeping processes, but also regulate protein functional folding and assembly which are necessary for activating/inhibiting various signaling pathways. In horsetail germinating spores, we found eight chaperones, including five proteoforms of heat shock protein 70 (HSP70), HSP90, T-complex protein 1 (TCP1) subunit alpha, and TCP1 subunit gamma. Among these HSP members, HSP70 was found to bind microtubules and interact with kinesin in tobacco (*Nicotiana tabacum*) pollen tubes (Parrott et al., 2013). Besides, Arabidopsis HSP90 was found to be active in mature and germinating pollen grains (Prasinos et al., 2005). Moreover, TCP1 plays a pivotal role in the folding and assembly of cytoskeleton proteins as an individual or complex with other subunits (Sternlicht et al., 1993; Bhaskar et al., 2012). The variations of chaperones in spores highlight that protein processing is important for diverse processes (e.g., photosynthesis, cytoskeleton, protein turnover, and various metabolisms) upon horsetail spore germination.

In addition to folding and assembly, active protein turnover occurred in germinating horsetail spores, which was reflected by the changes of eight degradation-related proteins (*Table 1*). These proteins included an alpha 7 proteasome subunit, three proteoforms of zinc dependent protease, a zinc metalloprotease, two proteoforms of FtsH protease, and an ATP-dependent Clp protease (*Table 1*). Among them, all the zinc dependent proteases, zinc metalloprotease, and FtsH are all ATP-dependent metalloproteases in chloroplasts, which play a major role in assembly and maintenance of the plastidic membrane system. The Clp protease system plays essential role in plastid development through selective removal of miss-folded, aggregated (Nishimura and van Wijk, 2014), or unwanted proteins. All these imply that active protein degradation and turnover in chloropyllous spores are crucial for spore germination.

**ROS Homeostasis is Crucial for Spore Germination**

In the tip-growing pollen tubes and root hairs, ROS act as regulators in diverse signal and metabolic pathways, modulating kinase cascades, ion channels, cell wall properties, nitric oxide levels, and G-protein activities (Wilson et al., 2008; Swanson and Gilroy, 2010). Interestingly, in fern spores, nitric oxide was shown to be a positive regulator for tip growth (Bushart and Roux, 2007). However, the ROS function in fern spores is poorly understood. Our proteomics results revealed that seven proteins in germinating spores function as ROS scavengers, including two proteoforms of 2-cys peroxiredoxin (Prx), a chloroplast drought-induced stress protein of 32 kDa (CDSP32), ascorbate peroxidase (APX), and three proteoforms of dehydroascorbate reductase (DHAR). The proteoforms of Prx, CDSP32, DHAR, and APX are predicted to be localized in chloroplasts and/or cytoplasm in horsetail spores (*Table 1*). CDSP32 is composed of two thioredoxin (Trx) modules and has been found to be involved in the protection of the photosynthetic apparatus against oxidative damage (Brien et al., 2002). PrxR/Trx pathway is a central antioxidant defense system in plants, in which PrxRs employ a thiol-based catalytic mechanism to reduce H$_2$O$_2$ and is regenerated using Trxs as electron donors. APX and DHAR are the key members in glutathione-ascorbate cycle. In this cycle, H$_2$O$_2$ is reduced to water by APX using ascorbate as the electron donor. The oxidized ascorbate is still a radical,
which can be converted into dehydroascorbate spontaneously or reduced to ascorbate by monodehydroascorbate reductase. Dehydroascorbate is then reduced to ascorbate by DHAR at the expense of glutathione, yielding oxidized glutathione. Finally, oxidized glutathione is reduced by glutathione reductase using NADPH as electron donor (Horling et al., 2003). Previous proteomics studies have revealed that the abundances of APX and Trx were changed in germinating pollen grains from O. sativa (Dai et al., 2007a), Arabidopsis (Ge et al., 2011), Brassica napus (Sheoran et al., 2009), P. meyeri (Chen et al., 2009), Picea wilsonii (Chen et al., 2012), and Pinus strobus (Fernando, 2005). This indicated that PrxR/Trx pathway and glutathione-ascorbate cycle were employed in germinating pollen grains. Thus, our results indicate chlorophyllous fern spores with active photosynthesis and respiration modulate ROS homeostasis by these enzymes during germination.

Prediction of Protein-protein Interaction Upon Spore Germination

The fern spore germination is a fine-tuned process regulated by temporal and spatial expression and interaction of a number of genes/proteins. To discover the relationship of DAPs during horsetail spore germination, the protein-protein interaction networks were generated by the web-tool STRING 9.1 (http://string-db.org). The DAPs homologs in Arabidopsis were found by sequence BLASTing in TAIR database (http://www.arabidopsis.org/Blast/index.jsp) (Supplementary Table S5), and then the homologs were subjected to the molecular interaction tool of STRING 9.1 for creation of proteome-scale interaction network. Among all the 80 DAPs, 63 proteins identified in germinating spores and represented by 38 unique homologous proteins from Arabidopsis, were depicted in the STRING database (Figure 5), according to the information from the published literature, genome analysis based on domain fusion, phylogenetic profiling/homology, gene neighborhood, co-occurrence, co-expression, and other experimental evidence (Supplementary Figure S4). Four functional modules are apparently illuminated in the network, which form tightly connected clusters (Figure 5). In the protein networks, stronger associations are represented by thicker lines (Figure 5). In Module 1 (green nodes), photosynthetic proteins (chlorophyll a/b-binding protein, RuBisCO large subunit, RuBisCO activase, and RuBisCO large subunit binding protein), three members of chloroplastic protein synthesis machine (EF-Ts, EF-G, and TypA), chloroplast-located Prx/Trx and zinc dependent protease, as well as a mitochondria-located glycine decarboxylase appeared linked closely. This implies that photosynthesis, photorespiration, chloroplastic protein synthesis and turnover, as well as ROS scavenging are active and cooperated closely in horsetail chlorophyllous spores. Besides, multiple metabolic enzymes (e.g., two malate dehydrogenases, enolase, transketolase, pyruvate dehydrogenase, and 6-phosphogluconate dehydrogenase), cytoplasmic members of protein synthesis machine (40S ribosomal protein SA,

![FIGURE 5](image_url)

**FIGURE 5** | The protein-protein interaction (PPI) network of proteins in *E. arvense* spores based on STRING analysis. A total of 63 differentially abundant proteins represented by 38 unique homologous proteins from Arabidopsis are shown in PPI network. Nodes in different colors belong to four main groups. The PPI network is shown in the confidence view generated by STRING database. Strong associations are represented by thicker lines. Detailed information on protein names and abbreviations can be found in Table 1.
Conclusion and Remarks

Although the molecular mechanism of pollen germination has been well-studied as a model for cell polar growth, our knowledge of fern chlorophyllous spore germination is lacking. In this proteomics study, we found some pollen homologous proteins and several novel components are pivotal for fern spore germination. The dynamics of photosynthesis, TCA cycle, glycolysis, and pentose phosphate pathway, as well as the variations of reserve mobilization pathways (fatty acid synthesis, amino acid metabolism, sulfur assimilation, and secondary metabolism) indicate that both heterotrophic and autotrophic metabolisms are triggered in chlorophyllous spores, which is obviously distinct with non-green spores and pollen grains. Besides, a number of proteins are suspected to be necessary for the cell nuclear migration, cytoskeleton dynamics, and cell wall modulation during fern spore germination. Importantly, the protein synthesis machines, protein processing, and proteasome-dependent protein degradation in cytoplasm and chloroplasts are active for the rapid protein synthesis and turnover. In addition, several members in ROS signaling and G protein-involved vesicle trafficking are crucial for polar rhizoid growth. All these provide invaluable information, however, further validation and characterization of these proteins in a model system (i.e., C. richardii), as well as the post-translational modification analysis are still necessary for ultimately discovering protein functions and interactions toward understanding of the underlying sophisticated cellular and molecular processes in fern germinating spores.

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Supplementary Material

The Supplementary Material for this article can be found online at: http://journal.frontiersin.org/article/10.3389/fpls.2015.00441/abstract

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