Reserve metabolism and ubiquitin proteasome system are involved in germination progress of wheat grains

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Abstract

To further elucidate the molecular mechanism of the germination in common wheat, a differential proteomic analysis was completed between the F1 hybrid RS5322 and its parental lines (RS53 and SN22) using isobaric tags for relative and absolute quantitation (iTRAQ) based strategy. Rapid grain germination and superior young seedling growth were observed in the F1 hybrid line. A total of 807 differentially abundant proteins (DAPs) were identified by iTRAQ analysis of grains at 48 h after imbibition in distilled water. Bioinformatics analysis shows that 638 DAPs were annotated in 38 Gene Ontology functional groups, 764 DAPs were classified into 23 clusters of orthologous groups of protein categories, and 538 DAPs were enriched in 65 Kyoto encyclopedia of genes and genomes pathways. Real time quantitative PCR of 12 genes encoding different important proteins showed certain transcriptional and translational expression similarities during grain development. In the F1 hybrid, the DAPs were particularly those involved in starch and sugar metabolism, protein metabolism, protein modification, and ubiquitin proteasome system (UPS). It was speculated that UPS might be responsible for a high germination ability in the F1 hybrid by regulating storage substance metabolism. The DAPs identified in this study provide a scope for improving the grain germination trait in agricultural crops.

Additional key words: differentially abundant proteins, F1 hybrid, iTRAQ, seed imbibition.

Introduction

Heterosis is a phenomenon whereby F1 offspring can display improved or increased function of some biological qualities that exceed those observed in their parents. The heterotic traits of an offspring are enhanced as a result of mixing the genetic contributions of its parents (Chen et al. 2015).

In plants, this phenomenon has been exploited for improvement of phenotypic and agronomic traits such as germination, growth, resistance to biotic/abiotic stresses, and yield (Fu et al. 2015). The hybrid utilization in maize, oil seed crops, and rice has greatly promoted production in the world (Premlatha et al. 2010, Mistry et al. 2016, Wang et al. 2017). The seed production cost of hybrid cultivar is generally higher than that of inbred cultivar. The seed germination quality directly affects the promotion and utilization of hybrid (Colombo et al. 2017, Derer et al. 2018). Therefore, it is very important to study the characteristic of seed quality and germination mechanism in hybrid cultivars.

Maize grains were used to study protein profile differences of seed germination between hybrid and its parents by proteomics based on two-dimensional electrophoresis. The results of Fu et al. (2011) indicated that dominance, partial dominance, and overdominance of the different proteins played roles in regulating seed germination in maize. The most significant are those proteins involved in germination-related hormone signal transduction, especially the abscisic acid (ABA) and gibberellin (GA) regulation networks. The results of Meena et al. (2018) indicated that the differentially...
abundant proteins, particularly involved in metabolic and energy processes, as well as hormone biosynthesis in the F₁ hybrid, might be responsible for heterotic seed germination in this hybrid.

The germination is considered an important critical stage in the crop life cycle. Seed germination involves a series of events, which begins with water imbibition by dry seeds followed by the extrusion of the radicle towards the end. The uniform performance of crop germination and the subsequent healthy seedlings are vital factors for crop production and are of both economical and ecological importance (Lamichhane et al. 2018). The plant seed is the reservoir of saccharides, oils, and proteins, which maintain its viability during dormancy and provides primary substances for the growth of the embryo during germination (Han et al. 2014). The degradation of seed storage is a complex physiological process, and many physiological factors or pathways are involved in germination mechanisms.

So far, there are many studies on the germination characteristics of plant seeds, including global protein expression trends (Jacobsen et al. 2013, Ziková et al. 2013), specific protein function, storage materials degradation, responses to stress, and post-transcriptional modification (Novikova et al. 2014, Wei et al. 2015, Eckstein et al. 2016, Singarayer et al. 2018), and some research focused on specific organs and developmental stages at the molecular level (Long et al. 2012).

Wheat (Triticum aestivum L.) is the second most widely produced crop in the world, and its production is considered vital for global food security (Taki et al. 2018). Several wheat hybrid cultivars have been developed and grown commercially (Sun et al. 2017, Tikhenko et al. 2017); however, seldom attempts have been made to investigate germination characteristics at the translational level in wheat hybrid seeds. In this study, differentially abundant proteins (DAPs) during grain germination were identified using isobaric tags for relative and absolute quantitation (iTRAQ) approach in a high-yielding grown F₁ wheat hybrid (RS5322) by comparing it with its parental lines (RS53 and SN22). We investigated the divergence of DAPs involved in physiological mechanisms in seed germination. Furthermore, the transcriptions of genes corresponding to some DAPs were also analyzed using real time quantitative PCR. This study added more information on the DAPs associated with higher seed germination ability in wheat F₁ hybrids and broadened the knowledge of molecular processes in F₁ grain germination progress.

Materials and methods

Plants and treatments: Grains of wheat (Triticum aestivum L.) F₁ hybrid (RS 5322) were generated by hand pollination from its female parent (RS53) and male parent Shannong 22 (SN22) and the grains were harvested at physiological maturity in June 2017 on the experimental farm of Shandong Agricultural University, Taian, China (E 117° 09′, N 36° 09′). The RS53 was friendly provided by Heterotic Crop Utilization Laboratory, Institute of Crops, Chinese Academy of Agricultural Sciences, Beijing, China. The SN22 was developed and preserved in our laboratory. The mass of 1 000 grains and the germination characteristics of the F₁ hybrid and its parents were represented an average of three replicates. The grains were imbibed in distilled water in the dark at 20 ± 2 °C for 12 h and then they were placed with the embryo side down on moist three-layered filter paper in a Petri dish for germination under a 12-h photoperiod, an irradiance of 250 µmol m⁻² s⁻¹, day/night temperatures of 25/20 °C, and a relative humidity of 75 %. Grains were examined at regular intervals for determining the onset of germination and were considered as germinated based on the emergence of the visible radicle. The germination rate was recorded from 12 to 96 h after imbibition (HAI). The germination experiments were repeated three times with each experiment consisting of five replicates and 200 grains were used in each replicate. For proteomics analysis, grains at 48 HAI were used for analyzing the differences in the protein pattern of the F₁ hybrid in comparison with those of its parental lines.

Sample preparation and iTRAQ labeling: Protein preparation was conducted according to Ma et al. (2014) with minor modifications. Grain samples were ground into fine powder in liquid nitrogen using a mortar and pestle, and 100 mg samples were extracted with 1 cm³ of extraction buffer [50 mM Tris-HCl (pH 8.0), 0.1 M KCl, 5 mM EDTA, 30 % (m/v) sucrose] containing 1 mM phenylmercuricsulphonium fluoride for 3 h. After centrifugation at 15 000 g for 20 min, supernatants were transferred to new tubes. Five-fold volumes of 10 % (m/v) cold trichloroacetic acid in acetone were added to the supernatants and stored at -20 °C for 2 h, followed by centrifugation at 15 000 g for 20 min. The pellets were rinsed with 90 % (v/v) cold acetone, left at -20 °C for 40 min, and then centrifuged at 15 000 g for 5 min. This rinse step was repeated three times. After freeze drying, the final pellets were stored at -80 °C or analyzed instantly.

Protein samples were incorporated into 500 mm² of STD buffer [4 % (m/v) sodium dodecylsulfate, 150 mM Tris-HCl, 1 mM diithiothreitol, pH 8.0], and incubated in boiling water for 8 min, then subjected to ultra-sonication for 10 times (duration: 4 min; time interval: 4 min). After centrifugation at 15 000 g for 60 min, protein concentrations were determined by Bradford method. About 150 µg protein samples were diluted with 150 mm² of UA buffer (8 mM urea, 150 mM Tris-HCl, pH 8.0), centrifuged at 15 000 g for 30 min, and then 150 mm² of UA buffer was added and centrifuged for another 30 min. After adding 100 mm² of UA buffer with 50 mM iodoacetamide, the samples were incubated in darkness for 30 min, and then centrifuged at 15 000 g for 20 min; this was repeated twice. Then, 100 mm² of DS buffer (50 mM triethylammonium bicarbonate at pH 8.5) were added and centrifuged at 15 000 g for 20 min. This step was repeated twice and then 40 mm² trypsin solution (2 µg trypsin from Promega in 0.04 cm³ of DS buffer) was added. The samples were incubated at 37°C for about 16 - 18 h. The resulting peptides were collected by centrifugation and the peptide content
was tested by Bradford method. The iTRAQ labeling of 100 μg peptide samples from hybrid and its parental lines grains was performed using three iTRAQ Reagent 4-plex kits according to the manufacturer’s protocol (AB Sciex, Foster City, CA, USA). Three biological replicates were iTRAQ-labeled. Protein samples of each line were equally mixed as an inner standard and labeled with 114, 115, 116, and 117 tags. Three 4plex aApplications kits were used to label the samples in this study (Fig. 1 Suppl.).

Liquid chromatography mass spectrometry / mass spectrometry and data analysis: The iTRAQ labeled and pooled peptide samples were analyzed on a tandem mass spectrometer Q Exactive equipped with an Easy nLC autosampler (Thermo Fisher Scientific, San Jose, CA, USA). Survey scans were acquired at resolution of 70 000 at m/z 200, and resolution for HCD spectra was set to 17 500 at m/z 200; the mass window was 2 Da for precursor ion selection; normalized collision energy was 30 eV; and dynamic exclusion duration was 60 s. Five μg of the peptide mixture was loaded on a packed capillary tip (C18-reversed phase column with 15 cm long, 75 μm inner diameter) with RP-C18 5 μm resin and washed in buffer A (0.1 %, m/v, formic acid in water). After washing, the peptide mixture was separated with a gradient of buffer B (0.1 % v/v) formic acid in acetonitrile at a flow rate of 0.25 mm³ min⁻¹. A data-dependent “top10” method was used to get the most abundant precursor ions (mass range 300 - 1 800 m/z). For protein identification, the MS raw files were processed by Mascot 2.4 (Matrix Science, 2012) and Proteome Discoverer 2.2 (Thermo Fisher Scientific, Shanghai, China). The acquired MS/MS spectra were automatically searched against the uniprot_pooideae_108267.fasta, and the total number of protein sequences used in this database was 108 267. A unique protein with at least two unique peptides, with a false discovery rate < 0.01, was qualified for further quantification data analysis. The parameters were set as: a peptide mass tolerance of ± 20 ppm, fragment mass tolerance of 0.1 Da, and number of allowed missed tryptic cleavage sites of 2. Protein quantification was based on the total intensity of the assigned peptides. The average of three labelled samples mixes was used as reference, based on the weighted average of the intensity of report ions in each identified peptide. When the abundance of a protein showed a difference corresponding to a > 1.6-fold or < 0.625-fold change in F1 compared with its parental line, and has a statistical significance (P ≤ 0.05), the protein was considered to be a differential abundance protein.

The functional annotations of the DAPs identified between RS5322 and its parents were performed using the Blast2GO program against the non-redundant protein database (NR; NCBI). Functional classification of proteins by clusters of orthologous groups (COG) database (http://www.ncbi.nlm.nih.gov/COG/) was performed on all identified DAPs using the Blastx 2.2.26+ software in the STRING 9.8 database. Then, all identified DAPs were mapped to a pathway in the KEGG database using Blast2GO/Blastp 2.2.26+. A P-value ≤ 0.05 was used as the threshold to judge the significance of GO, COG, and KEGG pathway analysis results.

Extraction of RNA and real-time quantitative PCR analysis: Total RNA of wheat grains were extracted using the RN8-EASY spin Plus Plant RNA kit (Aidlab, Beijing, China) according to the manufacturer’s instructions. Genomic DNA was removed by digesting each sample (20 - 50 μg of total RNA) with DNase I (Promega, Madison, WI, USA). According to the manufacturer’s instructions, approximately 1 μg RNA was reverse transcribed to cDNA using SuperScript III (Invitrogen, USA). Primer pairs for qPCR analysis (Table 3 Suppl.) were designed by the Primer3Plus program (http://www.bioinformatics.nl/cgi-bin/primer3plus/primer3plus.cgi) and checked by blasting primer sequences in the NCBI database (http://www.ncbi.nlm.nih.gov/tools/blastHome), and all primers were specifically consistent with the respective sequence of its targeted gene. All qPCRs were carried out using SYBR Green PCR Master Mix (Roche Applied Science, Penzberg, Germany) on an ABI 7500 real-time PCR system (Applied Biosystems, Foster City, CA, USA) with three replicates. Data were processed using the 2-△△Ct method. ADP-ribosylation factor gene was used as reference for normalization.

Statistical analysis: Statistical analyses were performed with the SPSS version 17.0 software package for Windows (IBM SPSS Statistics). Data are presented as mean values ± SD of independent experiments. Differences between treatments in this research were analyzed by one-way analysis of variance (ANOVA), and comparisons among means were made using the Duncan’s test calculated at P = 0.05.

Results

The germination pattern was examined in a wheat F1 hybrid (RS5322) in comparison with its parental lines (Fig. 1). Germination percentage of 96.2 % was achieved in the F1 hybrid at 96 HAI, which was slightly higher than those of its parents, which were 93.7 and 89.6 % in the female and male parent, respectively. Germination was initiated in F1 hybrid grains at 12 HAI with 18.5 % compared with 9.2 - 7.8 % germination rates observed in parental lines at the same time. In addition, 7-d-old seedling dry mass per plant of the F1 hybrid (53.6 ± 3.7 mg) was significantly higher than those of the female parent (46.2 ± 5.6 mg) and the male parent (47.6 ± 2.8 mg). The 1000-grain dry mass of the experimental lines were 43.3 ± 1.9, 45.3 ± 3.1, and 42.8 ± 2.7 g in F1 hybrid, female parent, male parent, respectively. This finding indicated that the F1 hybrid with the rapid grain germination and superior seedling growth did not have the highest grain mass before germination.

In this study, iTRAQ-based quantitative proteome characterization during grain germination of the wheat hybrid RS5322 and its parental lines (RS53 and SN22) was investigated to reveal the main DAPs involved in wheat grain germination. A global profiling of quantitative proteome was obtained at 48 HAI, at which the hybrid showed significantly different germination capability compared with its parental lines. The data collected from
these samples were analyzed using Mascot and Proteome Discovery software. Overall, 43,672 spectra identified were matched to known spectra, and 12,310 unique peptides and the corresponding 5,651 proteins were identified. According to false discovery rate $< 0.01$ and unique peptide numbers $\geq 2$, a total of 3,234 non-redundant proteins were obtained.

A 1.6-fold cut-off was used to implicate significant changes in the DAPs during grain germination between RS5322 and its parental lines. Of 3,234 non-redundant proteins identified, 807 showed more than 1.6-fold changes ($P \leq 0.05$) in protein abundance between hybrid and at least one of its parental lines, therefore, these proteins were identified as DAPs. Part of DAPs in hybrid compared with its parental lines is provided in Table 4 Suppl. More DAPs and their detailed information could be found in the research laboratory’s homepage (http://www.538hybridwheat.cn/Blog/html/blog.github.html) and iProX (https://www.iprox.org/) with the dataset identifier IPX0001950000. The distributions of 807 DAPs and their overlapping between the hybrid and its parental lines are illustrated using the Venn diagram (Fig. 2). In general, six clearly different expression patterns during grain germination were generalized among 807 DAPs. A total of 383 DAPs were down-regulated in the hybrid compared with at least one parental line (Fig. 2A), including 92 DAPs decreased compared with both parental lines (I), while 109 and 182 DAPs, particularly down-regulated compared with the female (II) and the male line (III), respectively. As indicated in Fig. 2B, among 424 up-regulated DAPs, 98 DAPs were shared by both parental lines (IV), 130 and 196 DAPs particularly up-regulated compared with the female (V) and male line (VI), respectively.

According to the molecular functions listed on the GO website, the 807 DAPs were classified into 14 functional categories (Fig. 3A). Among these functional protein categories, DAPs were involved in metabolism (27.76 %), stress/defence (11.28 %), transport (9.91 %), transcription/translation (8.42 %), storage proteins (7.81 %), protein synthesis/assembly (6.32 %), transport (4.99 %), signal transduction (3.84 %), and unknown function (9.29 %). Subcellular localizations of the 807 identified DAPs were predicted also using GO and 16.23 % of DAPs were located in the plasma membranes, 14.62 % in the cytoplasm, and 6.82 % in the ribosomes, but 35.94 % were unknown (Fig. 3B). The results of molecular functions of the 807 identified DAPs were: 21.93 % of DAPs had the function of binding, 14.62 % had catalytic activity, and 13.14 % had structural molecule activity, but the molecular function of 16.36 % DAPs were unknown (Fig. 3C).

These DAPs were further annotated based on the COG database. Of these DAPs, 766 proteins were assigned functional annotations grouped into 23 categories. A total of 13.84 % of 807 proteins were involved in posttranslational modification, protein turnover, chaperones, 11.32 % in general functional prediction only, 6.81 % in energy production and conversion, and 8.17 % in translation (Fig. 4).

To investigate the involvement of these proteins in biological functions, 514 (63.69 %) DAPs were mapped to 54 pathways in the KEGG database (Table 1 Suppl.). Metabolic pathways was the most represented pathway (37.21 %), followed by biosynthesis of secondary
metabolites (18.04 %), and ribosome (5.81 %). A few DAPs were involved in pyrimidine metabolism (0.92 %) and DNA replication (0.92 %). KEGG enrichment results showed that most of these DAPs were involved in saccharide and energy metabolism pathways, such as oxidative phosphorylation, glycolysis/gluconeogenesis, and pyruvate metabolism. However, many proteins could not be annotated according to the NCBI, GO, KEGG, and
To verify whether the differences in protein abundance were reflected at the transcriptional level, and to confirm the authenticity and accuracy of the proteomic analysis, 12 genes (two genes selected from each of the six clusters), were analyzed by qPCR at 48 HAI in F₁ and its parental lines. From these 12 genes, 5 genes encoding NAD(P)H-quinone oxidoreductase subunit 5 (NdhF), glucose-1-phosphate adenylyltransferase (GPAT), putative PDI-like protein, 40S ribosomal protein S9, and 26S proteasome non-ATPase regulatory subunit (PSMD) showed similar trends when compared with the iTRAQ analysis results from both parents. Four genes encoding β-glucosidase, triticain α, β-ATP synthase subunit, and 6-fructosyltransferase, showed similar trends when compared with the iTRAQ analysis result from at least one parent. However, 3 genes encoding aspartic proteinase, thioredoxin, and protein phosphatase 2A structural subunit, did not show any similar trends compared with iTRAQ analysis.

**Discussion**

The experiments of grain germination show that a higher germination ability was manifested in the F₁ hybrid (RS5322) as evidenced by an early radicle emergence along with the vigorous growth of the radicles compared with its parental lines (RS53 and SN22, Fig. 1). The rapid germination was associated with a vigorous growth in the F₁ hybrid throughout the germination period. In this study, the grain weight of the F₁ hybrid was significantly lower than of the female parent but did not differ from that of the male parent. The results indicate that the rapid grain germination and superior seedling growth observed in the F₁ hybrid did not correlate with grain mass in these wheat lines. The hybrid RS5322 has obvious advantages in germination rate and grain vigour, so it is suitable to be used as hybrid material in studying hybrid advantages in germination.

To further elucidate the molecular mechanism of grain germination in F₁ hybrid in wheat, a differential proteomic analysis was completed between the F₁ line RS5322 and its parental lines (RS53 and SN22) using iTRAQ-based strategy. As a result, 807 DAPs were identified, among which, 383 were down regulated and 420 were up-regulated in F₁ compared with its parental lines. Bioinformatics analysis showed that 636 DAPs were annotated in 38 functional groups, 764 DAPs were classified into 23 COG categories, and 538 DAPs were enriched in 65 KEGG pathways. The potential roles of some DAPs in grain germination were discussed as follows.

As it is well known, the grain germination is degradation progress of storage material as it is a high energy-requiring process. Grain reserve mobilization or degradation of starch (Job 2005), protein (Maria et al. 2014, Shaik et al. 2014, Zhao et al. 2018) occurs during seed germination. The saccharide degradation is one of the basic metabolic pathways and its main physiological function is to provide energy. In this study, iTRAQ results revealed that many of the pathways contained DAPs associated with saccharide and energy metabolism. As shown in Table 1 Suppl., DAPs significantly enriched in starch and sucrose metabolism (12 DAPs), glycolysis/gluconeogenesis metabolic pathway (8 DAPs), pyruvate...
metabolism (4 DAPs), TCA cycle (5 DAPs), and oxidative phosphorylation (14 DAPs). Glycolysis is the metabolic pathway that converts glucose into pyruvate. The proteome of germination between non-dormant cultivated rice and the dormant wild rice seeds was investigated and showed that the proteins involved in sucrose cleavage and glycolysis decrease, and those involved in ATP and CoQ synthesis and proteolysis increase in endosperm of germinated seeds (Xu et al. 2016). The TCA cycle is a series of chemical reactions used by all aerobic organisms to release stored energy through the oxidation of acetyl-CoA derived from sugars, fats, and proteins, into ATP and carbon dioxide. Proteomic study has been performed in dormant sunflower seeds during imbibition and germination at permissive and non-permissive temperatures for germination, respectively. The results showed that TCA cycle and glycolysis are more active in germinating seeds than in non-germinating seeds (Xia et al. 2018). Oxidative phosphorylation is the

Fig. 5. Comparison of mRNA expression patterns of 12 representative differentially abundant proteins in hybrid RS5322 and its parental lines RS53 and SN22 at 48 hours after imbibition by isobaric tags for relative and absolute quantitation and by quantitative PCR. The expressions of proteins and mRNA in the hybrid were taken as controls in each protein/gene, respectively. The expression of protein and mRNA in the parents were expressed as the ratio of the parent to the hybrid. Blue columns represent protein abundance patterns, red lines represent mRNA expression patterns. F - female parent RS 53, H - hybrid line RS5322, M - male parent SN22.
metabolic pathway in which the mitochondria in cells utilize energy released by the oxidation of nutrients to produce ATP. Mitochondrial oxidative phosphorylation resume shortly imbibed seed after dormancy period (Pergo et al. 2011). Our results showed that many of DAPs involved in starch degradation and energy metabolism were up-regulated in germinated F1 hybrid. Among these proteins, glyceraldehyde-3-phosphate dehydrogenase (GAPDH) plays an important role in glycolysis/gluconeogenesis and it is an enzyme of sugar phosphate metabolism that catalyzes the reversible interconversion of glyceraldehyde-3-phosphate and 3-phosphoglycerate. GAPDH plays an important role in seed ageing and programmed cell death in *P. sativum* (Chen et al. 2013) and in *Arabidopsis* (Rajjou et al. 2008) during seed germination. Our results showed that the relative abundance of two GAPDH protein isoforms increased (2.88- and 2.31-folds) in hybrid compared with its parents, which indicated it might play a great role in wheat germination. An *Arabidopsis* mutant, ndufs4 (for NADH dehydrogenase [ubiquinone] fragment S subunit 4), lacking complex I of the respiratory chain, has constitutively lowered phosphorylation efficiency, delayed germination and this phenotype can be rescued by application of GA (Meyer et al. 2009). Our results showed that NdHF, a NAD(P)H-quinone oxidoreductase subunit 5 involved in saccharide and energy metabolism pathway was significantly up-regulated in F1 hybrid compared with its parental lines. Meanwhile, a similar result was obtained by qPCR analysis (2.31- and 1.89-folds higher than those of parental lines, respectively). It can be speculated that the changed abundance of the NAD(P)H protein might had the promontional effects during F1 hybrid germination.

Proteins play essential roles in the development of cells, organs, or tissues through the interaction with other molecules or modifications (Gutteridge and Thornton 2005), and protein synthesis and proteolysis are vital for the growth and development of plant (Murray et al. 2006). During the development of grains, storage proteins are sequestered in protein storage vacuoles, which are transformed to protein bodies at the end of maturation and drying. Published data indicate that endopeptidases, including cysteine, serine, aspartic proteases and metalloproteases, are responsible for the degradation of storage proteins during the grain development (Chrispeels et al. 1975, Cornel et al. 1994). As shown in Table 1 Suppl., 12 DAPs significantly enriched in proteolysis group and these proteins included predicted metal-dependent protease of the PAD1/JAB1 superfamily, predicted cysteine protease, subtilisin-like protease, aspartic protease, and ATP-dependent zinc metalloprotease FTSH. In previous research, four cysteine endopeptidases were detected in wheat endosperm during the initial growth after seed imbibition and the activities of all of these endopeptidases increased continuously during germination (Shi et al. 2009). Asparaginyl-endopeptidase activity was detected in endosperms of germinating wheat seeds and the high activity was found during germination (Bottari et al. 1996).

As various kinds of proteins exist in grains including globulins, gliadins, and glutenins, these endopeptidases might have different substrate specificity to the storage proteins in mature wheat endosperm. In our study, several endopeptidase involved in proteolysis were significantly up-regulated in F1 compared with its parental lines, and these important and specific endopeptidases might be tightly correlated with grain germination in F1 hybrid.

After proteolysis, amino acids are decomposed into nitrogen-containing and non-nitrogen-containing parts. The nitrogen-containing part is eventually metabolized to urea and the nitrogen-free part is α-ketonic acid which enters the TCA cycle. Many studies have proved that amino acid metabolisms play a great role during grain germination. Seed germination of *Pinus laricio* was inhibited by humic substances through affecting metabolic processes, and, in particular, the amino acid metabolism (Nardi et al. 2002). Glutamine synthetase in concert with NADH-dependent glutamate synthetase (GOGAT)
constitute the major route of assimilation of ammonium derived from reserve mobilization and glutamic acid/glutamine synthesis in germinating *Medicago truncatula* seeds (Glewarcz et al. 2004). In our study, many identified DAPs were involved in pathways associated with amino acids metabolism, including cysteine and methionine metabolism (9 DAPs), glycine, serine, and threonine metabolism (7 DAPs), arginine and proline metabolism (6 DAPs) (Fig. 6). It has been reported that arginine metabolism is unbalanced in simultaneous biosynthesis and degradation of arginine, which could explain the lower accumulation of storage proteins observed during the late stages of germination (Llebres et al. 2018).

Several studies have investigated the role of ABA, GAs, and their interactions in the regulation of grain germination. ABA is the negative regulator of grain germination and promotes seed dormancy, whereas GA overcomes the effect of ABA by inducing germination (Gabriele et al. 2010, Liu et al. 2010, Nambara et al. 2010, Chen et al. 2017). Nine DAPs significantly enriched in hormones-related pathways were identified in our study. These pathways included DAPs in biosynthesis of plant hormones (4 DAPs) and plant hormone signal transduction (5 DAPs). NCED, 9-cis-epoxyprostenoid dioxygenase is a key enzyme in ABA biosynthesis. Previous studies have demonstrated that *LsNCED4*, a temperature-regulated gene, acts as a germination inhibitor (Argyris et al. 2011). In the present study, two NCED protein showed much lower expression in the F₁ hybrid compared with RS53 and SN22. Thus, lower accumulation of NCED4 in the F₁ hybrid could play an important role during germination even though the mechanism of its regulation and its precise role remain to be elucidated. On the contrary, GA3ox2-1, a 2-oxoglutarate-dependent dioxygenase, which is involved in GA biosynthesis (Pearce et al. 2015) showed much higher expression in the F₁ hybrid. Moreover, AIR12, an auxin-induced in root cultures protein 12, which might be involved in the synthesis and signalling of auxins was also detected in these DAPs. The specific roles and mechanisms of these DAPs in germination needs to be further studied.

Endoplasmic reticulum ER is a subcellular organelle where proteins are folded with the help of lumenal chaperones. Correctly folded proteins are packaged into transport vesicles transporting proteins to the Golgi complex. Misfolded proteins are retained within the ER lumen in complex with molecular chaperones and they are directed toward degradation through the proteasome (Swanton et al. 2003). The ubiquitin/proteasome system (UPS) is a highly regulated mechanism of intracellular protein degradation and turnover. Through the concerted actions of a series of enzymes, proteins are marked for proteasomal degradation by being linked to the polypeptide co-factor, ubiquitin (Eleanor et al. 2018). Proteasomes are important proteases in eukaryotes and regulate many cellular processes, including metabolism, cell cycle, and proteolysis of regulatory proteins (Salomons et al. 2010), and the optimal proteasome activities are required for seed germination in bean (Inès et al. 2014) and Arabidopsis (Chiu et al. 2016). As shown in Table 2 Suppl., 26 DAPs were significantly enriched in protein processing in ER and UPS, these metabolisms included protein processing in ER (16 DAPs), proteasome (7 DAPs), and ubiquitin mediated proteolysis (3 DAPs). In a previous study, a putative E3 ubiquitin ligase ECERIFERUM9 regulates ABA biosynthesis and response during seed germination and post germination growth in Arabidopsis (Zhao et al. 2014). In our iTRAQ results, four PSMD homologs were found to be up-regulated in the hybrid compared with its parental lines. Meanwhile, qPCR results showed the higher expression abundance of a PSMD1 in hybrid (Fig. 5). The higher amount of PSMD protein might improve the degradation of misfold proteins in seed, leading to degradation of the storage and germination. In addition, selective proteolysis of proteins mediated by the ubiquitin pathway is important in the regulation of many biological events. We found that two DAPs involved in ubiquitin-mediated proteolysis were up-regulated in the hybrid compared with its parental lines, which may improve the proteolysis of misfolded proteins and affect germination in hybrid. Our study showed that many proteins involved in modification, processing and degradation of proteins were significantly accumulated in F₁ hybrid, which indicated that protein processing in ER and UPS might play great roles in reserve degradation or plant hormones regulation during germination.

**Conclusions**

Our study led to the identification of DAPs in water-imbibed grains of a wheat F₁ hybrid compared with its parental lines. The identified DAPs were involved in various metabolic and cellular processes; the higher accumulation of proteins related with sugar and energy metabolism and proteolysis in the F₁ hybrid indicates that a higher energy supply is very important for higher germination ability. Notably, the abundance of NCED4 protein, involved in biosynthetic pathway for ABA, and acting as an inhibitor in grain germination was lower in germinated grains of the F₁ hybrid. In addition, more DAPs were found to be enriched significantly in protein processing in ER and UPS. Thus, differential accumulation of these proteins and their related metabolism processes appear to be a part of the tight regulation of molecular events associated with higher grain germination in the F₁ hybrid.

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