Transcriptome Analysis and Ultrastructure Observation Reveal that Hawthorn Fruit Softening Is due to Cellulose/Hemicellulose Degradation

Jiayu Xu¹, Yuhui Zhao¹, Xiao Zhang¹, Lijie Zhang², Yali Hou¹ and Wenxuan Dong¹

¹ College of Horticulture, Shenyang Agricultural University, Shenyang, China, ² College of Forestry, Shenyang Agricultural University, Shenyang, China

Softening, a common phenomenon in many fruits, is a well coordinated and genetically determined process. However, the process of flesh softening during ripening has rarely been described in hawthorn. In this study, we found that ‘Ruanrou Shanhong 3 Hao’ fruits became softer during ripening, whereas ‘Qiu JinXing’ fruits remained hard. At late developmental stages, the firmness of ‘Ruanrou Shanhong 3 Hao’ fruits rapidly declined, and that of ‘Qiu JinXing’ fruits remained essentially unchanged. According to transmission electron microscopy, the middle lamella of ‘Qiu JinXing’ and ‘Ruanrou Shanhong 3 Hao’ fruit flesh was largely degraded as the fruits matured. Microfilaments in ‘Qiu JinXing’ flesh were arranged close together and were deep in color, whereas those in ‘Ruanrou Shanhong 3 Hao’ fruit flesh were arranged loosely, partially degraded and light in color. RNA-Seq analysis yielded approximately 46.72 Gb of clean data and 72,837 unigenes. Galactose metabolism and pentose and glucuronate interconversions are involved in cell wall metabolism, play an important role in hawthorn texture. We identified 85 unigenes related to the cell wall between hard- and soft-fleshed hawthorn fruits. Based on data analysis and real-time PCR, we suggest that β-GAL and PE4 have important functions in early fruit softening. The genes Ffase, Gns,α-GAL, PE63, XTH, and CWP, which are involved in cell wall degradation, are responsible for the different textures of hawthorn fruits. Thus, we hypothesize that the different textures of ‘Qiu JinXing’ and ‘Ruanrou Shanhong 3 Hao’ fruits at maturity mainly result from cellulose/hemicelluloses degradation rather than from lamella degradation. Overall, we propose that different types of hydrolytic enzymes in cells interact to degrade the cell wall, resulting in ultramicroscopic Structure changes in the cell wall and, consequently, fruit softening. These results provide fundamental insight regarding the mechanisms by which hawthorn fruits acquire different textures and also lay a solid foundation for further research.

Keywords: hawthorn, TEM observation, RNA-Seq, different texture, cell wall
INTRODUCTION

Softening is a common phenomenon in many fruits and is an important factor that influences fruit quality. From horticultural and commercial viewpoints, ripening grants positive attributes to fruit and greatly impacts its various quality and nutritional characteristics, including fiber content and composition, lipid metabolism, and vitamin and antioxidant contents (Kalt, 2002). Ripening-related softening is a complex multi-component process. Fruit softening is attributed to the coordinated actions of several enzymes on different cell wall polymers. During the ripening process of fleshy fruits, it is generally accepted that textural changes, mainly associated with decreased firmness, result from the reduction of the cell to cell interactions due to the dissolution of the middle lamella, loosening of the primary cell wall and a reduction in cell turgor (Goulao and Oliveira, 2008; Mercado et al., 2011). During ripening, some fruits (e.g., kiwi, tomato, and European pear) soften to a melting texture, while other fruits (e.g., apple, watermelon, and Asian pear) partially soften and remain relatively firm and crisp (Ng et al., 2015). The extent of disassembly is linked to the chelator-soluble fraction of the cell wall. In one study, several transgenic plants with reduced cell mRNA levels were obtained, but no significant correlation was found with endoglucanase (EG) activity or fruit firmness (Woolley et al., 2001). Ortiz et al. (2010) indicated that a portion of the arabinose-rich side chains removed from the pectic polymers remained transiently linked to the chelator-soluble fraction of the cell wall. In addition, sugar analyses suggested that the disassembly of cell walls is aided by the previous elimination of galactan sidechains, which could facilitate pectin solubilization. The activity patterns of the considered cell wall-modifying enzymes were very similar. Given the complexity of the ripening process, the use of tools that allow global evaluations of the molecular processes triggered within the fruit are important. Some studies have already studied the softening of apple (Harb et al., 2012; Gwanpua et al., 2016), tomato (Lunn et al., 2013), peach (Yoshioka et al., 2011; Cao et al., 2014), strawberry (Posé et al., 2015), and pear (Hiwasa et al., 2004; Chiriboga et al., 2012; Wei et al., 2015) fruits.

The genus *Crataegus* (hawthorn), which is a common type of fruit, belongs to the Rosaceae family and is a genus of spiny trees or shrubs that exist in the northern hemisphere (Verma et al., 2007). Hawthorn that is planted for human consumption can be traced back to 300 A.D. In addition, hawthorn has been approved by the Ministry of Health of the People’s Republic of China as a raw material for functional foods and has been included in the Chinese Pharmacopeia as an herbal medicine (Liu et al., 2010). Furthermore, hawthorn is one of the most widely consumed horticultural products, either in fresh or processed form, and is an important component of many processed food products because of its excellent flavor, attractive color and high macro- and micro-nutrient contents (Cao et al., 1995; Özcan et al., 2005). To date, over 150 compounds, especially phenolic compounds, have been identified in *Crataegus pinnatifida* (Wu et al., 2014). Among these phenolic compounds, procyanidins (procyanidin B2, procyanidin B3, and procyanidin C1), flavonoids (epicatechin, hyperoside, quercetin, rutin, and isoquercitrin) and triterpenoids acid (ursolic acid, corosolic acid, oleanolic acid, and maslinic acid) are important bioactive components of hawthorn fruits (Liu et al., 2011; Yang and Liu, 2012; Chai et al., 2014; Wu et al., 2014).

In recent years, Illumina RNA-Seq and digital gene expression (DGE) have provided an opportunity to explore different phenomena in diverse species via de novo assembly, facilitating the identification and analysis of the majority of transcriptomes (Zhang et al., 2010). Using RNA-Seq technology, Dai et al. (2013) found that degraded fruit endocarp (stone) formation might be correlated with decreased lignin synthesis in a unique hawthorn variety, 'soft-endocarp,' which has a soft and broken endocarp (stone), whereas the endocarp of most hawthorn varieties are hard. Nevertheless, few data are available regarding the different textures of hard- and soft-fleshed hawthorn fruits. Almost 300 resources are present in the National Hawthorn Germplasm Repository of Shenyang, China. 'Ruanrou Shanlihong 3 Hao' is a wild cultivar of *C. pinnatifida* with lower flesh firmness during the fruit maturing and harvesting periods than hawthorn fruits with average firmness. Thus, 'Ruanrou Shanlihong 3 Hao' fruit is a preferable fruit for studying fruit softening. 'Qiu JinXing' is an existing cultivar with fruits that are firmer during fruit ripening and with good fruit qualities and properties. Therefore, 'Qiu JinXing' and 'Ruanrou Shanlihong 3 Hao' were chosen as experimental materials for this study. Transmission electron microscopy (TEM) was used to observe the microstructure of the fruit flesh cells and Illumina-based RNA-Seq was used to elucidate differential gene expression (Shin et al., 2015). For this analysis, ultrathin sections of hawthorn flesh were obtained and a de novo assembly of the fruit transcriptome of *C. pinnatifida* was performed. Using RNA-Seq, the hawthorn transcriptomes were sequenced, assembled, and annotated for the hawthorn fruits with hard and soft flesh. Using these platforms, the global transcriptional analysis of gene expression in the hawthorn fruits with hard and soft flesh were described. When determining the different textures of the hawthorn fruit cell walls, the cell wall metabolism was investigated at the molecular level in addition to the relationships between the important enzymes in the cell wall. Fruit softening is of theoretical significance and should be quantified to further reveal the mechanisms of hawthorn fruit softening during maturation.
MATERIALS AND METHODS

Plant Material
The hard-fleshed hawthorn ‘Qiu JinXing’ and the soft-fleshed hawthorn ‘Ruanrou Shanlihong 3 Hao’ were maintained in the National Hawthorn Germplasm Repository of China at Shenyang. Fruits were collected at the early developmental stage I (RRSI- 90 days before harvesting ‘Ruanrou Shanlihong 3 Hao’; QJXI- 105 days before harvesting ‘Qiu JinXing’), early developmental stage III (RRSII- 75 days before harvesting ‘Ruanrou Shanlihong 3 Hao’; QJXII- 75 days before harvesting ‘Qiu JinXing’), middle developmental stage I (RRSIII- 60 days before harvesting ‘Ruanrou Shanlihong 3 Hao’; QJXIII- 45 days before harvesting ‘Qiu JinXing’), middle developmental stage II (RRSIV- 45 days before harvesting ‘Ruanrou Shanlihong 3 Hao’; QJXIV- 30 days before harvesting ‘Qiu JinXing’), late developmental stage I (RRSV- 15 days before harvesting ‘Ruanrou Shanlihong 3 Hao’; QJXV- 15 days before harvesting ‘Qiu JinXing’), and late developmental stage II (RRSVI- the day of harvest of ‘Ruanrou Shanlihong 3 Hao’; QJXVI- the day of harvest of ‘Qiu JinXing’) (Figure 1). Fruits were selected for their uniform size, appearance, and lack of defects. Fifteen fruits were analyzed to determine their firmness and weight at different developmental stages. The remaining fruits were frozen in liquid N₂ and stored at −80°C for further analysis.

Measurement of Fruit Firmness and Weight of Hawthorn Fruits with Hard and Soft Flesh
Fruit firmness was measured on the opposite sides of each individual fruit among the 15 analyzed fruits at different developmental stages by using a non-damage texture analyzer (test anvil, cylinder 0.10 cm²; spring force, 806.50 mN; contact pressure, 12.50 N; resolution, 0.10 Ff; HPE II Ff, Bareiss, Germany), and the maximum force developed during the test was recorded. Firmness was determined as the average force in Newtons (N). Fruit total sugar, titrable acidity were measured using the method according to Ranganna (2001). An electronic analytical balance (YP1002N, Shanghai jingke, China) was used to measure the fruit weight at different developmental stages, and all statistical analyses were conducted using SPSS 19.0.

Observations of the Ultrastructures of the Fruit Cell Walls
The flesh tissues of the selected ‘Ruanrou Shanlihong 3 Hao’ and ‘Qiu JinXing’ fruits sampled at different developmental stages were fixed in 2.5% glutaraldehyde and 1% osmic acid, washed with PB buffer solution, dewatered using a gradient of ethanol concentrations, embedded with SPI812 resin, sliced using an Leica-EM-UC 7 ultramicrotome (thickness of 90 nm; Germany), dyed with uranyl acetate and lead citrate, and observed and photographed using a HT – 7700 TEM (Hitachi, Japan).

RNA Isolation, cDNA Library Preparation and Sequencing
Total RNA isolated from flesh was extracted using the modified CTAB method according to Chang et al. (2007). The RNA purity was assessed using a NanoPhotometer spectrophotometer (IMPLEN, USA), and the RNA concentration was measured using a Qubit® RNA Assay Kit (Q32852) with a Qubit2.0 Fluorometer (Q32857; Life Technologies, USA). The quality of total RNA was evaluated using an Agilent Bioanalyzer 2100 system (G2939AA, Agilent Technologies, USA). Only samples with an RNA integrity number (RIN) ≥ 7 and a 28S:18S RNA ratio ≥ 1.5 were used for deep sequencing. For mRNA library construction and deep sequencing, RNA samples were prepared using NEBNext Ultra™ RNA Library Prep Kit for Illumina® (E7530, NEB, USA) according to the manufacturer’s protocol, including poly-A mRNA purification and fragment, first-strand cDNA synthesis, second-strand cDNA synthesis.
and repair, adapter ligation, PCR enrichment, agarose gel purification, and library quality control. cDNA samples were prepared from the fruits of hard- and soft-fleshed hawthorns and sequenced using Illumina HiSeq purification, and library quality control. cDNA samples were and repair, adapter ligation, PCR enrichment, agarose gel purification, and library quality control. cDNA samples were prepared from the fruits of hard- and soft-fleshed hawthorns and sequenced using Illumina HiSeq purification methods.

**De novo Transcriptome Filtering, Assembly and Annotation**

Four samples, including samples of the two hard-fleshed hawthorns QJXIV (‘Qiu JinXing’ fruit at the middle developmental stage I) and QJXV (‘Qiu JinXing’ fruit at the late developmental stage I), the two soft-fleshed hawthorns RRSIV (‘Ruanrou Shanlihong 3 Hao’ fruit at the middle developmental stage I) and RRSV (‘Ruanrou Shanlihong 3 Hao’ fruit at the late developmental stage I), were sequenced. Each of the samples had two biology repeats. Raw data (raw reads) in the fastq format were filtered by removing the reads containing the adaptor, including the reads whose poly-N content was more than 10% and the reads with more than 50% bases and whose quality score Q was no more than five from the raw data. The remaining data were termed clean data (clean reads). Simultaneously, the Q20, Q30, GC-content and sequence duplication levels of the clean data were calculated. The following analyses were based on clean data. The left file was named left.fq, and the right file was named right.fq. Transcriptome assembly was accomplished based on left.fq and right.fq using Trinity (Grabherr et al., 2011) with min-kmer-cov set to two and all other parameters set to default values. Based on the NR (Deng et al., 2006), Swiss-Prot (Apweiler et al., 2004), gene ontology (GO) (Ashburner et al., 2011; KEGG (Kanehisa et al., 2004) databases, BLAST (Altschul et al., 1997) was used to achieve the amino acid sequences of the unigenes and the HMMER (Finn et al., 2013) software and Pfam (Eddy, 1998) database alignment were used to obtain annotated unigenic information. The raw data has been deposited at NCBI database, and the accession number of SRA database is: SRP082543.

**Differential Expression Analysis**

Gene expression levels were measured in the RNA-Seq analysis as reads per kilobase of exon model per million mapped reads FPKM (Trapnell et al., 2010). The DESeq software (Anders and Huber, 2010) was used to identify differentially expressed genes in a pairwise comparison, and the false discovery rate (FDR) and fold change (FC) were used to determine the DEGs. In our study, genes with FDR ≤ 0.01 and FC ≥ 2 in FPKM were considered as DEGs.

**Quantitative Reverse Transcriptase Real-time PCR Analysis**

First-strand cDNA was synthesized from 1 µg of RNA using PrimeScriptTM RT Reagent Kit with gDNA Eraser (RR047A, TaKaRa, China) following the manufacturer’s protocol. The synthesized cDNA was diluted 10-fold for the following Real-Time PCR analysis. Primers for all selected genes were designed using Primer Prime 5 software. The annealing temperature and primer sequence of the analyzed amplicons are listed in Supplementary Table S2, and amplification products were sequenced (Supplementary Figure S1) to check the product identity. For each gene, a cDNA serial dilution was used to generate a standard curve to calculate efficiencies. Quantitative PCR was performed using an iQ5 Real-Time PCR detection system (Bio-Rad, USA) with 20 µl reactions including 1 µl diluted cDNA sample, 1 µl each primer (10 µM) and 10 µl SYBR Green Master Mix (RR820A, Takara, China) under the following conditions: an initial hot start at 95°C for 3 min, followed by 40 cycles of 95°C for 10 s, the annealing temperature for each selected gene for 30 s and 72°C for 30 s, following melting curve range from 50 to 90°C. Two internal control genes, Actin and NADPH, were used for normalization, and relative gene expression was estimated using threshold cycles and the 2−ΔΔCT method (Livak and Schmittgen, 2001). Real-time PCR analysis was carried out with three biological and three technical replicates.

**RESULTS**

**Changes of Fruit Firmness, Weight, Total Sugar, and Titratable Acidity at Different Fruit Developmental Stages**

Fruit firmness is an important factor that affects fruit texture, ripening and softening. At the early and middle stages of fruit development, the firmness of ‘Ruanrou Shanlihong 3 Hao’ and ‘Qiu Jingxing’ fruits were maintained at a high level, with the value of firmness of the two cultivars almost equal. However, at developmental stages V and VI, the firmness of 'Ruanrou Shanlihong 3 Hao' fruits decreased rapidly, whereas that of ‘Qiu JinXing’ declined slowly and remained at a significantly higher level (Table 1). At maturity, the firmness of ‘Qiu JinXing’ fruits was higher than that of ‘Ruanrou Shanlihong 3 Hao’ fruits (Table 1). With fruit development, the fruit weights of ‘Ruanrou Shanlihong 3 Hao’ (soft flesh) and ‘Qiu Jingxing’ (hard flesh) gradually increased, accumulation of total sugar in the two cultivars were declined at early developmental stage I, and then increased gradually (Table 1). Titratable acidity contents of ‘Ruanrou Shanlihong 3 Hao’ fruit were increased gradually and reached to a peak value at middle developmental stage I and then decreased, whereas that in ‘Qiu Jinxing’ fruit were increased continuously (Table 1).

**Ultraprofifile Structure of the Fruit Flesh at Different Fruit Developmental Stages**

Plant cell wall structure is an important factor that determines fruit firmness, and plant cell walls are complex structures composed of cellulose, hemicellulose, pectin and protein. TEM...
can be used to observe the ultramicroscopic structure of fruit flesh tissue clearly. At the middle developmental stages of 'Ruanrou Shanlíhong 3 Hao' and 'Qiu JinXing,' the structures of the cell walls in the fruit flesh were complete, presenting a light-dark-light structure and a plasma membrane located next to the cell wall. Moreover, deep-colored and closely arranged microfilaments and dark middle lamella were observed (Figures 2A,B). At the late developmental stage, the middle lamella of 'Ruanrou Shanlíhong 3 Hao' and 'Qiu JinXing' fruit flesh tissue was nearly all degraded, the light-dark-light structure disappeared, and plasmolysis occurred (Figures 2C,D). However, the microfilaments in the 'Qiu JinXing' fruit tissue were arranged closely and colored deeply (Figure 2D), whereas the microfilaments in the 'Ruanrou Shanlíhong 3 Hao' fruit flesh tissue were arranged loosely, partially degraded and became lighter in color (Figure 2C). The trend of the changes in fruit firmness was nearly consistent with the TEM observations.

RNA-Seq and De novo Transcriptome Assembly

In the RNA-Seq experiment of 'Ruanrou Shanlíhong 3 Hao' and 'Qiu JinXing' fruits from two different developmental stages, approximately 46.72 Gb of clean data were obtained for all the samples in the RNA-Seq. In addition, 2,182,914 contigs, 199,204 transcripts, and 72,837 unigenes were obtained, and the N50s of the contigs, transcript and unigene were 88, 1,656, and 1,283, which indicated a high assembly integrity. The total lengths of the contigs, transcript and unigene were 168,024,365, 210,439,324, and 50,660,379, respectively. The mean lengths of the contigs, transcript and unigene were 76,971, 1056,40, and 695.53, respectively. Among the unigenes, 29,081 unigenes (39.93%) had lengths of 200–300 nt, 17,945 unigenes (24.64%) had lengths ranging from 300 to 500 nt, 11,447 unigenes (15.72%) had lengths ranging from 500 to 1,000 nt, 8,837 unigenes (12.13%) had lengths ranging from 1,000 to 2,000 nt, and 5,527 unigenes (7.59%) had lengths of more than 2,000 nt, resulting in a mean length of 695.53 (Table 2).

### Functional Annotation and Classification

Transcriptome analysis is important for elucidating the molecular components of cells and tissues and for interpreting the functional elements of the genome (Qiu et al., 2013). To classify the potential functions of the assembled unigenes, a sequence similarity search was conducted against the NR, Swiss-Prot, GO, COG, KOG, KEGG, and Pfam databases. A threshold magnitude of less than $10^{-5}$ was used for the NR, Swiss-Prot, GO, COG, KOG, KEGG databases and a threshold magnitude of less than $10^{-10}$ was used for the Pfam database. The results showed that 12,143, 27,411, 8,280, 20,603, 22,628, 22,240, and 39,248 unigenes exhibited significant similarity to known protein genes in the seven databases. Overall, 39,701 unigenes were annotated in the seven databases, including 28,466 unigenes with lengths of more than 300 nt, and 13,682 unigenes with length of more than 1000 nt (Table 3).

Based on the nr annotation, 19,887 of the distinct sequences showed top matches with sequences from Prunus persica, 3,393 of the distinct sequences showed top matches with sequences from Fragaria vesca, and only 737 and 704 of the distinct unigene sequences showed top matches with sequences from Bipolaris maydis and Theobroma cacao, respectively (Supplementary Figure S2).

Based on the GO databases, 27,411 putative unigenes of C. pinnatifida were categorized into 55 functional groups that belonged to three main GO ontologies: cellular component, molecular function, and biological process (Figure 3). Next, 71,981, 33,536, and 43,271 unigenes were annotated as the biological process, cellular component and molecular function, respectively. The 'Cell part,' 'Cell,' 'catalytic activity,' and 'metabolic process' terms were most prevalent. In addition, a high percentage of genes were classified under the 'Binding' and 'Cellular process' terms, and only a few genes were classified as 'protein tag,' 'Virion,' and 'synapse part' (Figure 3).

Based on the KOG databases, 12,143 putative unigenes of C. pinnatifida were categorized into 25 functional groups. The cluster for 'General function prediction only' was larger than any

### TABLE 1 | Variations in fruit firmness, weight, total sugar and titratable acidity in ‘Ruanrou Shanlíhong 3 Hao’ and ‘Qiu JinXing’ fruits.

| Cultivar            | Developmental stage | Fruit firmness (N) | Fruit weight (g) | Total sugar (%) | Titratable acidity (%) |
|---------------------|---------------------|--------------------|------------------|-----------------|------------------------|
| Ruanrou Shanlíhong 3 Hao | I       | 44.64 ± 0.19b     | 0.32 ± 0.01j     | 0.99 ± 0.01h    | 0.62 ± 0.01e           |
|                     | II      | 44.98 ± 0.33b     | 0.47 ± 0.01j     | 0.81 ± 0.02h    | 0.66 ± 0.00e           |
|                     | III     | 45.66 ± 0.35b     | 0.52 ± 0.01j     | 1.00 ± 0.03h    | 0.99 ± 0.02c           |
|                     | IV      | 45.38 ± 0.16b     | 0.57 ± 0.01ni    | 1.42 ± 0.08g    | 0.97 ± 0.01c           |
|                     | V       | 39.54 ± 0.41c     | 0.75 ± 0.02gh    | 2.23 ± 0.06e    | 0.85 ± 0.01d           |
|                     | VI      | 28.89 ± 0.58fg    | 1.16 ± 0.02nf    | 2.90 ± 0.03c    | 0.82 ± 0.01d           |
| Qiu JinXing         | I       | 44.26 ± 0.29b     | 0.94 ± 0.02g     | 1.53 ± 0.06g    | 0.66 ± 0.01e           |
|                     | II      | 47.97 ± 0.20a     | 1.27 ± 0.01e     | 0.70 ± 0.03e    | 0.88 ± 0.01d           |
|                     | III     | 46.37 ± 0.19ab    | 1.91 ± 0.03d     | 1.80 ± 0.06f    | 1.02 ± 0.01c           |
|                     | IV      | 46.11 ± 0.33ab    | 2.67 ± 0.03c     | 2.54 ± 0.17d    | 1.22 ± 0.02b           |
|                     | V       | 41.30 ± 0.65c     | 3.52 ± 0.05b     | 5.27 ± 0.16b    | 2.94 ± 0.04a           |
|                     | VI      | 40.03 ± 0.21c     | 4.93 ± 0.09a     | 7.25 ± 0.12a    | 2.99 ± 0.06a           |

I, early developmental stage; II, early developmental stage II; III, middle developmental stage I; IV, middle developmental stage II; V, late developmental stage I; and VI, late developmental stage II. Different letters indicate significant differences among the treatments at p < 0.05.
other group. However, few genes of the ‘cell motility’ category were found (Supplementary Figure S3).

The KEGG pathway is a collection of manually drawn pathway maps on the molecular interaction and reaction network. Pathway analysis provides information regarding biological functions and gene interactions. When all the annotated unigenes were searched for against the KEGG pathway, 8,280 unigenes were mapped to 120 pathways. Among the 120 KEGG pathways, galactose metabolism and pentose and glucuronate interconversions occurred in the cell wall, which suggested that these pathways may play a role in different hawthorn textures.

DEG Expression
Among the four samples, 195 common and different unigenes were found, with 6214 and 4789 different unigenes in RRSV and QJXV and in RRSIV and QJXIV, respectively (Figure 4). Some genes were down-regulated in RRSIV and QJXIV but obviously up-regulated in RRSV and QJXV, and some genes showed increased quantitative expression of RRSIV and RRSV but decreased transcript abundance in QJXIV and RRSVI. We identified 85 distinctly different unigenes with functions related to cell walls among the hawthorns with hard and soft flesh. Galactose metabolism and pentose and glucuronate interconversions were related to fruit conversion, with different degrees of softening in hawthorn fruits with soft and hard flesh.

Genes Related to the Different Texture of Hard- and Soft-Fleshed Hawthorn during Two Periods
To examine the biochemical basis of different textures between two types of hawthorns during the two periods, we performed transcriptome analysis. Expression analysis was conducted for two pairwise comparisons: RRSIV vs. QJXIV and RRSV vs. QJXV. Overall, 2377 up-regulated and 2412 down-regulated unigenes were observed between RRSIV and QJXIV (Supplementary Figure S4A), and 3434 up-regulated...
TABLE 2 | The length distribution of assembled unigenes.

| Length range | Contig       | Transcript | Unigene   |
|--------------|-------------|------------|-----------|
| 200–300      | 2,125,350 (97.36%) | 39,934 (20.05%) | 29,081 (39.93%) |
| 300–500      | 26,794 (1.23%)    | 34,334 (17.24%) | 17,945 (24.64%) |
| 500–1000     | 15,601 (0.71%)    | 43,252 (21.71%) | 11,447 (15.72%) |
| 1000–2000    | 9,646 (0.44%)     | 53,297 (26.75%) | 8,837 (12.13%)  |
| 2000+        | 5,523 (0.25%)     | 28,387 (14.25%) | 5,527 (7.59%)   |
| Total number | 2,182,914         | 199,204     | 72,837     |
| Total length | 168,024,365       | 210,439,324 | 50,660,379 |
| N50 length   | 88                 | 1,656       | 1,283      |
| Mean length  | 76.97              | 1056.40     | 695.53     |

The values in the tables show the numbers of contig/transcript/unigene in the corresponding range. The percentages in the brackets show the ratio of contig/transcript/unigene in the corresponding range. The table of * shows the numbers and percentages of contig with lengths of 0–300.

TABLE 3 | Summary of unigene annotations.

| Annotated databases | Unigene | ≥300 nt | ≥1000 nt |
|---------------------|---------|---------|---------|
| COG                 | 12,143  | 9,496   | 57.71   |
| GO                  | 27,411  | 20,515  | 11.163  |
| KEGG                | 8,280   | 6,293   | 3,305   |
| KOG                 | 20,603  | 15,689  | 8,557   |
| Pfam                | 22,628  | 18,704  | 11,477  |
| Swiss-Prot          | 22,240  | 17,996  | 10,150  |
| rr                  | 39,248  | 28,287  | 13,661  |
| All                 | 39,701  | 28,466  | 13,682  |

Annotated database, different functional databases; Unigene, unigene value of annotated databases; ≥300 nt, the unigene value of annotated databases with a length of more than 300 bases; ≥1000 nt, unigene value of annotated databases with a length of more than 1000 bases.

and 2780 down-regulated unigenes were observed between RRSV and QJXV (Supplementary Figure S4B). To appreciate the biological importance of the 4,789 differentially expressed genes between RRSIV and QJXIV, the lists of significantly down-regulated or up-regulated genes were analyzed with the database for annotation, and 43 unigenes (Supplementary Table S1) with functions related to cell wall structure were expressed differently. Overall, 10 different unigenes were used in galactose metabolism, 6 unigenes were used in pentose and glucuronate interconversions, and so on (Figure 5A). More unigenes were found to be expressed in chloroplasts (GO: 0009507) and the plasma membrane (GO: 0005886) (Supplementary Figure S5B). In the list of 45 up-regulated genes, ‘Gns’ and ‘Ffase’ are the most over-represented genes in terms of the different expressed genes. The expression of CPW in ‘Qiu JinXing’ fruit was higher than that in ‘Ruanrou Shanlihong 3 Hao’ fruit in both RRSIV vs. QJXIV and RRSV vs. QJXV. The expression of glycine-rich structural proteins and thin1 proteins in the cell walls in ‘Ruanrou Shanlihong 3 Hao’ fruit was higher than in ‘Qiu JinXing’ fruit in both RRSIV vs. QJXIV and RRSV vs. QJXV.

Verification of Gene Expression by Real-time PCR

To confirm the differential expression profiles of DEGs identified from RNA-Seq analysis, we selected 14 candidate DEGs with high expression levels for real-time assays. The expression levels of 14 DEGs at different fruit developmental stages were assayed. Real-time PCR indicated that the expression patterns of the 14 DEGs fitted those determined by RNA-Seq analysis. For instance, several up-regulated DEGs (β-GAL) with highly abundant transcripts also exhibited relatively high levels of expression in real-time PCR analysis (Figure 6). The FCs of some DEGs from real-time PCR were different from those obtained from RNA-Seq analysis, partially due to the different sensitivities of the two approaches and their different algorithms. The RNA-Seq generated absolute expression patterns, and real-time PCR showed relative expression values at the transcriptional level.

DISCUSSION

Cell wall degradation is the main factor affecting fruit softening and quality (Bennett and Labavitch, 2008). With the progression of ripening, complex carbohydrates are converted to simple sugars, and the acidity decreases with the accumulation of sugar. Meanwhile, as aromatic compounds accumulate, the cell wall dynamics change, which results in either dehiscence or softening (Klee and Giovannoni, 2011). In previous studies, degradation of the cell wall network structure was observed during the process of fruit maturation in tomato (Orfila et al., 2002). In addition, at the late stage of pineapple softening, the electron density of the middle lamella decreased, the cell wall became uneven, and microfibrils in the cell wall became loose and disordered (Khalil et al., 2007). In this study, the process of fruit softening during the maturation of ‘Ruanrou Shanlihong 3 Hao’ and ‘Qiu JinXing’ fruits was accompanied by fruit cell wall structural changes. The middle lamella of the ‘Ruanrou Shanlihong 3 Hao’ and ‘Qiu JinXing’ fruit flesh tissue were nearly completely degraded. However, the degree of microfibril degradation in the ‘Ruanrou Shanlihong 3 Hao’ fruit flesh was greater than that in ‘Qiu JinXing’ fruit flesh. These results indicated that the difference in fruit firmness between ‘Ruanrou Shanlihong 3 Hao’ and ‘Qiu JinXing’ fruits at maturity mainly resulted from cellulose/hemicellulose degradation rather than middle lamella degradation.
We obtained 46.72 Gb of clean data by RNA-Seq, with 2,182,914 contigs, 199,204 transcripts, and 72,837 unigenes in RNA-Seq. Overall, 12,143, 27,411, 8,280, 20,603, 22,628, 22,240, and 39,248 unigenes exhibited significant similarity to the known protein genes in the NR, Swiss-Prot, GO, COG, KOG, KEGG, and Pfam databases, respectively, and some KEGG pathways. With the application of massive and parallel sequencing technologies,
FIGURE 5 | The enriched pathways of significant DEGs. (A) For RRSV and QJXIV and (B) for RRSV and QJXV. Ordinate: nomenclature of the KEGG metabolic pathway; abscissa: gene numbers that annotated this pathway and the ratio of KEGG in all genes that annotated relative to the total KEGG.
transcriptome information has become generally abundant for not only model organisms on which international research efforts and funding are concentrated but also for non-model organisms. These results add to the current data on *C. pinnatifida* and provide data for future research in this species. In this study, RNA-Seq technology transcriptome analysis of hard- and soft-fleshed hawthorn revealed that 85 hydrolytic enzyme genes (Supplementary Table S1) and two different signaling pathways, galactose metabolism and pentose and glucuronate interconversions,
related to cell wall processes are associated with cell wall degradation.

In galactose metabolism, β-GAL was usually proposed to modify the cell wall via the removal of galactosyl and arabinosyl residues from polysaccharides in the cell wall at the initiation of fruit ripening (Mwaniki et al., 2007; Martinez and Civello, 2008; Chávez Montes et al., 2008; Posé et al., 2013). High β-GAL activity is very important during early ripening of Jonagold apples (Gwanpua et al., 2014), which is supported by our results. In our study, at the middle developmental stage I, a time when hawthorn fruits begin to soften, the expression levels of β-GAL were higher in ‘Ruanrou Shanlihong 3 Hao’ fruits than that in ‘QiujinXing’ fruits. Furthermore, α-GAL and Ffase, which are very important to such metabolism, were found for the first time in this study to be related to fruit softening. Before fruit firmness decreased, the level of α-GAL gene expression in ‘QiujinXing’ fruits was higher than that in ‘Ruanrou Shanlihong 3 Hao’ fruits. When the hawthorn fruits were softening, the level of α-GAL expression in ‘Ruanrou Shanlihong 3 Hao’ increased abruptly, though a continuous low level of expression was observed in ‘QiujinXing’ fruits, with higher levels of gene expression in ‘Ruanrou Shanlihong 3 Hao’ fruits compared to ‘QiujinXing’ fruits. The result suggested that α-GAL plays a role in changing the fruit texture. During the process of ‘QiujinXing’ fruit development in this experiment, the Ffase expression level was weak while the expression level of Ffase was high in the soft-fleshed hawthorn during the process of fruit development, which indicated that Ffase had an important impact on the fruit texture. In pentose and glucuronate interconversions, the level of PE63 expression generally decreased with increasing fruit development in ‘QiujinXing’ fruits. In contrast, in ‘Ruanrou Shanlihong 3 Hao’ fruits, the gene expression level was weak at the early and middle developmental stages and rapidly increased at the late developmental stage. The expression levels of PE63 were higher in ‘Ruanrou Shanlihong 3 Hao’ fruits at the late developmental stage, a time when hawthorn fruits were softening, showing that PE63 plays a role in hawthorn fruit softening. The PE4 expression level showed an increasing then decreasing tendency across the fruit developmental stages in the two cultivars. At middle developmental stage II, when hawthorn fruits begin to soften, the expression levels of PE4 were higher in ‘Ruanrou Shanlihong 3 Hao’ fruits than in ‘QiujinXing’ fruits. Posé et al. (2015) observed that silencing of PL genes reduced pectin degradation during strawberry fruit softening, though this is inconsistent with our results. In our study, the PL expression showed a decreasing, increasing and decreasing tendency at the early, middle and late fruit developmental stages, respectively, and PL expression in the two cultivars did not differ during fruit softening. Therefore, we hypothesize that PL does not affect hawthorn fruit softening. Overall, these results demonstrate PE4 and β-GAL has an important function in the early softening of different textures; α-GAL, Ffase, and PE63 might play key roles in textural changes, though the crucial participation of α-GAL and Ffase in fruit softening is reported for the first time in this study.

Costa et al. (2010) used QTL analysis to show that PG1 expression was associated with fruit firmness in apples, which was later validated by Longhi et al. (2013). Furthermore, Atkinson et al. (2012) showed that PG-suppressed ‘Royal Gala’ apples were firmer than wild-type after ripening. However, in this experiment, PG1 expression was weak during the process of ‘Ruanrou Shanlihong 3 Hao’ fruit development, but it was higher in the hard-fleshed hawthorn at the late stage of fruit development. This result is consistent with the TEM observations of cell walls at the late developmental stage. Another glycosidase, α-ARF, has been well documented for its involvement in cell wall modifications during fruit ripening. α-ARF has been proposed to release arabinosyl residues from the pectic fraction, and its functions have been widely discussed in melon (Nishiyama et al., 2007), apple (Ireland et al., 2014), and pear (Mwaniki et al., 2007). However, the patterns of α-ARF expression were similar to those of PG1 in this study, indicating that this enzyme is not part of the softening process. The level of EG17 expression exhibited a decreasing tendency in both ‘QiujinXing’ and ‘Ruanrou Shanlihong 3 Hao’ fruits, and the levels were nearly the same in the two cultivars at the late developmental stages. The level of GDP-man expression showed a decreasing tendency with fruit developmental stage in ‘QiujinXing’ fruits. At the middle developmental stage, the levels in the two cultivars were nearly the same. At the late developmental stage, the level of GDP-man expression in ‘QiujinXing’ fruits rapidly decreased and was lower than that in ‘Ruanrou Shanlihong 3 Hao’ fruits. XTH expression in ‘QiujinXing’ fruits was higher than that in ‘Ruanrou Shanlihong 3 Hao’ fruits in early developmental stage; however, the level of XTH expression in ‘Ruanrou Shanlihong 3 Hao’ fruits was higher than that in ‘QiujinXing’ fruits in the other developmental stages. Takeda et al. (2015) found that Gns plays a significant role in the degradation and modification of β-1, 3-glucan in fungal cell walls, and this is consistent with our results. The expression level of Gns in our study was high in soft-fleshed hawthorn fruits but weak in hard-fleshed hawthorns at the late developmental stage. Overall, PG1, α-ARF, and EG17 might not participate in textural changes in hawthorn fruits, whereas GDP-man, XTH, and Gns might be important for changing fruit texture in hawthorn fruits. Thus, the mechanisms by which hydrolytic enzymes affect fruit softening are distinct and complex for different types of fruit.

CONCLUSION

We hypothesize that different types of hydrolytic enzymes in cells interacted to degrade the cell wall, which results in different degrees of cell wall degradation and result in fruit softening, which can be assessed by TEM and RNA-Seq. These results of this study suggest that soft and hard flesh textures in C. pinnatifida fruits are resulted from different degrees of cellulose/hemicellulloses degradation. These results could provide fundamental insights regarding the mechanisms responsible for different hawthorn fruit textures and lay a solid foundation for further in-depth research.
AUTHOR CONTRIBUTIONS

JX and WD conceived this project and designed the work. JX, YZ, and LZ performed the research. JX, XZ analyzed data, JX, WD, and YH wrote the manuscript.

ACKNOWLEDGMENTS

This work was supported by the Program (project) of “The conservation and utilization of crop germplasm resources-Hawthorn (2014–2016)”.

SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: http://journal.frontiersin.org/article/10.3389/fpls.2016.01524

REFERENCES

Altschul, S. F., Madden, T. L., Schäffer, A. A., Zhang, J., Zhang, Z., Miller, W., et al. (1997). Gapped BLAST and PSI-BLAST: a new generation of protein database search programs. *Nucleic Acids Res.* 25, 3389–3402. doi: 10.1093/nar/25.17.3389

Anders, S., and Huber, W. (2010). Differential expression analysis for sequence count data. *Genome Biol.* 11:R106. doi: 10.1186/gb-2010-11-10-r106

Apweiler, R., Bairoch, A., Wu, C. H., Barker, W. C., Boeckmann, B., Ferro, S., et al. (2004). UniProt: the universal protein knowledgebase. *Nucleic Acids Res.* 32(Database issue), D115–D119. doi: 10.1093/nar/gkh131

Ashburner, M., Ball, C. A., Blake, J. A., Botstein, D., Butler, H., Cherry, J. M., et al. (2000). Gene Ontology: tool for the unification of biology. *Nat. Genet.* 25, 25–29. doi: 10.1038/75556

Atkinson, R. G., Sutherland, P. W., Johnston, S. L., Gunaseelan, K., Hallett, I. C., Mitra, D., et al. (2012). Down-regulation of POLYGALACTURONASE1 alters firmness, tensile strength and water loss in apple (*Malus × domestica*) fruit. *BMC Plant Biol.* 12:129. doi: 10.1186/1471-2229-12-129

Bennett, A. B., and Labavitch, J. M. (2008). Ethylene and ripening-regulated expression and function of fruit cell wall modifying proteins. *Plant Sci.* 175, 130–136. doi: 10.1016/j.plantsci.2008.03.016

Brummell, D. A. (2006). Cell wall disassembly in ripening fruit. *Funct. Plant Biol.* 33, 103–119. doi: 10.1071/FP05234

Brummell, D. A., and Harpster, M. H. (2001). Cell wall metabolism in fruit softening and quality and its manipulation in transgenic plants. *Plant Mol. Biol.* 47, 311–340.

Cao, G. Y., Feng, Y. X., and Qin, X. Q. (1995). Analysis of the chemical constituents of Hawthorn fruits and their quality evaluation. *Acta Pharmacol. Sin.* 30, 138–143.

Cao, L., Zhao, C., Su, S., Luo, C., Han, M., et al. (2014). The role of β-hexosaminidase in peach (*Prunus persica*) fruit softening. *Sci. Hortic.* 169, 226–233. doi: 10.1016/j.scienta.2014.02.015

Chai, W. M., Chen, C. M., Gao, Y. S., Feng, H. L., Ding, Y. M., Shi, Y., et al. (2014). Structural analysis of proanthocyanidins isolated from fruit stone of Chinese hawthorn with potent antityrosinase and antioxidant activity. *J. Agric. Food Chem.* 62, 123–129. doi: 10.1021/jf405385j

Chang, L., Zhang, Z., Yang, H., Li, H., Dai, H., et al. (2007). Detection of strawberry RNA and DNA viruses by RT-PCR using total nucleic acid as a template. *J. Phytopathol.* 155, 431–436. doi: 10.1111/j.1439-0434.2007.01254.x

Chávez Montes, R. A., Ranocha, P., Martinez, Y., Minic, Z., Jouanin, L., Marquis, M., et al. (2008). Cell wall modifications in *Arabidopsis* plants with altered l-arabinofuranosidase activity. *Plant Physiol.* 147, 63–77. doi: 10.1104/pp.107.110023

Chiriboga, M. A., Recasens, I., Schotmans, W. C., Dupille, E., Larrigaudière, C., et al. (2012). Cold-induced changes in ACC metabolism determine softening recovery in 1-MCP treated ‘conference’ pears. *Postharvest Biol. Technol.* 68, 78–85. doi: 10.1016/j.postharvbio.2012.02.006

Costa, F., Peace, C. P., Stella, S., Serra, S., Musacchi, S., Bazzani, M., et al. (2010). QTL dynamics for fruit firmness and softening around an ethylene-dependent polygalacturonase gene in apple (*Malus × domestica* Borkh.). *J. Exp. Bot.* 61, 3029–3039. doi: 10.1093/jxb/erq30

Dai, H. Y., Han, G. F., Yan, Y. J., Zhang, F., Liu, Z., Li, X., et al. (2013). Transcript assembly and quantification by RNA-Seq reveals differentially expressed genes between soft-endocarp and hard-endocarp hawthorns. *PLoS ONE* 8:e72910. doi: 10.1371/journal.pone.0072910

Deng, Y. Y., Li, J. Q., Wu, S. F., Zhu, Y., Chen, Y., and He, F. (2006). Integrated nr database in protein annotation system and its localization. *Comput. Eng.* 32, 71–74.

Eddy, S. R. (1998). Profile hidden Markov models. *Bioinformatics* 14, 755–763. doi: 10.1093/bioinformatics/14.9.755

Finn, R. D., Bateman, A., Clements, J., Coggill, P., Eberhardt, R. Y., Eddy, S. R., et al. (2013). Pfam: the protein families database. *Nucleic Acids Res.* 42(Database issue), D222–D230. doi: 10.1093/nar/gkt1223

Giovannoni, J. (2001). Molecular biology of fruit maturation and ripening. *Annu. Rev. Plant Biol.* 52, 725–749. doi: 10.1146/annurev.arplant.52.1.725

Goulao, L. F., and Oliveira, C. M. (2008). Cloning, characterisation and expression analyses of cDNA clones encoding cell wall-modifying enzymes isolated from ripe apples. *Postharvest Biol. Technol.* 48, 37–51. doi: 10.1016/j.postharvbio.2007.09.022

Goulao, L. F., and Oliveira, C. M. (2008). Cell wall modifications during fruit ripening: when a fruit is not the fruit. *Trends Food Sci. Technol.* 19, 4–25. doi: 10.1016/j.tifs.2007.07.002

Grabherr, M. G., Haas, B. J., Yassour, M., Levin, J. Z., Thompson, D. A., Amit, I., et al. (2011). Full-length transcriptome assembly from RNA-Seq data without a reference genome. *Nat. Biotechnol.* 29, 644–652. doi: 10.1038/nbt.1883

Gwanpua, S. G., Mellidou, I., Boeckx, J., Kyomugasho, C., Bessemansa, N., Verlinden, B. E., et al. (2016). Expression analysis of candidate cell wall-related genes associated with changes in pectin biochemistry during postharvest apple softening. *Postharvest Biol. Technol.* 112, 176–185. doi: 10.1016/j.postharvbio.2015.09.034

Gwanpua, S. G., Van Buggenhout, S., Verlinden, B. E., Christiaens, S., Shipgelman, A., Vicent, V., et al. (2014). Pectin modifications and the role of pectin-degrading enzymes during postharvest softening of Jonagold apples. *Food Chem.* 158, 283–291. doi: 10.1016/j.foodchem.2014.02.138

Harb, J., Gapper, N. E., Giovannoni, J. J., and Watkins, C. B. (2012). Molecular analysis of softening and ethylene synthesis and signaling pathways in a non-softening apple cultivar, ‘Honey crisp’ and a rapidly

FIGURE S1 | Sequence of the amplification products for real-time PCR.

FIGURE S2 | Nr function classification of assembled unigenes. The species distribution is shown relative to the total number of homologous sequences.

FIGURE S3 | KOG function classification of the assembled unigenes. Overall, 12,143 unigenes were grouped into 25 COG classifications.

FIGURE S4 | Differential gene expression profiles for different comparisons. (A) for RRSIVand QJXIV and (B) for RRSV and QJXV. Scatter plot of DEGs (FDR < 0.01 and FC ≥ 2) illustrating the full set of genes sampled. Red points: down-regulated genes; blue points: up-regulated; green points: genes that are not DEGs.

FIGURE S5 | (A) Enriched top GO directed acyclic graph (DAG) of significant DEGs (molecular function) between RRSIV and QJXIV. (B) Enriched top GO directed acyclic graph (DAG) of significant DEGs (molecular function) between RRSIV and QJXV.

TABLE S1 | DEGs related to the cell wall in different textures of ‘Ruannrou Shanlihong 3 Hao’ and ‘Qiu JinXing’ fruits at two developmental stages.

TABLE S2 | Primers for real-time PCR.
softening cultivar, 'McIntosh'. *Postharvest Biol. Technol.* 64, 94–103. doi: 10.1016/j.posthbt.2011.10.001

Hiwasa, K., Nakano, R., Hashimoto, A., Matsuzaki, M., Murayama, H., Inaba, A., et al. (2004). European, Chinese and Japanese pear fruits exhibit differential softening characteristics during ripening, *J. Exp. Bot.* 55, 2281–2290.

Ireland, H. S., Gunaseelan, K., Mudumgane, R., Tacken, E. J., Putterill, J., Johnston, J. W., et al. (2014). Ethylene regulates apple (*Malus × domestica*) fruit softening through a dose–time dependent mechanism and through differential sensitivities and dependencies of cell wall-modifying genes. *Plant Cell Physiol.* 55, 1005–1016. doi: 10.1093/pcp/pct034

Jiménez-Bermúdez, S., Redondo-Navado, J., Muñoz-Blanco, J., Caballero, J. L., López-Arándano, J. M., Valpuesta, V., et al. (2002). Manipulation of strawberry fruit softening by antisense expression of a pectate lyase gene. *Plant Physiol.* 128, 751–759. doi: 10.1104/pp.010671

Kalt, W. (2002). Health functional phytochemicals of fruit. *Hortic. Rev.* 27, 269–315.

Kanehsa, M., Goto, S., Kawashima, S., Okuno, Y., and Hattori, M. (2004). The KEGG resource for deciphering the genome. *Nucleic Acids Res.* 32, 277–280. doi: 10.1093/nar/gkh063

Khallil, H. P. S., Alwani, M. S., and Omar, A. K. M. (2007). Chemical composition, anatomy, lignin distribution, and cell wall structure of Malaysian plant waste fibers. *BioResources* 1, 220–232.

Klee, H. J., and Giovannoni, J. J. (2011). Genetics and control of tomato fruit ripening and quality attributes. *Annu. Rev. Genet.* 45, 41–59. doi: 10.1146/annurev-genet-110410-132507

Koonin, E. V., Fedorova, N. D., Jackson, J. D., Jacobs, A. R., Krylov, D. M., Makarova, K. S., et al. (2004). A comprehensive evolutionary classification of proteins encoded in complete eukaryotic genomes. *Genome Biol.* 5:R7. doi: 10.1186/gb-2004-5-2-r7

Liu, P., Kallio, H., Lü, D., Zhou, C., and Yang, B. (2011). Quantitative analysis of phenolic compounds in Chinese hawthorn (*Crataegus spp.*) fruits by high performance liquid chromatography-electrospray ionisation mass spectrometry. *Food Chem.* 127, 1370–1377. doi: 10.1016/j.foodchem.2011.01.013

Liu, P. Z., Yang, B. R., and Kallio, H. (2010). Characterization of phenolic compounds in Chinese hawthorn (*Crataegus pinnatifida* Bge. var. major) fruit by high performance liquid chromatography-electrospray ionisation mass spectrometry. *Food Chem.* 121, 1188–1197. doi: 10.1016/j.foodchem.2010.02.002

Livak, K. J., and Schmittgen, T. D. (2001). Analysis of relative gene expression using real-time quantitative PCR and the 2−ΔΔCT method. *Methods* 25, 402–408. doi: 10.1016/S1046-323X(01)00423-3

Longhi, S., Hamblin, M. T., Trainotti, L., Peace, C. P., Velasco, R., and Costa, F. (2013). A candidate gene based approach validates Md-PG1 as the main responsible for a QTL impacting fruit texture in apple (*Malus × domestica* Borkh). *BMC Plant Biol.* 13:37. doi: 10.1186/1471-2229-13-37

Lunn, D., Phan, T. D., Tucker, G. A., and Lyckett, G. W. (2013). Cell wall composition of tomato fruit softening changes during development and inhibition of vesicle trafficking is associated with reduced pectin levels and reduced softening. *Plant Physiol. Biochem.* 66, 91–97. doi: 10.1016/j.plaphy.2013.02.003

Martínez, G. A., and Civello, P. M. (2008). Effect of heat treatments on the softening characteristics during ripening of 'McIntosh' softening cultivar. *J. Exp. Bot.* 59, 85–90. doi: 10.1093/jxb/ert283

Mercado, J. A., Pliego-Alfaro, F., and Quesada, M. A. (2011). “Fruit shelf life and potential for its genetic improvement,” in *Edible Genomes: Utilizing Novel Genomes to Develop Novel Foods*, eds M. A. Jenks and P. J. Bebeli (Oxford: John Wiley & Sons, Inc.), 12ñ17.

Mercado, J. A. (2013). Insights into the effects of polygalacturonase FaPG1 gene silencing on pectin matrix disassembly, enhanced tissue integrity, and firmness in ripe strawberry fruits. *J. Exp. Bot.* 64, 3803–3815. doi: 10.1093/jxb/ert210

Qiu, Z., Wan, L., Chen, T., Wan, Y., He, X., Lu, S., et al. (2013). The regulation of cambial activity in Chinese fir (*Cunninghamia lanceolata*) involves extensive transcriptome remodeling. *New Phytol.* 199, 708–719. doi: 10.1111/nph.12301

Ranganna, S. (2001). “Sugar estimation,” in *Handbook of Analysis and Quality Control for Fruit and Vegetable Products*, 2nd Edn, ed. S. Ranganna (New Delhi: Tata McGraw-Hill), 12–17.

Sañudo-Barajas, J. A., Labavitich, J., Greve, C., Osuna-Enciso, T., Muy-Rangel, D., and Siller-Cepeda, J. (2009). Cell wall disassembly during papaya softening: role of ethylene changes in composition, pectin-derived oligomers (PDOs) production and wall hydrolases. *Postharvest Biol. Technol.* 51, 158–167. doi: 10.1016/j.posthbt.2008.07.018

Shin, H., Hong, S. J., Kim, H., Yoo, C., Lee, H., Choi, H. K., et al. (2015). Elucidation of the growth delimitation of Dunaliella tertiolecta under nitrogen stress by integrating transcriptome and peptidome analysis. *Bioresour. Technol.* 194, 57–66. doi: 10.1016/j.biortech.2015.07.002

Takeda, T., Nakano, Y., Takahashi, M., Konno, N., Sakamoto, Y., Arashida, R., et al. (2015). Identification and enzymatic characterization of an endo-1,3-β-glucanase from *Euglena gracilis*. *Phytochemistry* 116, 21–27. doi: 10.1016/j.phytochem.2015.05.010

Tatusov, R. L., Galperin, M. Y., Natale, D. A., and Koonin, E. V. (2000). The COG database: a tool for genome-scale analysis of protein functions and evolution. *Nucleic Acids Res.* 28, 33–36. doi: 10.1093/nar/28.1.33

Trappnell, C., Williams, B. A., Pertea, G., Mortazavi, A., Kwan, G., van, Baren M. J., et al. (2010). Transcript assembly and quantification by RNA-Seq reveals unannotated transcripts and isoform switching during cell differentiation. *Nat. Biotechnol.* 28, 511–515. doi: 10.1038/nbt.1621

Verma, S. K., Jain, V., Verma, D., and Khamesra, A. (2007). *Crataegus oxyacantha*-a cardioprotective herb. *J. Herb. Med. Toxicol.* 1, 65–71. doi: 10.1155/2013/149363

Vicente, A. R., Saladie, M., Rose, J. K. C., and Labavitich, J. M. (2007). The linkage between cell wall metabolism and fruit softening: looking to the future. *J. Sci. Food Agric.* 87, 1435–1448. doi: 10.1002/jsfa.2837

Wei, J., Qi, X., Cheng, Y., and Guan, J. (2015). Difference in activity and gene expression of pectin-degrading enzymes during softening process in two cultivars of Chinese pear fruit. *Sci. Hortic.* 197, 434–440. doi: 10.1016/j.scienta.2015.10.002
Woolley, L. C., James, D. J., and Manning, K. (2001). Purification and properties of an endo-β-1, 4-glucanase from strawberry and down-regulation of the corresponding gene, cel1. *Planta* 214, 11–21. doi: 10.1007/s004250100577

Wu, J., Peng, W., Qin, R., and Zhou, H. (2014). *Crataegus pinnatifida*: chemical constituents, pharmacology, and potential applications. *Molecules* 19, 1685–1712. doi: 10.3390/molecules19021685

Yang, B., and Liu, P. (2012). Composition and health effects of phenolic compounds in hawthorn (*Crataegus* spp.) of different origins. *J. Sci. Food Agric.* 92, 1578–1590. doi: 10.1002/jsfa.5671

Yoshioka, H., Hayama, H., Tatsuki, M., and Nakamura, Y. (2011). Cell wall modifications during softening in melting type peach “Akatsuki” and non-melting type peach “Mochizuki”. *Postharvest Biol. Technol.* 60, 100–110. doi: 10.1016/j.postharvbio.2010.12.013

Zhang, G., Guo, G., Hu, X., Zhang, Y., Li, Q., Li, R., et al. (2010). Deep RNA sequencing at single base-pair resolution reveals high complexity of the rice transcriptome. *Genome Res.* 20, 646–654. doi: 10.1101/gr.100677.109

**Conflict of Interest Statement:** 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.