Integrated Analysis of Transcriptome and Metabolome Reveals the Mechanism of Chlorine Dioxide Repressed Potato (Solanum tuberosum L.) Tuber Sprouting

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Sprouting is an irreversible deterioration of potato quality, which not only causes loss in their commercial value but also produces harmful toxins. As a popular disinfectant, ClO₂ can inhibit the sprouting of potato tubers. Using transcriptomic and metabolomic approaches to understand the repressive mechanism of ClO₂ in potato sprouting is yet to be reported. Sequencing the transcriptome and metabolome of potatoes treated with ClO₂ in this study revealed a total of 3,119 differentially expressed genes, with 1,247 and 1,872 genes showing down- and upregulated expression, respectively. The majority of the downregulated genes were associated with plant hormone signal transduction, whereas upregulated differential genes were associated primarily with biological processes, such as phenylpropanoid biosynthesis and the mitogen-activated protein kinase (MAPK) signaling pathway. Metabonomic assays identified a total of 932 metabolites, with 33 and 52 metabolites being down- and upregulated, respectively. Downregulated metabolites were mostly alkaloids, amino acids, and their derivatives, whereas upregulated metabolites were composed mainly of flavonoids and coumarins. Integrated transcriptomic and metabolomic analyses showed that many different metabolites were regulated by several different genes, forming a complex regulatory network. These results provide new insights for understanding the mechanism of ClO₂-mediated repression of potato sprouting.

Keywords: potato tuber, chlorine dioxide, repression of sprout, transcriptome, metabolome

INTRODUCTION

Potato is one of the most important food crops in the world, with a high nutritional value. However, its postharvest storage remains a challenging step to maintaining quality and minimizing market losses (Huang et al., 2014). During the potato storage in the fresh food market, or storage of tubers for seed harvesting and industrial processing, sudden sprouting occurs (Teper-Bamnolker et al., 2010). Tuber sprouting is controlled by several factors, such as plant hormones, genetic factors, signaling molecules, genotype, temperature, and other storage conditions (Wang et al., 2020).
Sprouting is also determined by metabolic activities that regulate phytohormones to exert growth control (Sonnewald, 2001). Therefore, the application of sprout suppressant is an urgent research target (Corsini et al., 1978; Campbell et al., 2006). Libraries were sequenced using the Illumina HiSeq 2500 platform (Jisi Huiyuan Biotechnology Co., Ltd., Nanjing, China), with 2 × 150 paired-end reads. The high-quality reads were assembled into unigenes following the methods by Conte et al. (2019). Chlorine dioxide (ClO$_2$) has broad and high-biocidal activity because of its strong oxidizing and sanitizing properties (Aday and Caner, 2011). ClO$_2$ was cleared by the Food and Drug Administration (FDA) (Food Drug Administration., 1998) as a safe antibacterial agent for cleaning and treating fruits and vegetables. Chlorine significantly reduces sprouting and effectively controls dry rot, soft rot, and silver scurf of the tubers (Tweddell et al., 2013). For example, cleaning fruit surfaces with ClO$_2$ reduced air-dried Salmonella in tomato and Escherichia coli O157:H7 strain in apples (Du et al., 2003; Pao et al., 2009). In addition, ClO$_2$ treatment maintained quality, reduced rot rate, and extended the shelf life of kiwifruit (Niu et al., 2009), mulberry (Chen et al., 2011), and litchi (Wu et al., 2011), and reduced pericarp browning in longan (Atinut et al., 2019).

Currently, most studies are focused on the use of ClO$_2$ in the control of pathogenic microbes, maintenance of fruit quality, and extending the shelf life of fruit. However, no studies on the repressive mechanism of ClO$_2$ in potato sprouting using transcriptome and metabolome approaches have been reported. This study analyzed the transcriptomic and metabolic profiles of ClO$_2$-treated tubers to uncover the molecular mechanism of ClO$_2$-mediated repression of potato sprouting.

**MATERIALS AND METHODS**

**Materials**

The tubers of the “Atlantic” cultivar were obtained from Kaikai Potato Seed Co., Ltd. (Gansu, China). After discarding the tubers with injuries or infections, potatoes were transferred to the laboratory and stored at 20 ± 3°C until use. A formulation with 12% ClO$_2$ as the active ingredient was purchased from Tianjin Zhangda Technology Development Co., Ltd. (Tianjin, China).

**Treatment of Tubers and Sampling**

After dormancy, tubers were cleaned and disinfected by immersion in 1% sodium hypochlorite for 3 min, rinsed with sterile water, and air-dried. Tubers were then soaked in 180 mg/L ClO$_2$ for 15 min, and a control group (CKH3) was soaked in sterile water. After drying, all treated tubers were stored in the dark at 20 ± 2°C with a relative humidity of 75–85%. Experiments were carried out in three replicates per treatment, with each containing 30 tubers.

**Methods**

**Sampling**

Sampling of the ClO$_2$H$_3$-treated tubers with no sprouts was performed after detecting sprouting in the CKH3 treatments. About 5 g of tuber tissue was cut 1 cm deep under the skin at the bud growth site (bud eye), with a 1 cm stainless steel punch. Samples were wrapped in tin foil paper, frozen in liquid nitrogen, and stored at −80°C until use.

**RNA Sequencing (RNA-Seq)**

Total RNA extraction (three duplicate samples) was performed following the method described by Woolfson et al. (2018). cDNA libraries were developed in three duplicates as described by Foutcart et al. (2006). Libraries were sequenced using the Illumina HiSeq 2500 platform (Jisi Huiyuan Biotechnology Co., Ltd., Nanjing, China), with 2 × 150 paired-end reads. The high-quality reads were assembled into unigenes following the methods by Haas et al. (2013). To count the assemblies, TopHat 2 software was used to map RNA sequence data to the Solanum tuberosum reference genome (DM_1-3_516_R44_potato.v6.1), downloaded from the Potato Genomics Resource (http://spuddb.uga.edu/dm_v6_1_download.shtml).

**Functional Annotation**

The unigenes were queried against the Kyoto Encyclopedia of Genes and Genomes (KEGG), Non-redundant (Nr), Eukaryotic Orthologous Groups of protein (KOG), Uniprot, and Gene Ontology (GO) public databases, with a threshold e-value of <10$^{-5}$ to obtain the functional annotations. The KEGG pathway analyses and GO functional classifications were visualized using the Web Gene Ontology Annotation Plot (WEGO) and KEGG automatic annotation servers, respectively.

**Metabolite Extraction**

The frozen samples were ground in a mixer mill for 1 min at 45 Hz. Approximately, 100 mg of samples were then mixed with 1.5 mL methanol–water (3:1). The samples were homogenized and sonicated for 15 min at 0°C, extracted with a shaker for 24 h at 4°C, followed by centrifugation with 13,800 × g at 4°C for 15 min. The supernatant was filtered through a membrane, the filtrate was diluted twice with methanol–water (3:1), and then transferred to glass vials; 20 µL of this solution was pooled as QC sample for ultra-high-performance liquid chromatography-mass spectrometry (UHPLC-MS) analyses (Sawada et al., 2009).

**UHPLC-MS Analyses**

Ultra-high-performance liquid chromatography-mass spectrometry separation was performed using an EXIONLC System, with mobile phases A and B as formic acid (0.1%) and acetonitrile, respectively. The column and the auto-sampler temperatures were set at 40°C. A Sciex QTrap 6500+ (Sciex Technologies) was applied. Ion source parameters were set as follows: temperature, 400°C; curtain gas, 35 psi; ion source gas 2, 60 psi; DP, ±100 V; ion source gas 1, 60 psi; and ion spray voltage, +5,500/−4,500 V (Zhang et al., 2015). The UHPLC-MS measurements were carried out in five replicates per sample.
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FIGURE 1 | The cluster heat map of the differentially expressed genes (A) and volcanic map (B) in control and ClO<sub>2</sub> treated tubers. Three biological repeats were shown in the figure. CKH3 represents the tubers of safflower treated with sterile water; ClO<sub>2</sub>H<sub>3</sub> represents the flowers of safflower treated with ClO<sub>2</sub>.

Data Pre-Processing, Annotation, and Screening of Differentially Accumulated Metabolites

The MRM data were acquired with the SCIEX Analyst Work Station Software (Version 1.6.3), then the raw (.wiff) MS data files were converted to .txt format using the MSconverter. MS peak detection and annotation were performed in the R program (Kuhl et al., 2012). Differentially accumulated metabolites (DAMs) obtained were screened with pairwise comparison using the Variable Importance in the Projection (VIP) with the first principal component of the OPLS-DA model as >1 and \( p < 0.05 \), Student’s \( t \)-test, as the threshold.

qRT-PCR Analyses

qRT-PCR analyses were performed following the methods of Woolfson et al. (2018), with the same RNA and cDNA samples used in the RNA-seq analysis. The accession numbers of selected genes based on the Potato Genomics Resource assembly database and their specific primers are presented in Supplementary Table 1. The primers for the housekeeping gene, \( efl \), were designed according to Nicot et al. (2005). The RT-qPCR reaction system was prepared as follows: Ultra SYBR mixture, 10 \( \mu l \); RNase-free water, 8.4 \( \mu l \); template cDNA, 0.8 \( \mu l \); and each primer, 0.4 \( \mu l \). RT-qPCR was performed with the Roche LightCycler 480 (Roche, Basel, Switzerland) thermocycler, with the program set as follows: initial 95°C for 10 min, followed by 40 reaction cycles of 95°C for 30 s, 72°C for 30 s, one final cycle of 72°C for 10 min, and one cycle of 40°C for 30 s. The relative gene expression was calculated using the \( 2^{-ΔΔCT} \) method (Livak and Schmittgen, 2001).

RESULTS

Quality Assessments of Sequencing Data and Differential Gene Expression Analyses

A total of 161.77 GB quality clean sequencing reads were obtained, with an average of 5.98 GB clean data per sample, and percentages Q20 and Q30 of bases above 97 and 93%, respectively, were observed. The guanine–cytosine (GC) content of each sample was above 42% after filtration (Supplementary Table 2). All clean reads were compared to the potato reference genome (DM_1-3_516_R44_potato.v6.1), and 89.25–90.10% of sequences from control or ClO<sub>2</sub> treatments were mapped to the reference genome, which demonstrated the high reliability of our data (Supplementary Table 3).

To further investigate the gene expression patterns, differentially expressed genes (DEGs) were screened by pairwise comparison with Fold Change (FC) \( ≥2 \) and False Discovery Rate (FDR) <0.05 and Fragments per Kilobase of transcript per Million fragments mapped (FPKM) values were calculated for each sample to normalize the gene expression. Hierarchical clustering analysis was performed on the filtered differentially expressed genes, and the genes with the same or similar expression behaviors were clustered, and the results of the clustering of differentially expressed genes are shown in Figure 1A. A total of 3,119 DEGs were detected, with 1,247 and 1,872 as down- and upregulated genes, respectively, after ClO<sub>2</sub> treatment (Figure 1B).

GO Annotation and Enrichment

GO classification was used to define the functions of DEGs from the comparisons between control and ClO<sub>2</sub>. As a result, 43
functional DEG categories were identified, including 12 “cellular components”, 10 “molecular functions”, and 21 “biological processes” (Figure 2A). Most DEGs were classified into the cell (1,728) and cell part (1,724) in the cellular components, whereas 1,288, 1,278, 1,534, and 1,425 DEGs were classified in binding, catalytic activity, molecular functions and cellular processes, and metabolic processes, respectively (Figure 2B).

**KEGG Annotation and Enrichment**
 DEGs were screened in the KEGG ortholog database for functional enrichment analyses. As a result, the comparison between the control and ClO$_2$-treated tubers assigned 19 pathways to different categories, including organismal systems, metabolism, environmental information processing, genetic information, cellular processes, and human diseases. In addition,
carbohydrate metabolism, signal transduction, biosynthesis of other secondary metabolites, amino acid metabolism, and lipid metabolism were significantly enriched in DEGs (Figure 3A). Notably, abundant DEGs were involved in phenylpropanoid biosynthesis, MAPK signaling pathway-plant, plant hormone signal transduction, plant-pathogen interaction, starch and sucrose metabolism, and glutathione metabolism (Figure 3B). Interestingly, only the plant hormone signal transduction pathway was enriched with downregulated genes, whereas the other two pathways were enriched with upregulated genes (Figures 3C,D).

Analyses of DEG in Phenylpropanoid Biosynthesis and Plant Hormone Signal Transduction Pathways

Phenylpropanoid biosynthesis and plant hormone signal transduction were the most enriched pathways, with upregulated and downregulated DEGs, respectively. The three differential genes of phenylpropanoid biosynthesis were downregulated, whereas the others were upregulated. RT-qPCR analyses using eight genes associated with the phenylpropanoid pathway and 12 genes of plant hormone signal transduction were used to further
FIGURE 4 | The phenylpropane biosynthesis pathway and the DEGs involved in this pathway (A) (PAL, phenylalanine ammonia-lyase; 4CL, 4-coumarate–CoA ligase; CYP73A, trans-cinnamate 4-monoxygenase; CAD, cinnamyl-alcohol dehydrogenase; COMT, caffeic acid 3-O-methyltransferase; POD, peroxidase; CCoAOMT, caffeoyl-CoA O-methyltransferase; CAG, coniferyl-alcohol glucosyltransferase) and The plant hormone signal transduction and the DEGs involved in this pathway (Continued)
confirm the RNA-seq results (Figure 4). As a result, consistent gene expression patterns were observed between qPCR and RNA-seq (Supplementary Figure 1).

**Metabolomic Data Analyses**

The metabolomic assay identified a total of 932 metabolites. Principal components analysis evidently separated metabolites into treated and control groups, and samples with 95% confidence intervals (Hotelling's t-squared ellipse) were selected for further analyses to screen DAMs (Figure 5A). As a result, 85 DAMs were detected, with 33 and 52 DAMs being down- and upregulated, respectively, after ClO$_2$ treatment (Figure 5B).

**Comparison of Metabolites**

The most abundant DAMs identified were flavonoids, followed by nucleotides and their derivatives, amino acids and their derivatives, alkaloids, coumarins, phenols, miscellaneous, and carboxylic acids and their derivatives (Figure 6A). A total of 52 DAMs were upregulated, whereas 33 DAMs were downregulated in the pericarp of ClO$_2$-treated tubers relative to that of control tubers. The most decreased metabolites were alkaloids, followed by amino acids and their derivatives, nucleotides and their derivatives, carboxylic acids and their derivatives, and phytohormones (Figure 6B). Most increased metabolites included flavonoids, followed by coumarins, nucleotides and their derivatives, phenols, and miscellaneous compounds (Figure 6C).

**Integrated Transcriptomic and Metabolomic Analyses**

The regulatory networks were analyzed by correlating DEGs and DAMs from each treatment. Results showed that multiple genes were positively or negatively regulated by multiple metabolites. For example, 17 DAMs were negatively or positively regulated by 20 or 30 DEGs, respectively, whereas 33 DAMs were negatively or positively regulated by 30 or 20 DEGs, respectively (Figure 7A). About three metabolites, 4-methylumbelliferyl acetate, methoxyindoleacetic acid, and DL-tyrosine, were strongly correlated with nine DEGs, with seven and two genes showing positive and negative correlations, respectively. Similarly, bruceine D and L-3-phenyllactic acid were significantly correlated with 10 DEGs, with eight and two genes showing positive and negative correlations, respectively, and 10 DEGs (three positively correlated, seven negatively correlated), respectively. In addition, (S)-2-acetolactate and vasicine were correlated with nine DEGs, with six and three genes showing positive and negative correlations, respectively, and eight DEGs (five positively correlated, three negatively correlated), respectively. Dalbergioidin and D-alpha-aminobutyric acid were markedly correlated with six DEGs, with one and five genes showing positive and negative correlations, respectively.
Zheng et al. Mechanism of ClO$_2$-mediated repression of potato tuber sprouting. RNA-seq results identified 3,119 DEGs after ClO$_2$ treatment, with the majority showing enrichment in phenylpropanoid biosynthesis, the MAPK signaling pathway plant, and plant hormone signal transmission pathways. Phenylpropanoid biosynthesis and MAPK signaling pathways were significantly activated, whereas the plant hormone signal transduction pathway was significantly inhibited. Consistently, ClO$_2$ treatment could activate phenylpropane metabolism to improve apple tolerance (Zhao et al., 2011) and promote the accumulation of MAPK.
which was shown to regulate browning in longan (Chumyam and Saengnil, 2020). Interestingly, ClO$_2$ treatment hindered signal transduction during potato tuber sprouting. Phytohormones are crucial endogenous regulators of tuber sprouting, and dynamic changes in endogenous hormones related to stages of sprouting were observed in hormone-treated tubers (Aksenova et al., 2013). This suggests that ClO$_2$ repressed potato sprouting by limiting plant hormone signal transduction. In addition, ClO$_2$ could improve the stress tolerance of tubers by activating phenylpropanoid biosynthesis and MAPK signaling pathways. This could be because of the ability of ClO$_2$ to induce the production of reactive oxygen species, which damage cell membranes, eventually leading to the obstruction of signal transduction during tuber sprouting.

Metabolites are important determinants of potato tuber sprouting (Aksenova et al., 2013). During the sprouting process, starch and protein degradation are initiated, and soluble sugars and amino acids are formed (Sonnewald, 2001). Metabolites are the final or intermediate regulatory products of the cell's biological processes (Pichersky and Lewinsohn, 2011), and their accumulation is controlled by numerous endogenous and exogenous factors. Plant growth and development are regulated by metabolites (Luca et al., 2012). Thus, integrated metabolomic and transcriptomic analyses can be used to identify key metabolic pathways and functional genes associated with plant growth and development (Matus, 2016; Wang et al., 2017). Potato sprouting is often accompanied by changes in a large number of metabolites (Sonnewald, 2001). Metabolomic results in this study revealed that most differentially upregulated metabolites associated with the repressing potato tuber sprouting were enriched in flavonoids and coumarins. In contrast, the most differentially downregulated metabolites were alkaloids, amino acids, and their derivatives. Flavonoids have health-promoting benefits because of their antioxidant capacity through cell signaling, pro-oxidative activity, or membrane characteristic mechanism (Mein et al., 2008). Potato tubers accumulate a high concentration of alkaloids, which might potentially compromise food safety because of their toxicity (Friedman,
For example, the accumulation of glycoalkaloids can inhibit acetylcholinesterase activity and damage stomach cell membranes and has been associated with systemic side effects and gastrointestinal disturbance in people (Yamashoji and Matsuda, 2013). Alkaloid intake in small quantities can lead to gastrointestinal symptoms, such as abdominal pain, vomiting, and diarrhea, whereas higher alkaloid doses can cause acute poisoning or severe symptoms, such as coma, neurological disorders, cardiac failure, and paralysis (Mensinga et al., 2005). Our results showed that ClO$_2$ could improve the antioxidant capacity and the safety of potato tubers by promoting the accumulation of flavonoids and inhibiting the synthesis of alkaloids, respectively (Deng et al., 2020; Wang et al., 2020).

Integrated metabolomic and RNA-seq were performed to determine the correlation between identified DEGs and DAMs. The results showed that different metabolites were regulated by different genes. For example, 4-methylumbelliferyl acetate and methoxyindoleacetic acid were regulated by nine genes, with seven and two genes as positive and negative regulators, respectively. The results suggest a complex regulatory mechanism between the changes in metabolite accumulation and gene expression abundance in potato tubers. Thus, we speculated that the mechanism of ClO$_2$-repressed sprouting is achieved by a coordinated complex process involving several metabolic pathways, which inhibit signal transduction, plant hormones, and alkaloid synthesis in potato tubers, leading to repressed sprouting. The activity of ClO$_2$ thus differs from other sprout inhibitors, such as chlorpropham, which is a mitotic inhibitor used to control tuber sprouting during storage (Vaughn and Lehnen, 1991). Essential oils instead act by damaging sprouts but do not really prevent their sprouting (Shukla et al., 2019). However, ClO$_2$ is a safer and inexpensive sprout inhibitor compared to chlorpropham and essential oils, which are environmentally harmful and expensive (Fu et al., 2019). This study provides a basis for understanding the mechanism of ClO$_2$-repressed sprouting; however, its specific regulatory roles still need further clarification.

CONCLUSION

This study used ClO$_2$-treated and control tubers to analyze the dynamic changes in metabolite accumulation and gene expression levels using UHPLC-MS and RNA-seq methods. The results revealed a total of 3,119 DEGs between the treated and control tubers. Most DEGs were enriched in phenylpropane metabolism, the MAPK signal pathway, and plant signal transduction pathways. ClO$_2$ could inhibit potato tuber sprouting by altering plant signal transduction. Metabolomic analyses identified 85 differential metabolites, with flavonoids and coumarins as the most upregulated differential metabolites, and alkaloids and amino acids and their derivatives as the downregulated differential metabolites. The ClO$_2$-mediated suppression of potato tuber sprouting is controlled by a complex regulatory network involving several different genes and metabolites.

DATA AVAILABILITY STATEMENT

The authors acknowledge that the data presented in this study must be deposited and made publicly available in an acceptable repository, prior to publication. Frontiers cannot accept an article that does not adhere to our open data policies.

AUTHOR CONTRIBUTIONS

XZ contributed to writing the original draft and writing, reviewing, and editing the manuscript. ML contributed to project administration and funding acquisition. ST contributed to supervision. SL contributed to formal analysis. JC contributed to project administration. XZ investigated the study. XW contributed to methodology. XG contributed to methodology software. JT contributed to data curation. YM contributed to resources. JS conceptualized the study. 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.2022.887179/full#supplementary-material

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