Transcriptome analysis provides insights into the regulation of metabolic processes during postharvest cold storage of loquat (Eriobotrya japonica) fruit

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Abstract
Loquat (Eriobotrya japonica) fruit accumulates lignin during postharvest storage under chilling conditions (0 °C), while low-temperature conditioning (LTC; 5 °C for 6 days followed by transfer to 0 °C) or heat treatment (HT; 40 °C for 4 h followed by transfer to 0 °C) can alleviate lignification. Here we compared transcriptome profiles of loquat fruit samples under LTC or HT to those stored at 0 °C at five time points from day 1 to day 8 after treatment. High-throughput transcriptome sequences were de novo assembled into 53,319 unique transcripts with an N50 length of 1306 bp. A total of 2235 differentially expressed genes were identified in LTC, and 1020 were identified in HT compared to 0 °C. Key genes in the lignin biosynthetic pathway, including EjPAL2, EjCAD1, EjCAD3, 4CL, COMT, and HCT, were responsive to LTC or HT treatment, but they showed different expression patterns during the treatments, indicating that different structural genes could regulate lignification at different treatment stages. Coexpression network analysis showed that these candidate biosynthetic genes were associated with a number of transcription factors, including those belonging to the AP2, MYB, and NAC families. Gene ontology (GO) enrichment analysis of differentially expressed genes indicated that biological processes such as stress responses, cell wall and lignin metabolism, hormone metabolism, and metal ion transport were significantly affected under LTC or HT treatment when compared to 0 °C. Our analyses provide insights into transcriptome responses to postharvest treatments in loquat fruit.

Introduction
Fruit firmness is an important fruit quality trait. How to maintain optimal fruit firmness is consistently a topic of postharvest or horticultural research. For loquat fruit, lignification is induced during low-temperature (0 °C) storage, which significantly decreases its commodity value1,2. Treatments, such as low-temperature conditioning (LTC; 5 °C for 6 days followed by transfer to 0 °C) and heat treatment (HT; 40 °C for 4 h followed by transfer to 0 °C), have been developed to alleviate chilling-induced lignification. However, the underlying regulatory mechanisms of these treatments have rarely been investigated.

Lignin is the main component of plant secondary cell walls and plays an essential role in many different types of cells to provide structural support for growth, development, and defense1–3. The biosynthetic pathway of lignin has been well studied in plants and involves the synthesis of monolignols followed by construction of lignin polymers through oxidative polymerization. The main biosynthetic route, the phenylpropanoid pathway, synthesizes the monolignols p-coumaryl, coniferyl, and sinapyl alcohol from phenylalanine. Studies in the model plant Arabidopsis have shown that cinnamate
4-hydroxylase (C4H), p-coumarate 3-hydroxylase, cinnamoyl-CoA reductase (CCR), and 4-coumarate:CoA ligase (4CL) play roles in regulating the lignin content, while 4CL simultaneously changes the lignin composition. Homologs of these genes have been identified in many woody trees and have been shown to have similar functions in lignin biosynthesis in these species. Several transcription factors (TFs) involved in regulating the lignification process have also been identified, including those from the NAC and MYB families. More than nine MYBs have been reported as lignin biosynthesis-related TFs in Arabidopsis, functioning as either activators or repressors of lignin pathway gene expression. Genome and transcriptome analyses have provided valuable information about the lignin biosynthetic pathway and its regulation in plant species.

Most studies on lignin focus on seedlings, roots, leaves, and stems from model plants and woody trees. However, lignin is also important in some fruits. Although most fruits only contain trace amounts of lignin, some fruits accumulate high levels of lignin in different layers, such as in loquat flesh, mangosteen pericarp, peach endocarp, and stone cells in pear pulp. In these fruits, lignification influences the quality and/or storability and thus consumer acceptance. Recent studies have shown correlations between the activities of lignin biosynthetic pathway and its regulation in plant species.

Plant materials and treatments
Loquat fruits (Eriobotrya japonica Lindl., cv. Luoyangqing) were harvested from an orchard in Luqiao, Zhejiang, China. Fruits were harvested at commercial maturity and transported to the laboratory on the day of harvest and then screened for uniform size and maturity and absence of disease and mechanical damage. Two separate pools from the same batch of fruits underwent HT and LTC, and another pool was stored at 0 °C. Physiological changes in fruits under LTC (5 °C for 6 days followed by transfer to 0 °C for 2 days) were described in our previous report. For HT, the loquat fruits were treated at 40 °C for 4 h and then transferred to 0 °C. Three biological replicates were performed for each treatment. The fleshy tissues were collected at days 0, 1, 2, 4, 6, and 8 during treatment, quickly frozen in liquid nitrogen, and stored at –80 °C until needed. To compare the different treatments, the initial day (day 0) was not included in the RNA-Seq analysis.

Measurement of fruit firmness and lignin content
Fruit firmness and lignin content are the main indices used to monitor postharvest lignification of loquat fruit. Fruit firmness was measured using a TA.XTplus Texture Analyser (Stable Micro Systems, UK), with a 5-mm diameter probe, a penetration rate of 1 mm s⁻¹, and a penetration depth of 4 mm. The firmness of each fruit was averaged from two measurements, 90° apart at the fruit equator, after removal of a small piece of peel. Fruit firmness was expressed as Newtons (N), and 10 individual fruit replicates were tested.

Lignin content was determined according to the method described by Shan et al. The frozen sample was ground into powder and homogenized in 5 ml of washing buffer (100 mM K₂HPO₄/KH₂PO₄, 0.5% Triton X-100, 0.5% PVP, pH 7.8). The mixture was cultured on a shaker at room temperature at 200 rpm for 30 min and then centrifuged (6000 × g, 25 °C) for 20 min. The pellet was suspended and washed twice in the washing buffer and then four times in 100% methanol. The pellet was dried at 80 °C in an oven overnight. Ten milligrams of the dry powder were dissolved in 1.0 ml of 2.0 M HCl and 0.1 ml of thioglycolic acid. The mixture was boiled in a water bath (100 °C) for 8 h and then cooled on ice for 5 min.
before centrifugation at 10,000 × g for 20 min at 4 °C. The pellet was washed with distilled water and suspended in 2.0 ml of 1.0 M NaOH. After agitating slightly at room temperature for 18 h, the solution was centrifuged at 10,000 × g for 20 min. The supernatant (0.5 ml) was transferred to a new tube with 0.1 ml of concentrated HCl. The tubes were left at 4 °C for 4 h to precipitate the lignin thioglycolic acid, followed by centrifugation at 10,000 × g for 20 min at 4 °C, and the precipitate was dissolved in 1 ml of 1.0 M NaOH. Absorbance was measured at 280 nm using 1.0 M NaOH as the blank. Data were expressed on a fresh weight basis, and all measurements were done in triplicate.

RNA-Seq library preparation
Total RNA was extracted from the fleshy tissues using the QIAGEN RNAeasy Plant Mini Kit according to the manufacturer’s instructions (QIAGEN, California, USA). RNA quality was evaluated via electrophoresis on 1% agarose gels, and RNA quantity was determined by a NanoDrop 1000 spectrophotometer (Thermo Scientific, Wilmington, DE, USA). Strand-specific RNA-Seq libraries were constructed using the protocol described in Zhong et al.26 and sequenced on the Illumina HiSeq 2500 platform in the single-end mode.

RNA-Seq data processing, de novo assembly, and comparative transcriptome profiling
Raw RNA-Seq reads were first processed with Trimmomatic27 to trim adapter and low-quality sequences. After trimming, reads <40 bp were discarded, RNA-Seq reads were aligned to the ribosomal RNA database28 using Bowtie29, and those that could be aligned were removed. The final cleaned reads were de novo assembled into contigs using Trinity30 with the minimum kmer coverage set to 10. The cleaned reads were then aligned to the assembled contigs using Bowtie29. Contigs in which the aligned reads in the sense direction were <10% of those aligned in the antisense direction were discarded since they might be false transcripts due to incomplete digestion of the second strand during the strand-specific RNA-Seq library construction step. Contigs having no match to plant sequences but matching sequences from bacteria, fungi, viruses, or archaea in the GenBank nt database were removed. The redundancies of Trinity-assembled contigs were removed using iAssembler31.

To functionally annotate the final assembled contigs, their sequences were blasted against the TrEMBL, Swiss-Prot, and Arabidopsis protein (TAIR) databases, with E-value cutoffs of 1E−10. Based on their hits in the three databases, gene ontology (GO) terms were assigned to the assembled contigs.

The final cleaned high-quality RNA-Seq reads were aligned to the assembled contigs using Bowtie29. Based on the alignments, raw counts of each contig were derived and normalized to RPKM (reads per kilobase exon model per million mapped reads). Raw counts were fed to edgeR32 to identify differentially expressed genes (DEGs) between LTC or HT and 0 °C with a cutoff of fold change ≥2 and false discovery rate ≤0.05. Clustering of gene expression profiles was performed using STEM33.

Entropy weight method
To evaluate the importance of DEGs based on their absolute expression levels (RPKM) and relative expression levels (fold change), an entropy weight method was used. For an m-indicator and n-gene problem, the entropy of the ith indicator is defined as

\[ H_i = \frac{1}{\ln n} \sum_{j=1}^{n} f_{ij} \ln f_{ij}, \quad i = 1, 2, ..., m \]

where \( f_{ij} = r_{ij} / \sum r_{ij} \). Thus the weight for each entropy is

\[ w_i = \frac{1 - H_i}{m - \sum_{i=1}^{m} H_i}. \]

Here five expression data points under LTC (or HT) and five-fold-change data points compared to the corresponding time points under 0 °C were used to calculate the entropy and generate a sorted list of DEGs in LTC or HT. This method was implemented in MATLAB.

Real-time PCR analysis
Gene-specific oligonucleotide primers were designed for real-time PCR and are listed in Supplementary Table 6. The LightCycler 1.5 instrument (Roche) was used for real-time PCR. It was initiated by 5 min at 95 °C followed by 45 cycles of 95 °C for 5 s, 60 °C for 5 s, and 72 °C for 10 s. Finally, it was completed with a melting-curve analysis. The PCR mixture (10 μl total volume) contained 2 μl of 5× LightCycler FastStart DNA MasterPLUS SYBR Green I Master Mix (Roche), 0.5 μl of each primer (10 μM), 1 μl of diluted cDNA, and 6 μl of PCR-grade H2O. Each run included no-template controls and a melting-curve analysis.

Results
Regulation of loquat fruit lignification
Ripe loquat fruits (cv. Luoyangqing) were sampled and stored under three different conditions: low temperature (0 °C), HT (40 °C for 4 h followed by transfer to 0 °C), and LTC (5 °C for 6 days followed by transfer to 0 °C). The lignin content increased from 3.43 × 10^3 A280/kg FW to 4.58 × 10^3 A280/kg FW over a period of 8 days at 0 °C. Fruit stored under HT had a lower lignin content than fruit stored at 0 °C (Fig. 1). Similar results, showing a reduction in both lignin and firmness of loquat fruits during storage at LTC, were reported previously22.
RNA-Seq, de novo assembly, and functional annotation

To obtain a better understanding of the molecular mechanisms underlying the lignin regulation of loquat fruits stored under these three different conditions, we performed comparative transcriptome analysis using RNA-Seq. A total of 48 RNA-Seq libraries were constructed and sequenced. These libraries were from samples at five or six time points for each of the three treatments (0 °C, LTC, and HT), with three biological replicates for each time point (Supplementary Table 1). After removal of adapters, low-quality sequences and ribosomal RNA reads, a total of approximately 517 million cleaned reads were obtained. Since no reference genome is available for loquat, we de novo assembled these reads into 53,319 unique transcripts with an N50 length of 1306 bp and an average length of 798 bp.

Of these 53,319 assembled transcripts, 36,252 (68.0%) were successfully annotated by homologous proteins in at least one of the three protein databases (TrEMBL, Swiss-Prot, and TAIR), and 26,382 transcripts (approximately 72.8% of the total annotated transcripts) were annotated by homologous proteins in all three databases (Supplementary Figure 1). GO terms were assigned to 35,183 (66.0%) transcripts.

DEGs in response to HT and LTC

We identified 5824 DEGs between LTC and 0 °C samples and 3981 between HT and 0 °C samples. To generate a more focused candidate DEG set, we selected genes differentially expressed at a minimum of two of the five time points that showed the same trend (either an increase or decrease) in any of the treatments. This method reduced the DEG numbers to 2235 and 1020 in LTC vs. 0 °C and HT vs. 0 °C, respectively. The numbers of DEGs at each time point are shown in Fig. 2. The LTC treatment resulted in more DEGs than the HT treatment at all five time points, and there were more upregulated genes than downregulated ones in both LTC and HT. Both LTC and HT treatments showed the largest number of DEGs at day 4.

Genes differentially expressed throughout nearly the entire time course (from day 1 to day 8) were the most responsive to the LTC or HT treatment. Therefore, we selected genes differentially expressed at least at four time points in each treatment: 101 and 50 upregulated and 79 and 13 downregulated genes were detected in LTC and HT, respectively (Supplementary Table 2). A comparison of these genes between the two treatments was performed, and six genes appeared in both LTC and HT (Supplementary Table 3). Of these six genes, two were downregulated and annotated as encoding an alcohol dehydrogenase (UN68301) and an auxin-binding protein (UN21207). The four upregulated genes were annotated as encoding a CBL-interacting protein kinase 08 (UN00386), an RNA-binding protein (UN30230), a UDP-glucuronate decarboxylase protein 1 (UN35156), and a gibberellin 3-beta hydroxylase (UN68191).

When checking DEGs, those with both high absolute and high relative expression levels should be a priority. Here we applied an analysis on both the expression and the fold changes on all five time points (a total of ten values) under LTC or HT using the entropy weight method (see the Materials and methods section for details). The ten-dimensional data were reduced to one value, and a new sorted index was generated for the DEGs to identify the top gene list (Supplementary Table 4). Under the HT condition, genes encoding the heat shock proteins dominated the top list. After excluding these heat shock genes, the top 10 genes (Supplementary Table 5) contained some DEGs mentioned above, such as UN00386 (CBL-CIPK) in the top list of LTC and UN68191 (gibberellin 3-beta hydroxylase) under HT.
Lignin and cell wall biosynthesis

Differentially expressed lignin-related TFs and structural genes

Genes encoding enzymes in the lignin biosynthetic pathway have been widely studied in model plants as well as in some fruit species. In this study, six highly differentially expressed structural genes in the lignin biosynthetic pathway were identified. Except for EjPAL2, the other five structural genes are marked in the late steps of the pathway (Fig. 3). The quantitative reverse transcriptase–PCR validation of these candidate genes’ expression is shown in Supplementary Fig. 3. Their different expression patterns across the LTC or HT treatment indicated that multiple structural genes could act differently at different stages of chilling injury and could be candidate structural genes contributing to loquat fruit lignification at different time points. TFs play crucial roles in regulating lignin biosynthesis, as reported in model plants, woody trees, and crops. We screened our assembled transcripts and predicted a total of 1163 TFs from 62 families, of which 120 and 32 were DEGs under LTC and HT, respectively, which were distributed in 31 families (Fig. 4). Of these TF families, the AP2/ERF family contained the most DEGs (20), followed by the bZIP family with 14 DEGs. Eighteen TFs belonging to ten families were differentially expressed under both LTC and HT.

Coexpression network between lignin-related TFs and structural genes

Using the gene expression profile data, we constructed coexpression networks between different TFs and structural genes based on the Pearson product–moment correlation coefficient. The correlation coefficient was calculated separately using the expression data under LTC and HT. Owing to the limited number of samples, we applied a very stringent cutoff for the correlation coefficient (0.9 for positive correlation and −0.9 for negative correlation). The resulting coexpression networks indicated that EjCAD3 (UN68301) and COMT (UN14792) were highly correlated since most of their connected TFs were in common in the LTC and HT samples (over 80% of all their connected TF nodes) (Fig. 5). Both CAD and COMT catalyze the final steps of the lignin biosynthesis. Their connected TFs included an ERF (UN48864), an NAC (UN68419), a bZIP (UN48861), and an MYB (UN68719). Our DEG analysis showed that UN68419 and UN68719 were differentially expressed at the early time points under LTC or HT, and UN48861 showed high expression levels throughout the entire time course. The HCT (hydroxycinnamoyl transferase) (UN28739) also shared many connected TFs with EjCAD3 and COMT under LTC, while under HT, EjCAD1 (UN68110) shared more TFs connected to EjCAD3 and COMT. The highly differentially expressed AP2 (UN33535) and MYB (UN19253) genes were also highly correlated with EjCAD3 and COMT under the HT condition.

Cell wall metabolism-related DEGs

Genes involved in the biosynthesis of different components of the cell wall other than lignin, including pectin, cellulose, and xylans, were also found to be responsive to LTC or HT treatment. Pectin was the most obviously affected in terms of gene expression changes. We obtained three DEGs related to pectin metabolism (Supplementary Tables 2 and 4): UN12673 (encoding a pectate lyase) and UN35268 (encoding a pectinacetyl esterase) that were upregulated in LTC and UN48794 (encoding a pectinesterase) that was first downregulated and then upregulated throughout the time course in HT (Fig. 6a).

GO term enrichment analysis

We further performed GO term enrichment analysis and identified lignin-/cell-wall-related biological processes enriched in DEGs under both LTC and HT. These processes included cell-wall-related processes, lignin metabolic processes, and phenylpropanoid biosynthetic processes (Table 1). The detailed relationship between the enriched lignification-related biological processes under LTC and HT is shown in Supplementary Fig. 2.

Gene expression patterns

Gene expression clustering was applied to identify genes with similar expression patterns. All the DEGs were clustered into 50 different groups (Fig. 7). Genes in the same clusters are very likely involved in the same biological processes. Therefore, once we confirm the function of a gene in the cluster, other genes in the same cluster could be considered to have similar functions. For the two
identified structural genes in lignin biosynthesis, *EjCAD3* (UN68301) and *COMT* (UN14792), we found several genes in the same cluster that were also identified to be in the same group by GO annotation analysis (Fig. 7), further supporting their potential roles in lignin biosynthesis.

**Other affected biological processes**

In-depth analysis of the filtered DEG sets (Supplementary Tables 2 and 4) identified genes involved in interesting biological processes other than lignin-/cell-wall-related ones under the LTC or HT treatment compared to 0 °C. These biological processes included those related to stress responses (including responses to cold, heat, and water deprivation), regulation of fatty acid biosynthetic processes, hormone metabolic processes, and several metal-ion-related processes (mainly calcium and cadmium transport), all of which were significantly enriched in the DEG sets.

**Stress responses**

CBL-CIPKs and LEA (late embryogenesis abundant) genes have been reported to be induced by cold stress. A number of differentially expressed LEA genes were identified, and the number of these LEA genes at each time point is shown in Fig. 6b. Under HT, more LEA genes were differentially expressed at the early time points (day 1 and day 2). In addition, we identified three LEA genes that were differentially expressed at day 1/day 2 under both LTC and HT treatments, and their expression levels were decreased under both LTC and HT compared to 0 °C (Fig. 6a). Furthermore, a number of heat-shock-related genes were also identified under HT; as expected, they were all upregulated.

**Hormone metabolic processes**

Plant hormones are highly correlated with stress responses under temperature-based postharvest treatments. Changes in plant hormone contents can affect fruit quality. A number of plant-hormone-related genes were responsive to HT or LTC in loquat fruit. The numbers of DEGs related to gibberellin (GA), ethylene (ETH), and auxin (IAA) are shown in Fig. 6c. LTC affected the expression of more IAA- and ETH-related genes than HT, while the expression of a similar number of GA-related genes was affected by LTC and HT.

**Enzymatic browning**

Enzymatic browning is one of the symptoms of loquat fruit chilling injury, which is usually displayed after the fruit are transferred from cold storage to room.
temperature. Enzymatic browning was not observed in the present study due to the short-term storage. However, the transcriptome analysis indicated that two enzymatic-browning-related polyphenol oxidase genes, UN48803 and UN48858, were downregulated under LTC throughout the entire time course (Fig. 6a), which may contribute to the lower browning index in LTC-treated fruits.

Sugar- and acid-related genes

In loquat fruit, fructose, sucrose, glucose, and sorbitol are the four main sugar compounds correlated with fruit taste. From the DEG analysis, we identified the gene UN49695 (encoding a fructokinase) that was upregulated under LTC but not differentially expressed under HT, and another gene, UN49801 (encoding a sorbitol dehydrogenase), that was downregulated under both LTC and HT (Fig. 6a). For fruit acidity, UN00648 (encoding a malic acid enzyme) and UN41276 (encoding a fruit-acidity-related protein) were found to be downregulated under HT, while under LTC, UN00648 was upregulated (Fig. 6a). These results indicated that LTC and HT could affect fruit sugar- and acid-related genes, which would eventually influence the fruit flavor, although LTC and HT could have different impacts.

Discussion

Loquat fruit is one of the best systems for fleshy fruit lignification research. Here we performed deep transcriptome sequencing of 48 loquat fruit samples under three different postharvest storage conditions using RNA-Seq. The transcriptome sequences were de novo assembled into 53,319 transcripts, of which 70.2% were
annotated. In addition to a recently reported loquat transcriptome assembly by Song et al.\textsuperscript{37}, our assembled and annotated loquat transcripts provide a valuable resource for loquat research and improvement.

The primary purpose of the present study was to compare gene expression patterns in loquat fruits stored at 0 °C after an HT or LTC treatment, both of which cause significantly reduced lignification, in order to identify structural and regulatory genes that contribute to loquat fruit lignification during cold storage. Some studies have focused on structural genes to understand the lignin regulation pathway in fruits, such as CCR in pear\textsuperscript{38} and 4CL in mulberry\textsuperscript{39}. Here we have also identified several structural genes, such as those encoding PAL, 4CL, COMT, HCT, and CAD, that contribute to loquat fruit lignification during cold storage. Some studies have focused on structural genes to understand the lignin regulation pathway in fruits, such as CCR in pear\textsuperscript{38} and 4CL in mulberry\textsuperscript{39}. Here we have also identified several structural genes, such as those encoding PAL, 4CL, COMT, HCT, and CAD, that contribute to loquat fruit lignification during cold storage (Fig. 4). We identified the PAL gene \textit{EjPAL2}, which we have previously shown is involved in phenylpropanoid synthesis during early stages of fruit development when there is a considerable increase in vascular tissues\textsuperscript{25}. Phenylpropanoid synthesis is a very early step of lignin biosynthesis, and low temperature mainly causes damage to the vascular tissues in the early stages of fruit development. Therefore, the upregulated expression of \textit{EjPAL2} could reinforce the vascular tissues for improved low-temperature tolerance. The \textit{HCT} and 4\textit{CL} genes were downregulated under both LTC and HT at day 1/day 2, while \textit{COMT} and \textit{EjCAD3} were downregulated at all sampled time points under both treatments. It is also notable that the \textit{HCT} and 4\textit{CL} genes have more significant expression differences under LTC compared to 0 °C than under HT, while \textit{EjPAL2} and \textit{COMT} have more significant expression differences under HT compared to 0 °C than under LTC. Thus these results indicated that multiple structural genes are involved in fruit lignification; however, similar repression effects of LTC and HT treatment may have distinct main targets. As mentioned above, \textit{COMT} and \textit{EjCAD3} were highly correlated. The \textit{COMT} gene merits a more in-depth functional analysis since there is no report on \textit{COMT} genes regulating loquat fruit lignification, while evidence has shown the importance of \textit{COMT} in lignin biosynthesis in Arabidopsis\textsuperscript{40,41}. Collectively, our results indicate that various lignin biosynthetic genes contribute to loquat fruit lignification during cold storage.

Studies in model plant species have shown that lignin biosynthesis is regulated by multiple TFs, such as those in
the NAC and MYB families. Our previous studies also showed that several TFs, including *EjMYB1/2*, *EjMYB8*, and *EjAP2-1*, are involved in the regulation of loquat fruit lignification. In the present study, RNA-Seq analysis identified a total of 134 TFs in loquat fruit belonging to 31 families that were responsive to either low-temperature conditioning (LTC) or high temperature (HT). Table 1 shows the enriched GO terms related to lignification, cell wall metabolism, and abiotic stress responses under LTC and HT.

| GO ID      | GO annotation                                         | Corrected P (LTC) | Corrected P (HT) |
|------------|-------------------------------------------------------|-------------------|------------------|
| GO:0009414 | Response to water deprivation                         | 5.695E−13         | 1.115E−06        |
| GO:0009409 | Response to cold                                       | 2.365E−09         | 0.0007           |
| GO:0009408 | Response to heat                                       | 7.346E−06         | 0.007            |
| GO:0071555 | Cell wall organization                                | 3.262E−06         | 3.732E−08        |
| GO:0042545 | Cell wall modification                                | 0.0105            | 0.0388           |
| GO:0052386 | Cell wall thickening                                  | 0.0133            | 0.0046           |
| GO:2000652 | Regulation of secondary cell wall biogenesis          | 0.0064            | 0.0281           |
| GO:0009808 | Lignin metabolic process                              | 0.0001            | 0.0038           |
| GO:1901141 | Regulation of lignin biosynthetic process             | 0.0001            | 0.011            |
| GO:0009699 | Phenylpropanoid biosynthetic process                  | 0.0006            | 0.0026           |
| GO:0009809 | Lignin biosynthetic process                           | 0.0047            | 0.0079           |
| GO:0046274 | Lignin catabolic process                              | 0.0047            | 0.0037           |
| GO:0070588 | Calcium ion transmembrane transport                   | 0.0001            | 0.0284           |
| GO:0035584 | Calcium-mediated signaling using intracellular calcium | 0.024             | 0.0299           |
| GO:0030001 | Metal ion transport                                   | 0.0017            | 0.0041           |
| GO:0015691 | Cadmium ion transport                                 | 0.0074            | 0.0299           |
| GO:0042445 | Hormone metabolic process                             | 0.0139            | 0.0286           |
| GO:0042304 | Regulation of fatty acid biosynthetic process         | 0.0159            | 0.0328           |

GO gene ontology, HT heat treatment, LTC low-temperature conditioning.

Fig. 7 Gene expression data clustering. Expression profile clustering of genes differentially expressed under low-temperature conditioning (LTC, a) or high temperature (HT, b) compared to 0 °C. Clusters with colored backgrounds have a statistically significant number of differentially expressed genes assigned vs. the number expected. The black line in each box is the extracted expression pattern for each cluster. Genes in the same group with *EjCAD3* under LTC and COMT under HT, which were also identified by the gene ontology annotation analysis, are listed.
LTC or HT. The previously reported TFs, which include *EjAP2-1* (*UN17693*) and three MYB TF genes including *EjMYB8* (*UN08246*), are among the 134 identified TFs. In addition to these previously reported TFs, other TFs that showed similar responses to LTC and HT, such as the NAC gene (*UN68419*), are also candidate regulators of loquat fruit lignification, although their exact functions require future analyses. The NAC gene (*UN22247*) identified from expression clustering, which appears in the coexpression networks of HT, and another NAC gene (*UN04037*) correlated with *EjPAL2* under LTC are also candidate regulators. Our previous study indicated that *EjNAC3* directly binds to the *EjCAD1*-like promoter. Based on the RNA-Seq results, this NAC (*UN04037*) may regulate fruit lignification via *EjPAL2*, which may reveal a novel mechanism for NAC regulation of lignification. The present omics-based analyses have expanded the view of possible TF activities in loquat fruit. The predicted correlations between TFs and structural genes have given us useful information that can guide us in the performance of more in-depth analyses of loquat fruit lignification induced by chilling injury.

Unlike the lignin biosynthetic genes, other functional genes involved in fruit lignification have rarely been studied. For instance, there are three LEA genes down-regulated by both LTC and HT (Fig. 6a), which are also positively correlated with loquat fruit lignin content. These LEA genes showed similar responses to both LTC and HT treatments, especially in the early stage (day 1 and day 2), which may be the main stress-related genes in loquat lignification. However, in loquat fruit, relationships between LEA genes and lignification have not been reported. The roles of LEA in lignin biosynthesis have been rarely reported in other plants, but when *IbLEA14* was overexpressed, sweet potato calli had higher lignin contents of monolignol and the expression of related genes. Other notable information came from our GO enrichment analysis. Although many DEGs were annotated as being related to the cold response and lignin biosynthesis, as expected, a group of metal-ion-related genes were also differentially expressed. The correlation of metal ion and cold responses have been widely studied in other plant species, and it has been reported that the increased cold tolerance of wheat leaves occurs not only at a low hardening temperature (4 °C) but also in the presence of cadmium. In addition, cadmium treatment can increase the activity of PAL, an important enzyme in lignin biosynthesis. There are also research findings on the effect of Ca on the cell wall in fruit. Divalent metal ions contribute to the cold resistance of the cell wall in fruit, and the dynamic equilibrium of the ion-binding state in the cell wall is affected by the temperature and the ion concentration in the environment. These genes, as well as some others, have rarely been studied in relation to loquat fruit lignification but may contribute to it.

**Conclusions**

Loquat fruits accumulate high amounts of lignin during postharvest storage under chilling conditions, which substantially decreases their quality and storability and thus consumer acceptance. Loquat fruits treated with LTC or HT showed reduced lignification. By comparing the transcriptome profiles of loquat fruits at different time points under LTC or HT to those under chilling (0 °C), we identified a large number of DEGs and found that biological processes such as stress responses, cell wall and lignin metabolism, hormone metabolic processes, and metal ion transport were significantly affected by LTC or HT compared to 0 °C. Key genes in the lignin biosynthetic pathway that were differentially expressed under LTC or HT compared to 0 °C were identified, which may contribute directly to the regulation of lignification in loquat fruit under chilling. A number of TFs showed differential expression under LTC or HT compared to 0 °C, and their expression profiles were highly correlated with those of the key lignin biosynthetic genes. These TFs included *EjMYB1/2*, *EjMYB8*, and *EjAP2-1*, which were reported previously to be involved in the regulation of loquat fruit lignification. Further functional studies are needed to elucidate the regulatory roles of the newly identified TFs in loquat fruit lignification. The findings of this work contribute to a better understanding of the molecular mechanisms underlying the lignification process during the postharvest storage of loquat fruit under chilling conditions.

**Data availability**

The raw RNA-Seq data have been deposited into the NCBI Sequence Read Archive (SRA) under the accession number SRP128075. This Transcriptome Shotgun Assembly project has been deposited at DDBJ/EMBL/GenBank under the accession number GGEW00000000. The version described in this paper is the first version, GGEW01000000. The assembled transcriptome sequences can also be downloaded from [http://www.ppilixyz.loquat/download.html](http://www.ppilixyz.loquat/download.html).

**Acknowledgements**

This research was supported by grants from the National Natural Science Foundation of China (31630067, 11771393, and 11632015), the Natural Science Foundation of Zhejiang Province, China (LR16C150001 and LZ14A010002), the Project of the Science and Technology Department of Zhejiang Province (2016C04001), and the US National Science Foundation (IOS-1539831).

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**Conflict of interest**

The authors declare that they have no conflict of interest.
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