Comparative Transcriptome Analysis Provides Insights Into the Mechanism by Which 2,4-Dichlorophenoxyacetic Acid Improves Thermotolerance in Lentinula edodes

Ruiping Xu, Shasha Zhou, Jiaxin Song, Haiying Zhong, Tianwen Zhu, Yuhua Gong, Yan Zhou and Yinbing Bian*

Institute of Applied Mycology, College of Plant Science and Technology, Huazhong Agricultural University, Wuhan, China

As the widest cultivated edible mushroom worldwide, Lentinula edodes suffers serious yield and quality losses from heat stress during growth and development, and in our previous study, exogenous 2,4-Dichlorophenoxyacetic acid (2,4-D) was found to improve the thermotolerance of L. edodes strain YS3357, but the molecular mechanism remains unclear. Here, we explored the potential protective mechanism of exogenous 2,4-D against heat stress by transcriptome analysis. 2,4-D possible improve the thermotolerance of L. edodes through regulating antioxidant genes, transcription factors, energy-provision system, membrane fluidity, and cell wall remodeling. Furthermore, 2,4-D was also found to regulate the saturation levels of fatty acids and ATP content in L. edodes mycelium under heat stress. This study proposed a regulatory network of 2,4-D in regulating L. edodes response to heat stress, providing a theoretical basis for improving L. edodes thermotolerance, and facilitating the understanding of the molecular mechanism of exogenous hormones in alleviating abiotic stress damage to macrofungi.

Keywords: comparative transcriptome, Lentinula edodes, 2,4-dichlorophenoxyacetic acid, heat stress, fatty acid metabolism

INTRODUCTION

As the most widely cultivated edible mushroom with the highest output in the world (Royse et al., 2017), Lentinula edodes provide valuable benefits for the biotransformation of agricultural residues, but its production can be significantly affected by thermal stress. Previous studies have reported that the fruiting bodies of Agaricus bisporus were obviously reduced and became smaller after exposure to high temperatures for a few days. Heat stress not only affects the growth rate of fungal mycelium but also causes changes in the cell wall integrity and the increase of membrane fluidity (Liu et al., 2017; Qiu et al., 2018). Additionally, high temperature causes changes in the extracellular metabolites of Pleurotus ostreatus to promote Trichoderma asperellum growth. Furthermore, high temperatures can cause the rotting of logs or bags of...
L. edodes infected with Trichoderma spp., leading to yield and quality losses (Cao et al., 2015; Wang et al., 2016). All these reports suggest that improving the thermotolerance of L. edodes will help the mushroom cultivation industry.

Previous studies have shown that Para-aminobenzoic acid (PABA) synthase, nitric oxide, and trehalose can reduce the accumulation of reactive oxygen species (ROS) to alleviate oxidative damage induced by heat stress, thereby improving the thermotolerance of a variety of macrofungi (Kong et al., 2012; Yin et al., 2015; Luo et al., 2021). In our previous studies, LetrpE, LeYUCCA8, LeDnaJ07, auxin, and 2,4-Dichlorophenoxycetic acid (2,4-D) were found to participate in the response of L. edodes to heat stress (Zhou et al., 2018a; Wang et al., 2018b, 2020; Zhong et al., 2021). 2,4-D, a functional analogue of the auxin indole-3-acetic acid (IAA) is widely used as an IAA substitute because it can induce IAA like activities (Maher and Martindale, 1980). Meanwhile, 2,4-D is more stable than IAA under blue and ultraviolet light (Stasinopoulos and Hangarter, 1990), making it more compatible than IAA as an exogenous auxin. In previous studies, researchers have tried to improve the thermotolerance of crops by using auxins, such as IAA, 2,4-D, or the 1-naphthaleneacetic acid (NAA; Sakata et al., 2010; Ishida et al., 2016). In our previous work, exogenous 2,4-D was shown to improve the thermotolerance of L. edodes strain YS3357 by increasing the mycelial antioxidant enzyme activity and reducing the cellular lipid peroxidation degree (Zhou et al., 2018b), but the molecular mechanism remains to be elucidated.

The purpose of this study was to explore the molecular mechanism by which 2,4-D improves the thermotolerance of L. edodes by using 0.01 mM 2,4-D to pre-treat YS3357 strain, and the cDNA library was constructed and sequenced according to the manufacturer’s instructions (Illumina, San Diego, CA, United States). The 12 cDNA libraries were sequenced at Beijing Genomics Institute (BGI)-Shenzhen (Wuhan, China), with three independent biological replicates of CK_NH, CK_HS, 24D_NH, and 24D_HS to sequence the YS3357 samples. No less than 3GB of clean data was obtained for each sample, and all the RNA-seq raw data were uploaded to NCBI SRA with the BioProject number: PRJNA813710.

Data pre-treated and quality control, mapping the RNA-seq reads to the L. edodes genome, gene expression levels calculation, DEGs screening, and transcription factors (TFs) identifying were all conducted following the description of our previous study (Wang et al., 2018b). 12 raw datasets of all samples were pre-treated by Trimmomatic. 2,4-D was observed to modify the saturation levels of fatty acids and ATP contents in L. edodes mycelium response to heat stress. This study not only explores the potential molecular mechanism of 2,4-D in regulating L. edodes response to heat stress but also identifies the saturation levels of fatty acids as the response of metabolites in this process.

MATERIALS AND METHODS

Sample Preparation

The sawdust medium (78 g of hardwood sawdust, 20 g of wheat bran, 2 g of lime and 20 g of agar in 11 of distilled water) was prepared as the basic medium. Six plates were added with a final concentration of 0.01 mmol/l 2,4-D (group of 24D_), and the other six were added with equal methanol (group of CK_). L. edodes wild strain YS3357 was cultivated for 9 days at 25°C in the two groups of medium. Then, three plates (means three repeats) of both groups were treated at 40°C for 24h (divided into groups of CK_HS and 24D_HS), and the remaining three plates of both groups continued to grow at 25°C for 24h (divided into groups of CK_NH and 24D_NH). After cultivation, all the samples were frozen in liquid nitrogen and used for RNA extraction.

cDNA Library Construction, RNA Sequencing, and Transcriptome Data Processing

Total RNA was extracted using the RNAiso Plus method (TAKARA, Shanghai, China), and the cDNA library was constructed and sequenced according to the manufacturer's instructions (Illumina, San Diego, CA, United States). The 12 cDNA libraries were sequenced at Beijing Genomics Institute (BGI)-Shenzhen (Wuhan, China), with three independent biological replicates of CK_NH, CK_HS, 24D_NH, and 24D_HS to sequence the YS3357 samples. No less than 3GB of clean data was obtained for each sample, and all the RNA-seq raw data were uploaded to NCBI SRA with the BioProject number: PRJNA813710.

qRT-PCR Analysis

To validate the transcriptome sequencing results, nine genes were randomly selected for qRT-PCR analysis using AceQqPCR SYBR Master SYBR (Vazyme, Nanjing, China), with the actin gene as the internal control. The results were analyzed
using the CFX Connect Real-Time PCR system (BIO-RAD), and the relative expression was calculated using the 2^(-ΔΔCT) method (Livak and Schmittgen, 2002).

**Fatty Acid Extraction and Analysis**

Fatty acid (FA) extraction followed a previously method (Welti et al., 2002). Fatty acid methyl esters (FAMEs) were analyzed by gas chromatography (GC, Beckman Coulter) and identified by their retention times relative to the mix standard of 37 components of FAMEs. The chromatographic peak area was measured as the relative content of the FAMEs.

**Determination of ATP Content**

The ATP content of each sample was measured using an ATP assay kit (Solarbio, Beijing, China) as instructed by the manufacturer. The ATP extract from the kit was used to lyse the mycelium, followed by centrifugation at 8000 g for 10 min at 4°C, transferring supernatant to another EP tube, adding 500 μl of chloroform and well shaking, centrifugation at 10000 g for 3 min at 4°C, collecting the supernatant to determine the ATP content for different treatment groups.

**Statistical Analysis**

Data were expressed as mean and standard deviation (SD). The significant differences between analyzed samples were determined by one-way ANOVA and evaluated through Duncan’s multiple range test at \( p < 0.05 \).

**RESULTS**

**Transcriptome Analysis of the Response of DEGs to Heat Stress Under 2,4-D Treatment**

In our previous study, 2,4-D was found to improve the thermotolerance of the *L. edodes* strain YS3357 (Figure 1A; Zhou et al., 2018a). To obtain an overview of the *L. edodes* transcriptomic responses to 2,4-D treatment under heat stress, CK_NH, CK_HS, 24D_NH, and 24D_HS samples with 12 RNA-Seq cDNA libraries, obtaining 20,566,345, 19,509,383, 20,336,913 and 20,310,599 150-bp paired-end reads for the four groups of samples, respectively, with Q30 > 94.93% for clean reads. Using the splice-aware aligner TopHat2, about 60% of the reads in YS3357 were mapped to the *L. edodes* (W1-26) genome sequence (Supplementary Table S1), and the matching rate was related to the genetic characteristics of different strains: W1 (W1-26 parent) is a cultured strain, while YS3357 is a wild-type strain. The Pearson correlation coefficients of different replicates per group were calculated and their values were over 86% (YS3357) in each sample (Supplementary Figure S1), indicating that the sequencing results were reliable and could be used for subsequent analysis.

For better understanding of the biological mechanism of 2,4-D in enhancing the thermotolerance of *L. edodes* mycelium, we analyzed the DEGs in CK_HS/CK_NH and 24D_HS/24D_NH, with 1,281 and 2,363 DEGs (more than 2-fold or less than 0.5-fold) identified in the two groups, respectively. The number of DEGs between the two groups were shown in Figures 1B,C. These DEGs were considered as important candidate genes response to the heat stress by 2,4-D and used for further analysis.

The up-regulated and down-regulated DEGs in CK_HS/CK_NH and 24D_HS/24D_NH were classified by GO term annotation. The up-regulated DEGs in the two groups showed the top three enriched terms, the categories of cellular component (catalytic activity, cell part, and organelle), molecular function (binding, transcription regulator activity, and transporter activity), and biological process (metabolic process, cellular process, and localization; Figure 1D). Among the down-regulated DEGs in the two groups, the top three enriched terms included cellular component (catalytic activity, cell part, and cell), biological process (metabolic process, response to stimulus, cellular process, and biological regulation), and molecular function (binding, transcription regulator activity, and transporter activity; Figure 1E).

Based on the GO annotation of DEGs, the up-regulated genes of 24D_HS/24D_NH showed unique functions in supramolecular complex, multicellular organismal process, obsolete GTP catabolic process, obsolete ATP-dependent proteolysis, and carbon utilization (Supplementary Table S2), while downregulated genes showed unique functions in the other organism, other organism part, obsolete gamma-glutamyltransferase activity, and obsolete electron transport (Supplementary Table S3).

The functions of 2,4-D in enhancing the thermotolerance of *L. edodes* were further investigated by KEGG pathway analysis induced by 2,4-D and heat stress exposure. Between CK_HS/CK_NH, 762 DEGs were enriched in 107 KEGG pathways, and between 24D_HS/24D_NH, 1876 DEGs were enriched in 115 KEGG pathways. KEGG pathway enrichment analysis revealed that 24D_HS/24D_NH was different from CK_HS/CK_NH in most of the enrichment pathways, such as tryptophan metabolism, oxidative phosphorylation, biosynthesis of unsaturated fatty acids, FA metabolism, non-homologous end-joining, ribosome, RNA transport, and the degradation of valine, leucine and isoleucine (Figures 1F,G). These different enrichment pathways were considered as essential for improving mycelial thermotolerance.

The RNA-seq data were confirmed by qRT-PCR analysis of the expression of nine randomly selected genes (Supplementary Table S4). The information for each gene and primer is shown in Supplementary Table S4. Correlation analysis was performed on gene folding changes between the two treatment groups (RNA-seq and qRT-PCR). The qRT-PCR data were seen to agree with the RNA-seq data, with a significant positive correlation \( (R^2 = 0.8918) \), indicating the reliability of the RNA-seq data (Supplementary Figure S2A). We selected four genes with significant difference expression at 24D_HS/24D_NH for the figure, which were relatively consistent with the transcriptome data (Supplementary Figure S2B).

**2,4-D Regulated the Expression of Antioxidant Genes in Response to Heat Stress**

Oxidative stress was reported directly induce cell damage after heat stress (Luo et al., 2021). Here, transcriptome analysis
showed that the expressions of genes encoding ROS scavenging enzymes were significantly changed by exogenous addition of 2,4-D under heat stress, and we identified the genes encoding such enzymes, including catalase (CAT), glutathione-S transferase (GST), ascorbate peroxidase (APX), lipoxygenase (LOX), and thioredoxin (TRX).

Among them, one APX gene (LE01Gene03530), two GST genes (LE01Gene11416 and LE01Gene05851), and one TRX gene (LE01Gene03334) were not differentially expressed between CK_HS/CK_NH but down-regulated by 2,4-D under heat stress. Additionally, the following genes were not differentially expressed between CK_HS/CK_NH but up-regulated between 24D_HS/24D_NH, such as two APX genes (LE01Gene08720 and LE01Gene08723), one GST gene (LE01Gene10661), one CAT gene (LE01Gene04245), one LOX gene (LE01Gene08979), and five TRX genes (LE01Gene02450, LE01Gene04578, LE01Gene10661, LE01Gene12376, LE01Gene12821; Figure 2A). These results suggested that 2,4-D may improve *L. edodes* thermotolerance by regulating the expression of antioxidant genes.

2,4-D Regulated Oxidative Phosphorylation in Response to Heat Stress

As previously reported, mitochondria are important organelles for ATP production in eukaryotic cells and oxidative phosphorylation-dependent energy conversion, including the respiratory chain and ATP synthase (Suhm and Ott, 2016). Moreover, oxidative phosphorylation plays a very important
role in cellular stress response (Jin et al., 2021). KEGG analysis revealed the enrichment of oxidative phosphorylation in the 24D_ HS/24D_ NH group, and analyzing the expression of genes related to oxidative phosphorylation. A total of 18 oxidative phosphorylation-related genes were found to be differentially expressed under 2,4-D exposure and heat stress (Figure 2B), with only two genes (LE01Gene00401 and LE01Gene04661) were down-regulated in 24D_ HS/24D_ NH but not differentially expressed in CK_HS/CK_NH, 16 genes up-regulated in 24D_ HS/24D_ NH but not differentially expressed in CK_HS/CK_NH. Furthermore, we measured the ATP content of CK_NH, CK_HS, 24D_NH, and 24D_HS, and the ATP content was found to increase significantly under heat stress, but this increase was significantly inhibited by exogenous 2,4-D (Supplementary Figure S3). These results indicated that 2,4-D may improve L. edodes thermotolerance by regulating oxidative phosphorylation-related genes and ATP production.

2,4-D Regulated IAA Biosynthesis Pathway in Response to Heat Stress

In the present study, the effects of 2,4-D on the endogenous auxin biosynthesis pathway in improving L. edodes thermotolerance were evaluated by analyzing the expression of genes related to auxin biosynthesis (Shellet, 2008). Interestingly, 30 TFs were differentially expressed between CK_HS/CK_NH and 24D_ HS/24D_ NH, including bZIP, zf-C2H2, zf-clus, Fungal_trans, HMG_box, and other TF family members (Figure 2C). These TFs were not induced by heat stress, but by 2,4-D under heat stress, may participate in L. edodes thermotolerance.

2,4-D Regulated Genes Involved in TCA Cycle and Glycolysis in Response to Heat Stress

KEGG enrichment analysis showed that TCA cycle-related genes were related to 2,4-D response under heat stress, and further analysis revealed that 10 genes were differentially expressed in 24D_ HS/24D_ NH, but not differentially expressed in CK_HS/CK_NH, including nine up-regulated genes (LE01Gene01135, LE01Gene03404, LE01Gene04224, LE01Gene07959, LE01Gene08443, LE01Gene10348, LE01Gene10979, LE01Gene11013, LE01Gene11377), and one down-regulated gene (LE01Gene04661; Figure 3).
Increasing the expression of these genes accelerates the TCA cycle and generates energy, notably phosphoenolpyruvate carboxykinase (PEPK), which produces oxaloacetate from pyruvate to sustain the TCA cycle.

Pyruvate is a key metabolite linking glycolysis and the TCA cycle (Fernie et al., 2004), and the enzyme genes in the glycolytic pathway from glucose to pyruvate were not differentially expressed in CK_HS/CK_NH and 24D_HS/24D_NH. Under aerobic respiration conditions, pyruvate can be converted to acetyl CoA through pyruvate dehydrogenase, while under anaerobic respiration, pyruvate can produce lactate or ethanol through lactate dehydrogenase and ethanol dehydrogenase, respectively (Sakihama et al., 2019). In the transcriptome results, we found four alcohol dehydrogenase (ADH) genes (LE01Gene06191, LE01Gene1430, LE01Gene12440, LE01Gene12441) up-regulated between 24D_HS/24D_NH but not differentially expressed between CK_HS/CK_NH (Figure 3), with the lactate dehydrogenase genes being not differentially expressed in either 24D_HS/24D_NH or CK_HS/CK_NH, leading to a possible reduction in the acetaldehyde contents. We also found the pyruvate dehydrogenase (PDH) genes were not differentially expressed between CK_HS/CK_NH but were up-regulated between 24D_HS/24D_NH (Figure 3).

Citrate synthase (CS) is the key rate-limiting enzyme in the TCA cycle (Li et al., 2018b), and we observed that citrate synthase genes (LE01Gene01135 and LE01Gene10348) were up-regulated in 24D_HS/24D_NH, but not differentially expressed in CK_HS/CK_NH (Figure 3). These results suggested that 2,4-D may improve L. edodes' thermotolerance by regulating the key genes related to the TCA cycle.

2,4-D Regulated the Fatty Acid Saturation Level in Response to Heat Stress

In the KEGG pathway analysis, we found the enrichment of the biosynthesis of unsaturated FAs and FA metabolism in 24D_HS/24D_NH, and Supplementary Table S5 shows the changes in transcript levels of key genes in CK_HS/CK_NH.

![Figure 3](https://biorender.com)
and 24D_ HS/24D_ NH. Peroxisome plays an essential part in cellular metabolism, including FA oxidation (Van Roermund et al., 2003). In this study, we found that seven peroxisome-related genes were not differentially expressed between CK_HS/CK_NH, but were up-regulated between 24D_ HS/24D_ NH (Supplementary Table S6), suggesting that 2,4-D may regulate the FA metabolism under heat stress.

At the same time, in order to explore whether the FA content of *L. edodes* in response to 2,4-D under heat stress, we measured the FA content in CK_NH, 24D_NH, CK_HS, and 24D_HS by GC–MS. The results showed that the FA saturation degree changed significantly in 24D_HS versus CK_HS (Figure 4A). Interestingly, compared with CK_NH, 24D_NH showed a significant decrease in the FA saturation degree, suggesting that the addition of 2,4-D affected the saturation degree of FA, while CK_HS showed no significant change compared to CK_NH (Figure 4B). The results indicate that 2,4-D can regulate the FA saturation degree under heat stress, and has a potential relationship with *L. edodes* thermotolerance.

**Co-expression Network Analysis Obtain Key Genes in 24D_HS**

The soft threshold was screened as 14, and the genes with different expression patterns were clustered into 11 modules and distinguished by different color blocks (Supplementary Figure S5A). The module with the highest correlation to the 24D_HS was MEbrown (Supplementary Figure S5B). A total of 877 genes were clustered in MEbrown module, with 141 genes in the co-expression network. The top three hub genes, co-expression networks, and GO terms are presented in Supplementary Figures S5C,D.

The top three hub genes (LE01Gene01937, LE01Gene05096, and LE01Gene05680) encode sulfate adenylyltransferase, condensin complex, and ubiquinol-cytochrome C reductase core protein 2, respectively. GO term analysis revealed that the 141 genes in the co-expression network were mainly enriched in cell part, organelle, protein-containing complex, and membrane part in the term of cellular component; binding and catalytic activity in the term of molecular function; metabolic process, cellular process, and localization in the term of biological process (Supplementary Figures S5C,D).

**DISCUSSION**

**2,4-D Affects Antioxidant Genes in *Lentinula edodes* Heat Stress**

Heat stress was reported to cause excessive ROS accumulation, DNA damage, protein denaturation, damage to lipid membrane permeability, and changes in cell wall integrity (Li et al., 2018a; Lippmann et al., 2019). High temperature could increase the auxin content in plants, so auxin plays a significant role in regulating plant adaptation to high temperature through auxin biosynthesis, signaling, and transport pathways (Franklin et al., 2011). As reported previously, the application of IAA, NAA, or 2,4-D could completely reverse male sterility in barley and *Arabidopsis* (Sakata et al., 2010). The pre-treatment of pea plants with auxins TA-12 and TA-14 could mitigate the oxidative stress provoked by high-temperature treatment (Sergiev et al., 2018). This is consistent with our previous results: (1) exogenous addition of IAA could significantly increase the thermotolerance of heat-sensitive strain YS3357, with a significant increase in the IAA content of mycelium under heat stress, and higher endogenous IAA content in the heat-tolerant strain than the heat-sensitive strain (Wang et al., 2018b, 2020); (2) exogenous 2,4-D could improve the thermotolerance of *L. edodes* strain YS3357 by increasing mycelial antioxidant enzyme activity and reducing cellular lipid peroxidation degree (Zhou et al., 2018a).

In this study, 2,4-D was found to regulate the expression of antioxidant genes to alleviate *L. edodes* heat stress. Based on RNA-seq data, exogenous application of 2,4-D can change the expression of Trx genes under heat stress. TRX is an...
important functional protein involved in scavenging intracellular ROS toxicity. Thioredoxin is a biomarker for oxidative stress, and many studies have shown the involvement of thioredoxins in a variety of redox-dependent cellular processes, such as gene expression, signal transduction, cell growth, and apoptosis (Matsuo and Yodoi, 2013). Trx2 was reported to be essential for maintaining the redox status when Schizosaccharomyces pombe was exposed to starvation and mild-heat stresses, indicating that cytoplasmic thioredoxins have an important role in adaptation to environmental change (Yano et al., 2010; Oku et al., 2013). These results indicated that 2,4-D may improve L. edodes thermotolerance through anti-oxidation reactions.

Collectively, exogenous 2,4-D may regulate the expression of CAT, GST, APX, LOX, and TRX genes to enhance the detoxification effect of ROS and reduce cell damage and cell death. As an antioxidant defense system, 2,4-D may improve the thermotolerance of L. edodes by scavenging active oxygen.

2,4-D Regulates Energy-Provision System and Fatty Acid Metabolism in Lentinula edodes Under Heat Stress

Stresses cause high energy consumption (De Block et al., 2005), leading to cell damage and energy imbalance under prolonged heat stress. The typical energy-provision pathways such as glycolysis, TCA cycle, and oxidative phosphorylation are cellular energy metabolic pathways, providing ATP for plant growth or defense (Fernie et al., 2004; Chen and Russo, 2012). As previously reported, heat stress could down-regulate energy metabolisms, decreasing the intermediate product of glycolysis and TCA cycle in the response of L. edodes to heat stress (Zhao et al., 2019). Exogenous L-Arginine was reported to enrich the DEGs involved in glycolysis, TCA cycle, and oxidative phosphorylation in Gracilariapis lemianiformis under heat stress (Zhang et al., 2021b). In the present study, 2,4-D could remarkably enhance the energy metabolism by activating the expression of genes encoding the TCA cycle (e.g., pyruvate dehydrogenase, citrate synthase, phosphoenolpyruvate carboxykinase, and fumarate reductase) and electron transport chain NADH dehydrogenase, cytochrome c reductase, cytochrome c oxidase, and ATPase (Figures 2B, 3) under thermal stress, it can provide energy for subsequent mycelial regeneration. Additionally, ATP content analysis showed that there may be a potential correlation between ATP content and 2,4-D to alleviate mycelial heat stress.

Pyruvate can be metabolized to lactate and ethanol in the glycolytic pathway, resulting in the accumulation of acetaldehyde as a DNA-damaging metabolite in the pathway to ethanol (Noguchi et al., 2016). In this study, we found that pyruvate decarboxylase and ethanol dehydrogenase were up-regulated while lactate dehydrogenase remained unchanged under 2,4-D exposure heat stress, which may have led to a reduction in acetaldehyde content, thus reducing cellular toxicity under heat stress. Meanwhile, pyruvate metabolism produces less energy during anaerobic respiration, but this energy is crucial for cell survival. The expression of these genes induced by 2,4-D at heat stress may play an important role in the recovery of L. edodes mycelium from thermal damage.

Citrate synthase, a central enzyme in carbon metabolism, is essential in the TCA cycle, amino acid synthesis, and the glyoxalate cycle. As a first step of the TCA cycle, citrate synthase catalyzes the condensation of oxaloacetate and acetyl-CoA to form citric acid (Ren et al., 2020). Citric acid could improve thermotolerance in P. ostreatus mycelium, nitric oxide (NO) inhibiting the expression of aconitase gene, resulting in citric acid accumulation under heat stress, inducing the expression of Aox gene to activate the alternative oxidative pathway (Hou et al., 2021). Based on transcriptome data analysis, under heat stress, 2,4-D could induce the up-regulated expression of two citrate synthase genes and Aox gene agreeing with NO alleviating heat stress damage in P. ostreatus mycelia, inferring that 2,4-D may regulate energy metabolism to improve the thermotolerance of L. edodes.

Moreover, citric acid is defined as a key precursor in lipid accumulation (Magoulis et al., 2020). In our study, three delta 12 desaturase genes showed up-regulated expression under 2,4-D exposure heat stress. Interestingly, we measured the FA content and found an increase in the FA saturation degree at 24D_HS, it probably due to variations in FA metabolism. As the key genes for FA β-oxidation, 2,4-D under heat stress induced the up-regulated expression of ketoacyl-CoA thiolaic (LE01Gene02324), acylcarbamate carrier protein (LE01Gene12299), and hydroxyacyl-CoA dehydrogenase (LE01Gene07809; Supplementary Table S5). The saturation level of fatty acids may be an important factor in controlling plant thermotolerance (Larkindale and Huang, 2004; Zhang et al., 2021a), which was consistent with our results. Additionally, the expression of peroxisome related genes was up-regulated, promoting FA β-oxidation while producing a large amounts of acetyl-CoA, contributing to the TCA cycle. Our FA data suggested that the saturation levels of fatty acids were regulated by 2,4-D under heat stress, it could be a potential factor for the L. edodes mycelium to improve thermotolerance.

2,4-D Serves as a Regulator of Transcription Factors Involved in Heat Stress

Transcription factors play an important role in controlling various aspects of fungal metabolism, development, stress tolerance, members of the Zn2Cys6, C2H2, GATA, bZIP, and heat shock factor transcription factor (HSF) families were involved in abiotic stress response (John et al., 2021). HSFs with the HSF structural domain are regulators of the fungal heat shock protein (HSP) genes, to which the structural domain binds in response to heat shock and other stresses (Zhou et al., 2018a). In the present study, we found nine TF families between CK_HS/CK_NH and 24D_HS/24D_NH, with a total of 30 genes differentially expressed, including 21 TFs not induced by heat stress, but differentially expressed under 2,4-D exposure heat stress, such as Fungal_trans, GATA, zf-C2H2, and zf-clus and other TFs (Figure 4). Notably, the expression of DEGs involved in HSF was down-regulated in 24D_HS/24D_NH, but with no differential expression in CK_HS/CK_NH, indicating the negative regulation of HSF in L. edodes mycelium thermotolerance, inferring that the regulation of TFs on HSF may contribute to 2,4-D induced thermotolerance in L. edodes.
**2,4-D May Improves *Lentinula edodes* Thermotolerance Through a Complex Network**

Heat stress has emerged as one of the most destructive abiotic stresses on crops (Glazebrook, 1999). For macrofungi, heat stress may affect the production of fruiting bodies and even cause significant yield decrease (Lu et al., 2014). The effects of heat stress on fungi mainly include mycelial growth, cell wall structure, and membrane fluidity (Luo et al., 2021). Many previous studies have revealed the important roles of auxin and auxin signaling in improving the abiotic stress of plants (Piotrowska-Niczyporuk et al., 2018; Sergiev et al., 2018; Wang et al., 2021). However, the signal transduction pathway of auxin in fungi remains unknown.

In this study, the molecular role of 2,4-D in enhancing the thermotolerance of *L. edodes* mycelium was demonstrated by a complex regulatory network in combination with our findings (Figure 5). KEGG pathway analysis revealed Ribosome and RNA transport pathways exclusively in 24D_HS/24D_NH, suggesting that the 2,4-D may improve protein synthesis and increases the ability of RNA transport under heat stress. In eukaryotes, the spatio-temporal articulation of gene expression is mainly determined by the intertwined pathways of RNA transport and local translation regulation (Kindler et al., 2005).

For ROS elimination, 2,4-D can perform the functions of regulating antioxidant genes in heat stress (CAT, GST, APX, LOX, TRX). Heat stress can induce an increase in the level of endogenous IAA in *L. edodes* mycelium (Wang et al., 2018a). 2,4-D can also regulating the expression of auxin biosynthesis pathway-related genes under heat stress, thereby affecting downstream genes regulated by auxin. 2,4-D inducing the expression of some transcription factors (bZIP, Zf-C2H2, Zf-clus, etc.) to resist heat stress. 2,4-D inducing the expression of key genes involved in the TCA cycle, glycolysis and oxidative phosphorylation (citrate synthase, pyruvate dehydrogenase, alcohol dehydrogenase, etc.) under heat stress, thus affecting ATP production and FA synthesis.

Heat stress can also disrupt the integrity of the cell wall and damage the cell membrane, thereby altering the function of the plasma membrane structure and leading to increased membrane fluidity and reduced permeability (Ashraf and Harris, 2004). However, exogenous 2,4-D can reduce the membrane fluidity of cells through increase FA saturation degree in response to heat damage under heat stress. Micro-tubules are cytoskeletal elements and the αβ tubulin heterodimer is a structural subunit of micro-tubules essential for intracellular transport and cell division in all eukaryotes (Nogales et al., 1998). In GO annotation, the up-regulated genes in unique GO term enrichment at 24D_HS/24D_NH included α-tubulin and β-tubulin (Supplementary Table S2), suggesting that tubulin is involved in 2,4-D induced thermotolerance of *L. edodes*. Fungal chitinases are hydrolytic enzymes responsible for the degradation of chitin, with chitinases and chitosanase playing an essential role in cell wall remodeling and cell wall synthesis, respectively (Seidl, 2008). In the transcriptome

![FIGURE 5](www.figdraw.com)
data, the two enzyme genes were both up-regulated in 24D_HS/24D_NH, suggesting 2,4-D may be involved in cell wall remodeling after heat stress (Supplementary Table S7).

For the three hub genes in the co-expression network in 24D_HS, sulfate adenylyltransferase (LE01Gene01937) was the key energy gene of four moderate thermophiles and catalyze adenosine phosphosulfate (APS) to generate ATP and sulfate, which is the final stage of the sulfate oxidation so as to obtain energy (Zhou et al., 2015); the function of condensin complexes (LE01Gene05096) is versatile in gene regulation and chromosome segregation (Csankovszki et al., 2009); ubiquinol-cytochrome C reductase core protein 2 (LE01Gene05680) is a key subunit of the mitochondrial electron transport chain complex III (Sun et al., 2003). Collectively, these genes in MEbrown module may contribute to the 2,4-D induced thermotolerance of L. edodes.

CONCLUSION

As an analogue of the endogenous auxin IAA, 2,4-D plays an important role in regulating plant responses to abiotic stresses. The content of endogenous IAA in L. edodes can affect its thermotolerance, and exogenous 2,4-D can affect the antioxidant enzyme activity thus the thermotolerance of L. edodes. In this study, we used exogenous 2,4-D to increase the thermotolerance of L. edodes and explored the potentially molecular mechanism of 2,4-D enhanced thermotolerance in L. edodes by transcriptome analysis. 2,4-D improve L. edodes’ thermotolerance possibly through regulating antioxidant genes, transcription factors, energy-provision system, membrane fluidity, cell wall remodeling, and other mechanisms. Future research can focus on functional analysis of the key genes through over-expression and gene editing. This study development of an potential molecular mechanism pathway to improve L. edodes thermotolerance through 2,4-D regulation and has also genetic basis for improving its thermotolerance, meanwhile, the results provided a reference for the research of auxin improving abiotic stress of fungi.

DATA AVAILABILITY STATEMENT

The original contributions presented in the study are publicly available. This data can be found at NCBI SRA (accession number: PRJNA81371).

AUTHOR CONTRIBUTIONS

YB, YZ, and YG conceived and designed the experiments. RX and SZ performed the experiments and wrote the paper. JS, HZ, and TZ contributed to modify and polish the paper. All authors contributed to the article and approved the submitted version.

FUNDING

This research was funded by the National Natural Science Foundation of China (grant no. 31972476).

SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2022.910255/full#supplementary-material
Nogales, E., Wolf, S. G., and Downing, K. H. (1998). Structure of the alpha beta tubulin dimer by electron crystallography. *Nature* 391, 199–203. doi:10.1038/34465
Noguchi, C., Grothusen, G., Anandarajan, V., Martinez-Lage Garcia, M., Terlecky, D., Corzo, K., et al. (2016). Genetic controls of DNA damage avoidance in response to acetaldehyde in fission yeast. Cell Cycle 16, 45–58. doi:10.1080/15384101.2016.123732
Oku, M., Hoseki, J., Ichiki, Y., and Sakai, Y. (2013). A fluorescence resonance energy transfer (FRET)-based redox sensor reveals physiological role of thioredoxin in the yeast *Saccharomyces cerevisiae*. FEBS Lett. 587, 793–798. doi:10.1016/j.febslet.2013.02.003
Piotrowska-Niczyporuk, A., Bajguz, A., Zambryszcka-Szlewa, E., and Bralska, M. (2018). Exogenously applied auxins and cytokinins ameliorate lead toxicity by inducing antioxidant defence system in green alga *Acutodesmus obliquus*. Plant Physiol. Biochem. 132, 535–546. doi:10.1016/j.plaphy.2018.09.038
Qu, Z., Wu, X., Gao, W., Zhang, J., and Huang, C. (2018). High temperature induced disruption of the cell wall integrity and structure in *Pleurotus ostreatus* mycelia. *Appl. Microbiol. Biotechnol.* 102, 6627–6636. doi:10.1007/s00253-018-9090-6
Ren, M., Yang, X., Bie, J., Wang, Z., Liu, M., Li, Y., et al. (2020). Citrate synthase desuccinylation by SIRT5 promotes colon cancer cell proliferation and migration. *Biomed. Res. Int.* 2020, 1018. doi:10.1155/2020/1018
Rouse, D. J., Baars, J., and Tan, Q. (2017). “Current overview of mushroom production in the world” in *Edible and Medicinal Mushrooms: Technology and Applications*. Eds. C. Tedesco and A. Parodi-Giménez (United States: John Wiley and Sons), 5–13.
Sakata, T., Oshino, T., Miura, S., Tomabechi, M., Tsunaga, Y., Higashitani, N., et al. (2010). Auxins reverse plant male sterility caused by high temperatures. *Proc. Natl. Acad. Sci. U. S. A.* 107, 8569–8574. doi:10.1073/pnas.0906810107
Sakihama, Y., Hidese, R., Hasunuma, T., and Kondo, A. (2019). Increased flux in acetyl-CoA synthetic pathway and TCA cycle of *Klyveromyces marxianus* under respiratory conditions. *Sci. Rep.* 9:5319. doi:10.1038/s41598-019-41863-3
Seidl, V. (2008). Chitinases of filamentous fungi: a large group of diverse proteins with multiple physiological functions. *Fungal Biol. Rev.* 22, 36–42. doi:10.1016/j.fbr.2008.03.002
Sergiev, I., Todorova, D., Shopova, E., Jankauskienė, J., Jankovska-Bortkevič, E., and Jurkoniene, S. (2018). Effects of auxin analogues and heat stress on garden pea. *Zemědělský listy* 105, 243–248. doi:10.1080/12143757.2018.1150531
Shelест, E. (2008). Transcription factors in fungi. *FEBS Microbiol. Lett.* 286, 145–151. doi:10.1111/j.1574-6968.2008.01293.x
Stasinopoulos, T. C., and Hangarter, R. P. (1996). Preventing photochemistry in culture media by Long-pass light filters alters growth of cultured tissues. *Plant Physiol.* 93, 1365–1369. doi:10.1104/pp.93.4.1365
Suhin, T., and Ott, M. (2016). Mitochondrial translation and cellular stress response. *Cell Tissue Res.* 367, 21–31. doi:10.1007/s00441-016-2460-4
Sun, G., Knitter, M. T., and Anderson, V. E. (2003). Mass spectrometric characterization of mitochondrial electron transport complex subunits of the rat heart ubiquinol-cytochrome c reductase. *J. Mass Spectrom.* 38, 531–539. doi:10.1002/jms.667
Van Roermund, C. W. T., Waterham, H. R., Jilst, L., and Wanders, R. (2003). Fatty acid metabolism in *Saccharomyces cerevisiae*. *Cell. Mol. Life Sci.* 60, 1838–1851. doi:10.1007/s00018-003-3076-x
Wang, G., Cao, X., Ma, X., Guo, M., Liu, C., Yan, L., et al. (2016). Diversity and effect of *Trichoderma* spp. associated with green mold disease on *Lentinula edodes* in China. *Microbiology 5*, 709–718. doi:10.1002/mbe.3364
Wang, Q., Guo, M., Xu, R., Zhang, J., Bian, Y., and Xiao, Y. (2019). Transcriptional changes on blight fruiting body of *Flammulina velutipes* caused by two new bacterial pathogens. *Front. Microbiol.* 10:2845. doi:10.3389/fmicb.2019.02845
Wang, G., Luo, Y., Wang, C., Zhou, Y., Mou, C., Kang, H., et al. (2020). Hsp40 protein LeDnaJ07 enhances the thermotolerance of *Pleurotus ostreatus* mycelia. *Appl. Microbiol. Biotechnol.* 102, 6627–6636. doi:10.1007/s00253-018-9090-6
Wang, G. Z., Ma, C. J., Luo, Y., Zhou, S. S., Zhou, Y., Ma, X. L., et al. (2018b). Citrate synthase isoform contributes infection and stress resistance of the stripe tuoliensis. *Plant Physiol.* 177, 145–151. doi:10.1093/txb/tbx200
Wang, F., Niu, H., Xin, D., Long, Y., Wang, G., Liu, Z., et al. (2021). OsIAA18, an aux/IAA transcription factor gene, is involved in salt and drought tolerance in *Oryza sativa* L. *Cell. Physiol. Biochem.* 100:408. doi:10.1159/000494784
Wang, Z., Luo, Y., Wang, C., Zhou, Y., Mou, C., Kang, H., et al. (2020). Hsp40 protein LeDnaJ07 enhances the thermotolerance of *Pleurotus ostreatus* mycelia. *Appl. Microbiol. Biotechnol.* 102, 6627–6636. doi:10.1007/s00253-018-9090-6
Wang, Z., Luo, Y., Zhou, S. S., Zhou, Y., Ma, X. L., et al. (2018b). Citrate synthase isoform contributes infection and stress resistance of the stripe tuoliensis. *Plant Physiol.* 177, 145–151. doi:10.1093/txb/tbx200
Wang, Z., Luo, Y., Zhou, S. S., Zhou, Y., Ma, X. L., et al. (2018b). Citrate synthase isoform contributes infection and stress resistance of the stripe tuoliensis. *Plant Physiol.* 177, 145–151. doi:10.1093/txb/tbx200
Wang, G., Zhou, S., Luo, Y., Ma, C., Gong, Y., Zhou, Y., et al. (2018a). The heat shock protein 40 LeDnaJ regulates stress resistance and indole-3-acetic acid biosynthesis in Lentinula edodes. *Fungal Genet. Biol.* 118, 37–44. doi: 10.1016/j.fgb.2018.07.002

Weih, R., Li, W., Li, M., Sang, Y., Biesiada, H., Zhou, H. E., et al. (2002). Profiling membrane lipids in plant stress responses. *J. Biol. Chem.* 277, 31994–32002. doi: 10.1074/jbc.M205375200

Yano, T., Oku, M., Akeyama, N., Itoyama, A., Yurimoto, H., Kuge, S., et al. (2010). A novel fluorescent sensor protein for visualization of redox states in the cytoplasm and in peroxisomes. *Mol. Cell. Biol.* 30, 3758–3766. doi: 10.1128/MCB.00121-10

Ye, J., Fang, L., Zheng, H., Zhang, Y., Chen, J., Zhang, Z., et al. (2006). WEGO: a web tool for plotting GO annotations. *Nucleic Acids Res.* 34, W293–W297. doi: 10.1093/nar/gkl031

Yin, C., Zheng, L., Zhu, J., Chen, L., and Ma, A. (2015). Enhancing stress tolerance by overexpression of a methionine sulfoxide reductase A (MrA) gene in Pleurotus ostreatus. *Appl. Microbiol. Biotechnol.* 99, 3115–3126. doi: 10.1007/s00253-014-6365-4

Zhang, J., Liu, S., Hu, C., Chen, X., Sun, X., and Xu, N. (2021b). Physiological and transcriptome analysis of exogenous l-arginine in the alleviation of high-temperature stress in Gracilaria lemaneiformis. *Front. Mar. Sci.* 8:784586. doi: 10.3389/fmars.2021.784586

Zhang, H., Zhang, Z., Xiong, Y., Shi, J., Chen, C., Pan, Y., et al. (2021a). Stearic acid desaturase gene negatively regulates the thermotolerance of Penicillium ternata by modifying the saturated levels of fatty acids. *Ind. Crop. Prod.* 166:113490. doi: 10.1016/j.indcrop.2021.113490

Zhao, X., Chen, M., Zhao, Y., Zha, L., Yang, H., and Wu, Y. (2019). GC(−)-MS-based non-targeted and targeted metabolic profiling identifies changes in the Lentinula edodes mycelial metabolome under high-temperature stress. *Int. J. Mol. Sci.* 20:2330. doi: 10.3390/ijms20092330

Zhong, H. Y., Luo, Y., Gong, Y. H., Xu, R. P., and Bian, Y. B. (2021). Functional analysis of Indole-3-pyruvate Monoxygenase gene YUCCA8 involved in heat resistance of Lentinula edodes by RNAi. *Acta Edulis Fungi* 28, 11–18. doi: 10.16488/j.cnki.1005-9873.2021.03.002

Zhou, D., Peng, T., Zhou, H., Liu, X., Gu, G., Chen, M., et al. (2015). Expression of critical sulfur- and iron-oxidation genes and the community dynamics during bioleaching of chalcopyrite concentrate by moderate thermophiles. *Curr. Microbiol.* 71, 62–69. doi: 10.1007/s00284-015-0817-7

Zhou, S. S., Wang, G. Z., Luo, Y., Ma, C. J., Gong, Y. H., and Bian, Y. B. (2018a). Auxin and auxin analogues enhancing the thermotolerance of Lentinula edodes. *Mycosystema* 37, 1723–1730. doi: 10.13346/j.mycosystema.180145

Zhou, G., Ying, S. H., Hu, Y., Fang, X., Feng, M. G., and Wang, J. (2018b). Roles of three hsf domain-containing proteins in mediating heat-shock protein genes and sustaining asexual cycle, stress tolerance, and virulence in Beauveria bassiana. *Front. Microbiol.* 9:1677. doi: 10.3389/fmicb.2018.01677

**Conflict of Interest:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

**Publisher's Note:** All claims expressed in this article are solely those of the authors and do not necessarily represent those of their affiliated organizations, or those of the publisher, the editors and the reviewers. Any product that may be evaluated in this article, or claim that may be made by its manufacturer, is not guaranteed or endorsed by the publisher.

**Copyright © 2022 Xu, Zhou, Song, Zhong, Zhu, Gong and Bian. This is an open-access article distributed under the terms of the Creative Commons Attribution License (CC BY). The use, distribution or reproduction in other forums is permitted, provided the original author(s) and the copyright owner(s) are credited and that the original publication in this journal is cited, in accordance with accepted academic practice. No use, distribution or reproduction is permitted which does not comply with these terms.**