Metabolomic Profiling of Dongxiang Wild Rice Under Salinity Demonstrates the Significant Role of Amino Acids in Rice Salt Stress

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Dongxiang common wild rice is a precious rice germplasm resource for the study and improvement of salt tolerance in rice. The metabolism profile of Dongxiang wild rice (DXWR) under salinity was determined by high performance liquid chromatography-mass spectrometry (HPLC-MS) to find differential metabolites and screen potential biomarkers for salt-tolerant rice varieties. A global untargeted metabolism analysis showed 4,878 metabolites accumulated in seedlings of Dongxiang wild rice. Principal component analysis (PCA) and orthogonal partial least squares-discriminant analysis (OPLS-DA) results provided a clear metabolism discrimination between DXWR under control and DXWR under salinity. A total of 90 metabolites were significantly changed (49 upregulated and 41 downregulated) under salinity, of which the largest increase was in DL-2-Aminoadipic acid (27.08-fold) and the largest decrease was in L-Carnitine (0.014-fold). Amino acids and nuclear glycosides were mainly upregulated, while carbohydrates and organic acids were mainly downregulated in the salt-treated group. Among the top 10 upregulated metabolites, five kinds of differential metabolites were amino acids. According to the survival rates of the seedlings under salinity, we selected three backcross inbred lines of DXWR with survival rates above 80% as salt-tolerant progenies (pro-DS) and three backcross inbred lines with survival rates below 10% as non-salt-tolerant progenies (pro-NDS) for an amino acid change analysis. This analysis found that the change in L-Asparagine (2.59-fold) was the biggest between pro-DS and pro-NDS under salinity, revealing that the contents of L-Asparagine may be one of the indices we can use to evaluate the salt tolerance of rice varieties.

Keywords: salinity, dongxiang wild rice, metabolome, amino acid, L-asparagine

INTRODUCTION

Rice is a crucial cereal crop in developing countries as it feeds more than half of the global population. However, soil salinity is one of the major environmental stresses that influences rice growth and development in many rice cultivation regions because rice is a glycophyte and, therefore, highly susceptible to salinity stress, especially at the seedling and reproductive stages (Huang et al., 2020; Zhang et al., 2021b). Furthermore, there are $1 \times 10^9$ hectares of saline-alkali...
land worldwide, accounting for about 20% of total agricultural land (Munns and Tester, 2008; Liu et al., 2017). In addition, the salinization area is still expanding. It has been estimated that nearly 50% of all the cultivated land in the world will be salinized by 2050 (Zhu, 2016), posing a major threat to sustainable agricultural development and food security. Thus, the development and cultivation of salt-tolerant cultivated rice are some of the effective strategies to increase the utilization rate of salinity land, enlarge the area of arable soil, and ensure the national food supply.

As an ancestor of modern rice, cultivated rice species were mostly domesticated from common wild Oryza rufipogon species, but some abiotic stress-related traits were lost in the process of domestication. One particular variety that descended from this process was Dongxiang common wild rice, which was discovered in Dongxiang County (N 28°14’), Jiangxi, China, in 1978 (Tian et al., 2006; Xie et al., 2010) and is the northernmost region where any species of wild rice grows. This variety is a rich source of potential genes related to high grain yield, disease and insect resistance, cytoplasmic male sterility, fertility restoration, wide cross-compatibility, cold, and drought tolerance (Quan et al., 2018; Li et al., 2019; Qi et al., 2020; Wu et al., 2020). Zhou et al. (2016) also found that, compared with cultivated rice, Dongxiang wild rice had a very high survival rate under salt stress, suggesting that it is a precious plant germplasm resource for the genetic improvement of salt tolerance in cultivated rice.

Salt stress is the combined effect of osmotic and ion toxicity (primary effect) and oxidative stress (secondary effect). Plants respond to salt stress through many strategies, including the selective preferential uptake and exclusion of K+ ions over Na+ ions (Silveira et al., 2003; Hanin et al., 2016; Su et al., 2020) and the elimination of reactive oxygen species (ROS) through the antioxidant defense system (Tofighi et al., 2017; Wu et al., 2017). Moreover, the synthesis of small organic solutes such as proline, glycine, polyamines, glucose, inositol, glycerol, sugars, and betaine enhances the osmotic pressure of the plant cell against salinity (Latz et al., 2013; Nan et al., 2016).

Metabolomics plays a vital role in studying the changes of metabolites caused by external environment changes or genetic modification, which is the closest research field of the phenotype, and is also one of the hotspots in the field of genomic research. For instance, under drought, high temperature, and multiple salt stress, Sun et al. (2016) found significant changes in citric acid, malic acid, aspartic acid, glucose, fructose, proline, alanine, and threonine in maize leaves by NMR. Supporting this, Guo et al. (2017) systematically analyzed the changes in the metabolic groups of maize seedlings under salt stress by gas chromatography-mass spectrometry (GC-MS), which found that salt changed plant gluconeogenesis and inhibited photosynthesis, nitrogen metabolism, glycolysis, and the synthesis of many kinds of amino acids. Thus, the identification and quantitative study of plant tissue metabolic changes under salt stress will provide insight into the salinity tolerance mechanisms of rice (Yaish and Kumar, 2015). Furthermore, the metabolic profiling of salt-stressed rice from roots and leaves revealed that salt affected the xylem sap metabolic group and significantly decreased the concentrations of tricarboxylic acid (TCA) cycle intermediates and the shikimic acid pathway (Zuther et al., 2007). The study of different rice genotypes by Hakim et al. (2014) also showed that, with the increase of salt stress, the contents of proline and reducing sugars increased while the levels of chlorophyll content, non-reducing sugars, and grain yield decreased, implicating a high degree of genotypic variation in metabolomes under salt stress. Other previous studies have shown that salt increases ROS and some antioxidants in rice (Formentin et al., 2018; Jana and Yaish, 2020). Although technological progress has enabled the usage of various analytical techniques, such as gas chromatography–time-of-flight mass spectrometry (GC-TOF-MS), to study the global metabolites in specific stressed cells or tissues, few studies have clarified the metabolomic changes of O. rufipogon due to salt stress.

The discovery and identification of key metabolites play an essential role in the mechanism of salt tolerance. As such, the purpose of this study was to determine the mechanism of salt tolerance based on non-targeted metabolomic analysis. Therefore, in this study, ultra-high performance liquid chromatography-quadrupole time-of-flight mass spectrometry (UHPLC-Q-TOF-MS) and ultra-high performance-multiple reaction monitoring-mass spectrometry (UHPC-MRM-MS) analysis techniques were used to analyze and evaluate the functions of the differentially accumulated metabolites in the seedlings of Dongxiang wild rice and their introgressive lines under control and salt treatments. The results obtained from this study revealed that Dongxiang wild rice could accumulate a large number of metabolites in varying degrees under salinity, while a few of them played a potential role in the mechanism of salt tolerance.

**MATERIALS AND METHODS**

**Plant Materials**

The seeds of Dongxiang common wild rice (Oryza rufipogon Griff.) DY80 were collected in an experimental paddy field at the Rice Research Institute, Jiangxi Academy of Agricultural Sciences, Jiangxi, China. Cultivated rice R974 (recurrent parent) and DY80 (donor parent) were used to produce the F1 hybrid. Then, the F1 hybrid was backcrossed to R974 to form a BC1F1 population. Finally, the BC1F1 population (backcross inbred lines, BILs) was obtained using the BC1F1 population for nine consecutive inbred times. Dongxiang common wild rice (DXWR) and BILs were used to study rice metabolism under salt stress.

Mature seeds were germinated in the dark on multiple filter papers moistened with double-distilled water in a petri dish, and the uniformly germinated seeds were sown in 96-well plates supported by a plastic container. The seedlings were grown under photoperiodic conditions (light for 16 h and darkness for 8 h) at 28°C in a lightproof chamber, and the growing culture solution was updated two times a day. After 14 days of seedling growth, the culture solution was replaced with the same solution supplemented with 0 and 200 mM of NaCl, as previously described (Zhou et al., 2016). For the metabolomic analysis, 20 seedlings were collected and mixed to minimize the effect of metabolome unevenness among the plants. The experiment was designed with a random complete block and repeated six times.
Metabonomic Analysis

Extraction of Tissue Metabolites

Plant tissues (60 mg) homogenized in liquid nitrogen were ultrasonically broken with 1 ml of a methanol/acetonitrile (2:2:1, v/v) solution at a 4°C temperature for 30 min (Villaret et al., 2020). The broken tissues were incubated for 1 h to precipitate protein, then centrifuged at 13,000 × g with a micro-centrifuge at 4°C for 10 min. Afterward, the supernatant was collected and dried in a speed vacuum concentrator. The process was then repeated six times for each sample (biological replicates).

UHPLC-Q-TOF MS Analysis

Rice extracts were analyzed using the Agilent 1290 ultra-high-performance liquid chromatography system (Agilent Technologies, California, USA) coupled with electrospray ionization quadrupole time-of-flight mass spectrometry system (AB SCIEX, Framingham, USA). The samples were separated by chromatography and randomly injected into an ACQUITY UPLC BEH column (2.1 × 100 mm, with a particle size of 1.7 µm) (Acquity, Waters, Milford, MA, USA). In the process of analysis, the sample tray was kept at 4°C. The injection volume was 2 µl and the column was kept at 25°C. Quality control (QC) samples are injected five times for analytical control.

The flow rate for the mobile phase was set at 0.3 ml/min throughout the gradient. For eluent A, 25 mM of ammonium acetate, 25 mM of ammonium hydroxide, and double-distilled water were used, while acetonitrile was used as eluent B. The following gradient profile was employed: 0–1 min, 95% B; 1–14 min, 95–65% B; 14–16 min, 65–40% B; 16–18 min, 40% B; 18–18.1 min, 40–95% B; 18.1–23 min, 95% B.

A dual electronically stored information source was operated in positive and negative ionization modes. The detailed mass spectrometry (MS) conditions were set according to the method of Julijana et al. (2013). For data acquisition, the time-of-flight mass spectrometry (TOF MS) scan at a mass-to-charge ratio of 60–1,000 Da was set at an accumulation time of 0.2 s/spectra, and product ion scan at a m/z of 25–1,000 Da was set at an accumulation time of 0.05 s/spectra. The secondary MS was performed by information-dependent acquisition (IDA). The IDA parameters were set as follows: exclude isotopes within 4 Da and have six candidate ions to monitor per cycle.

Data Processing and Multivariate Statistical Analysis

The raw data files were converted to a mass spectrometer output file format using the software tool mzConvert (ProteoWizard, proteowizard.sourceforge.net). Retention time alignment, peak integration, and peak alignment were then performed with XCMS software. Afterward, peaks were picked using the cenWave algorithm [(parameters: m/z = 25 ppm, prefilter = c (10, 100), peak width = c (10, 60)] and grouped using the Obiwarp algorithm (parameters: bw = 5, mzwid = 0.25, minfrac = 0.5) (Benton et al., 2015). The peak of the missing values >50% in the group were then removed from the data matrix. After data processing, a two-dimensional matrix was generated, which was composed of m/z and retention time (RT) data pairs, and the mass value and intensity of the peak were output to Excel for further chemometric analyses.

A multivariate statistical analysis of the dataset was carried out using the SIMCA-P software (version 14.1, UMETRICS, Umeå, Sweden) (Nicholson et al., 1999). A principal component analysis (PCA), which consists of score plots and loading plots, showed the contrast between the different samples, while the load chart explains the inherent changes in the data matrix. The identification markers of different rice varieties were screened by an orthogonal partial least square discriminant analysis (OPLS-DA). A hierarchical clustering analysis (HCA) was performed by Pictometry (Caraux and Pinloche, 2005) using z-score transformed metabolite abundances and clustering on their Pearson distances (Minoura et al., 2012).

Pathway Enrichment and Clustering Analysis

To clarify the metabolic pathway of secondary metabolites, the compounds identified in the above databases were submitted to the Kyoto Encyclopedia of Genes and Genomes (KEGG) (http://www.genome.jp/kegg/pathway.html). A hierarchical clustering analysis (HCA) was performed by PermutMatrix (Caraux and Pinloche, 2005) using z-score transformed metabolite abundances and clustering on their Pearson distances (Minoura et al., 2012).

Amino Acid Determination

Metabolite Extraction

The sample was ground into powder after freeze-drying. Then, 60 mg of each sample were accurately weighed and transferred to a centrifugal tube. After adding two steel balls and 1,000 µl extraction solvent (acetonitrile-methanol-water 2:2:1, including the mixture of internal standards of isotopically-labeled) pre-cooled at −20°C, the samples were mixed with a vortex, homogenized at 40 Hz for 4 min, and sonicated for 5 min in an ice-water bath. The homogenate and ultrasonic cycles were repeated three times and then incubated at −40°C for 1 h and centrifuged at 12,000 rpm at 4°C for 15 min (Mushtaq et al., 2014). Finally, the clear supernatant of 80 µl was transferred to an auto-sampler vial for UHPLC-tandem MS (MS/MS) analysis.

UHPLC-MRM-MS Analysis

Rice extracts were separated by chromatography and randomly injected into a Waters ACQUITY UPLC BEH amide column (100 × 2.1 mm, 1.7 µm) using an Agilent 1290 Infinity II series UHPLC System (Agilent Technologies). In the process of analysis, the sample tray temperature was kept at 4°C. The injection volume was 1 µl, while the column temperature was kept at 35°C. Mobile phase A was a 1% formic acid aqueous solution, while the mobile phase B was a 1% formic acid acetonitrile solution.

An Agilent 6460 triple quadrupole mass spectrometer (Agilent Technologies) equipped with a Jet Stream Technology Ion Source (AJS) electrospray ionization (Agilent Technologies, California, USA) (AJS-ESI) interface was applied for MS. The parameters of the ion source were set by Dunn et al. (2011).

Data Processing and Multivariate Statistical Analysis

The lower detection limits and lower limits of quantitation were determined by the signal-to-noise ratio. According to the US FDA guidelines for the validation of the bioanalytical method
(Saccenti et al., 2014), lower detection limits (LLODs) and lower limits of quantitation (LLOQs) are defined as the analyte concentrations that lead to peaks of signal-to-noise ratio (S/N) at 3 and 10, respectively. In the sample, the content of the target metabolite was calculated according to the following formula.

\[ c_M [nmol \cdot g^{-1}] = \frac{C_F [nmol \cdot L^{-1}] \cdot V [mL]}{m [mg]} \]

Where \( C_F \) is the final concentration, \( C_M \) is the metabolite concentration, \( V \) is the final volume, and \( m \) is the quality of the sample.

RESULTS

Metabolic Profiling of Salt-Stress Responses in Wild Rice
To investigate the influence of salt conditions on rice seedling metabolites, both rice seedlings were analyzed under non-salt and salt conditions using an ultra-high performance liquid chromatography-quadrupole time-of-flight mass spectrometry-based metabolomic approach. As shown in Supplementary Figure 1, there was a significant difference in the intensities of the compounds peaks in the total ion chromatography (TIC) plot obtained by the UHPLC-Q-TOF-MS analysis between the two samples, especially for peaks between the retention times of 4 and 12 min in the TIC plot (Supplementary Figure 1). According to the data treatment procedures, for an efficient comparison of the two organic rice samples under non-salt and salt conditions, 4,484 positively correlated variables were detected and 3,441 negatively correlated variables were detected (Supplementary Tables 1, 2).

PCA and OPLS-DA
The 4,484 positively correlated variables and 3,441 negatively correlated variables were normalized based on internal standards and intensities and used to perform multivariate statistical analyses. The quality of the model was evaluated by R2 and Q2 values. R2, which explains the proportion of variance, explained, and predicted in the PCA model separately. Specifically, 65.7 and 69.5% of the positively correlated variances (R2) and negatively correlated variances (R2), respectively, were explained by the first two components. On the other hand, the Q2 values were 45.2 and 51.5%, respectively (Figure 1), for the same variances. All these values showed the validity of the model. These results suggested that the, and DXWR under salinity (DS) and DXWR without salt treatment (DN) samples were distributed in different groups according to the first two principal components, while there were no outlier samples. This indicated that there were metabolic differences between the DN and DS groups.

To strengthen the distinction between the experimental groups and improve the validity and reliability of the results, an OPLS-DA was used to accurately evaluate the metabolic patterns of these samples. In Supplementary Figure 2, the OPLS-DA parameters of the model were expressed by the R2 of 0.834 and Q2 of −0.329 (Supplementary Figure 2). These parameters exhibited the better stability and predictability of the model and effectively reflected the metabolic differences between the DN and DS responses to salt stress.

Hierarchical Cluster Analysis of Total Metabolites
By clustering all the identified metabolites, the metabolites of six samples in the DS group were clustered into one group (Figure 2), indicating that the metabolic patterns of the samples were similar after salt treatment. At the same time, the DN and DS groups formed relatively clear high and low expression regions, while the high and low expression regions of the DN group were opposite to those of the DS group, indicating that salt stress had a significant effect on rice metabolism.

Differential Metabolite Analysis in Rice Under Salt Stress
The results showed that salt stress significantly affected the distribution of metabolites in rice. As shown in Figure 3, the volcano map of differential metabolites (fold change > 2 and \( p < 0.05 \)) showed significant differences between the DN and DS groups.

Using a variable importance in the projection screening >1 at a level of \( p < 0.05 \), a total of 90 differential metabolites were isolated in the positive ion mode and negative ion mode. Compared with DN, 49 differential metabolites in DS were upregulated and 41 were downregulated. The 10 differential metabolites with the most extensive upregulated range were DL-2-Aminoacidic acid (27.08-fold), Uracil (26.89-fold), D-Quinovose (23.38-fold), 2'-O-Methyluridine (22.4-fold), 2-Oxoadipic acid (13.438), Indole-3-carboxylic acid (13.26-fold), L-Asparagine (10.67-fold), L-Proline (9.12-fold), L-Histidine (10.49-fold), L-Tryptophan (6.881-fold), and L-Glutamine (6.32-fold). The most downregulated differential metabolite was L-Carnitine (0.014-fold).

According to Figure 4, there were 88 differential metabolic species, including 18 amino acids and their derivatives (all are up-regulated), 27 organic acids and their derivatives, 16 nucleosides and their metabolites, 5 sugars, 5 alcohols, 4 organic amines, 2 lipids, and 11 other substances. Similarly, in the negative ion mode, a total of 54 differential metabolites were screened. Compared with the control group, there were 35 upregulated metabolites in DS, including 15 amino acids and their derivatives, of which S-Methyl-5'-thioadenosine (40.107) was the most upregulated (Supplementary Table 3). There were 19 downregulated metabolites, of which the largest decrease was in D-Galactarate (0.015) (Supplementary Table 3). From the differences in the types and expression of different metabolites, salt stress had the greatest effect on the amino acid metabolism of rice.

KEGG Pathway Analysis for Differential Metabolites
The Kyoto Encyclopedia of Genes and Genomes is one of the databases commonly used in communication research, as researchers use it to read a large amount of literature to describe many metabolic pathways and their relationships to a specific
graphical language. In this study, a KEGG analysis was used to annotate differential metabolites. Compared with the metabolites of the DN group, the DL-2-Aminoadipic acid with the largest upregulation of the metabolites in the DS group was annotated in the amino acid biosynthesis and metabolic pathway. As shown in Figure 5, the KEGG pathway enrichment analysis of the differentially expressed metabolites of DN/DS compared with the control group by Fisher precise test showed that significant changes had taken place in important pathways such as Aminoacyl-tRNA biosynthesis, ABC transporters, Alanine, Aspartate and Glutamate metabolism, Purine metabolism, and Galactose metabolism. Among the top 20 KEGG pathways, 9 were related to amino acid synthesis. Next, a conjoint analysis of the metabolome and transcriptome was performed (Zhou et al., 2016), which found that 6 pathways (251 genes) were involved in amino acid metabolism among top the 10 KEGG pathways. The results indicated that amino acid metabolism could be used as a physiological indicator to evaluate rice salt tolerance.

Morphological Traits in the Progenies of Dongxiang Wild Rice Under Salinity

According to previous studies, more than 80% of Dongxiang wild rice under seedlings survived under a 200-mM NaCl treatment for 12 days, which is a much higher amount than what common cultivated rice can survive (Zhou et al., 2016). To study the changes in the metabolites in the introgression lines of DXWR, 2-week-old seedlings of 130 BILs were exposed to a hydroponic solution with 200 mM of NaCl for 12 days. Under normal growth conditions, the 130 BIL seedlings survived and retained their green color. Under salt stress conditions, however, most BIL leaves became curled up, wilted, and lost their green color. Thereafter, the seedlings were returned to normal growth conditions, resulting in most of BILs not being able to restore normal growth and only a small number of BILs resuming growth. According to the survival rates of the seedlings under salinity, 3 BILs with survival rates above 80% were selected as AP17, AP92, and AP114 and 3 BILs with survival rates below 10% as AP91, AP290, and AP10, respectively, for a differential metabolite analysis (Figure 6). Additionally, AP17, AP92, and AP114 were defined as salt-tolerant progenies (pro-DS), whereas AP91, AP290, and AP10 were non-salt-tolerant (pro-NDS).

Changes of Amino Acid Content in the Progenies of Dongxiang Wild Rice

According to the metabonomic analysis, salt stress significantly affected the amino acid metabolism of Dongxiang wild rice. In order to study the changes in the amino acids in the progenies of Dongxiang common wild rice under salt stress, Student’s t-test ($p < 1$) was used to screen the differential metabolites between salt-tolerant and non-salt-tolerant progenies under salinity (Table 1). In the two groups, the five amino acids with concentrations more than 4,000 (nmol/g) were the same, namely, L-Serine, L-Glutamic acid, L-Alanine, L-Glutamine, and L-Aspartic acid. Moreover, the three amino acids with the lowest expression were the same, namely, 4-Hydroxyproline, 3-Methyl-L-histidine, and 1-Methyl-L-histidine. The change in L-Asparagine was the
biggest between pro-DS and pro-NDS under salinity, while the fold change of pro-DS/pro-NDS was 2.5-fold. Moreover, the contents of L-Aspartic acid (1.44-fold) and L-Glutamine (1.77-fold) were also upregulated in pro-DS. On the contrary, Alanine, Methyl-L-histidine, and L-Citrulline were downregulated in pro-DS. Thus, the changes in amino acid content in the progenies were consistent with the metabonomic characteristics of DXWR.

**DISCUSSION**

In rice salinity phenotyping related to the seedling, the major visible phenotypic symptoms include plant survival rates, phenotypic expression (yellowing tip burning, browning, and rolling in younger leaves), and poor root growth. Salt stress also influences numerous biochemical processes, i.e., chlorophyll content, CO2 emission, malondialdehyde (MDA), and superoxide dismutase (SOD) (Hussain et al., 2019). The symptoms of salinity have also been often found to overlap with those of other abiotic stresses. Therefore, there is, unfortunately, no single clear-cut trait or criterion in rice salinity phenotyping. Previous studies have shown that physiological and biochemical processes and more than 6,800 genes of DXWR were changed under salt stress (Zhou et al., 2016; Ai et al., 2020), while the overexpression of miR1861h from DXWR in particular increased tolerance to salt stress in cultivated rice (Ai et al., 2020). In this study, a total of 68 metabolites were significantly changed under salinity. Most of the upregulated metabolites were nuclear glycosides and amino acids. Carbohydrates and organic acids, in contrast, were mainly downregulated. Furthermore, 15 differential metabolites significantly or extremely significantly affected 20 metabolic pathways such as phenylalanine, tyrosine, and tryptophan biosynthesis, alanine-aspartic acid-glutamic acid metabolism, and the TCA cycle. Of these significant pathways, nine were amino acid metabolic pathways, indicating that amino acid metabolism can be used as a physiological indicator to evaluate rice salt tolerance.

The accumulation of osmotic protective agents is very important to the alleviation of the intracellular osmotic imbalance caused by salt stress in plants. Proline, in particular,
is an important osmotic regulator that maintains the osmotic balance between the protoplast and the environment. Previous studies have shown that salt stress significantly increased levels of proline accumulation (Forlani et al., 2019; Gerona et al., 2019). Specifically, the fold changes of D-Proline and L-Proline were 7.893 and 9.124 in DXWR under salinity, respectively. Hussain et al. (2019) also found that proline contents were increased in two cultivated rice varieties under salt stress. Additionally, Ganie et al. (2019) revealed that the activity of the proline biosynthesis gene OsP5CS was inhibited by the feedback of proline content and had a significant effect on rice resistance to salt stress. Furthermore, glutamic acid, the substrate for the synthesis of proline, was also upregulated, a finding that was consistent with the findings of a study by Zhang et al.
FIGURE 4 | Differential metabolites of DN and DS. The sizes of the circles reflect FC in abundance between DN and DS; red indicates significantly upregulated in DS, green indicates significantly downregulated. DN is DXWR without salt treatment and DS is DXWR under salinity.
FIGURE 5 | The Kyoto Encyclopedia of Genes and Genomes (KEGG) classification of the differential metabolites (A) and transcriptomes (B) between DN and DS. The metabolic pathways annotated in KEGG are classified according to the type of pathway. The Abcissa is the number of metabolites annotated to that pathway and the ratio of the number of metabolites annotated to the total number of annotated metabolites in the corresponding pathway. DN is DXWR without salt treatment and DS is DXWR under salinity.
FIGURE 6 | Comparison of phenotypes between saline-treated plants and blank controls (CK).
on the metabonomics of salt-tolerant rice varieties in Changbai.

The content of basic and neutral amino acids was increased in Dongxiang wild rice under salinity. Moreover, S-Methyl-5′-thioadenosine (40.107) was the most upregulated in negative ion differential metabolites. Wang et al. (2019) found that more basic (Lys, Arg) and neutral (Val, Ala) amino acids were transported and accumulated in the amino acid permease 5 (OsAAP5) overexpression lines than in the wild type. Additionally, OsAAP5 could regulate tiller bud outgrowth by affecting zeatin levels. Zeatin is ubiquitous in most bioassays with extremely high activity and is considered to play an important role in plant growth and development (Mok and Mok, 2001). Adenosine and S-Methyl-5′-thioadenosine are also synthetic precursors of trans zeatin formed by adenosine phosphate-isopentenytransferase (IPT). There are seven IPT genes in Arabidopsis (Arabidopsis thaliana), of which IPT3 is upregulated by nitrate (Sakakibara, 2006). The absorption and assimilation of nitrate by plants are usually closely combined with nitrogen (N) metabolism. Nitric oxide, produced as part of N metabolism, is one of the most widely used signal molecules in organisms. Furthermore, polyamines and nitric oxide are related to the salt tolerance of plants (Ahmad et al., 2016; Khushboo and Gyan, 2019).

Asparagine is an important amino acid for the long-distance transport of nitrogen in plants. Degenkolbe et al. (2013) reported that higher levels of asparagine (Asn) were predominantly found in drought-sensitive cultivars under drought stress. However, little is known about the effect of Asn on rice salt tolerance. In this study, the leaves of DXWR and pro-DS presented slighter wilting than pro-NDS after salt stress. The contents of L-Asparagine were also increased 10.675-fold in DXWR under salt stress. Moreover, the leaves of DXWR and pro-DS presented slighter wilting than pro-NDS after salt stress. The contents of L-Asparagine were also increased 10.675-fold in DXWR under salt stress. More interestingly, the change in L-Asparagine (2.59-fold) was the biggest between pro-DS and pro-NDS. Asparagine has a high concentration of C/N and is more stable compared with other amide compounds. Furthermore, two asparagine synthase genes were identified in rice, with one OsASN1 located on chromosome 3 and the other annotated as OsASN2 located on chromosome 6. The analysis of the T-DNA insertion mutant showed that OsASN1 is involved in the regulation of rice development and is specific for tiller outgrowth (Luo et al., 2019). In a previous study, OsASN1 and cyanolalanine synthetase were upregulated under salt stress in DXWR seedlings (Zhou et al., 2016). Cyanolalanine synthetase in salt-tolerant rice catalyzes the conversion of cyanide to cyanolalanine, which is further transformed into asparagine and enters the amino acid pool of rice. This may also be one of the reasons why, among the top ten upregulated metabolites, five kinds of differential metabolites were amino acids. Therefore,
the contents of L-Asparagine may be one of the indices to evaluate the salt tolerance of rice varieties. However, the related speculations need to be further studied and verified.

DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article/Subplementary Material, further inquiries can be directed to the corresponding authors.

AUTHOR CONTRIBUTIONS

YC and WH: investigation, methodology, formal analysis, and writing of the original draft. FZ: methodology and supervision. XL: resources and methodology. BH: conceptualization, resources, methodology, writing, reviewing and editing, supervision, and project administration. JX: conceptualization, resources, methodology, writing, review and editing, supervision, and project administration. All authors contributed to the article and approved the submitted version.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fpls.2021.729004/full#supplementary-material

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