Mitochondria: Key Organelles Accelerating Cell Wall Material Accumulation in Juice Sacs of Pummelo (Citrus grandis L. Osbeck) Fruits during Postharvest Storage

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1. Introduction

Citrus is an important fruit tree crop around the world. Juice sac granulation of citrus fruits is a severe physiological disorder during the late growing season and postharvest storage of citrus fruits. This disorder is characterized by an increase in firmness and a decrease in the flavour quality of juice sacs. A high rate of juice sac granulation leads to severe decline not only in fruit quality but also in the economic effectiveness of the citrus industry. Granulation rates of citrus fruits vary widely. For lime, orange, and mandarin, they range from 10% to over 80% [1, 2].

Studies have demonstrated that a secondary cell wall is formed in juice sacs during citrus fruit granulation [3, 4]. Also, sugar contents decrease in granulated juice sacs [5, 6]. Moreover, the abundance of organic acids varies strongly.

Citric acid and isocitric acid contents decrease but malic acid levels increase in granulated juice sacs of “Guanxi” pummelo fruits [7].

Transcriptome data of granulating juice sacs of ponkan fruits provided evidence explaining the changes in carbohydrate contents: the expression levels of genes encoding enzymes involved in the sugar and citric acid degradation pathways significantly increased, while transcript levels of enzymes that participate in the synthesis pathways of sugar and citric acid greatly decreased [8]. Phenols are precursors of lignin [9], which decrease in juice sacs during granulation development of citrus fruits [3, 10]. Lignin accumulation is positively associated with the degree of juice sac granulation [11]. Besides, cellulose and hemicellulose contents also exhibit increasing trends during granulation [12]. Phenylalanine ammonia lyase (PAL) and cinnamyl alcohol
dehydrogenase (CAD) are enzymes involved in the lignin biosynthesis pathway. Increases in the activities and transcript levels of the two enzymes improve lignin accumulation [13–15]. Reports demonstrated that elevated PAL and CAD activities were accompanied by an increase in the degree of granulation of pummelo fruits [11]. In addition, activities of enzymes that participate in the cell wall degradation pathway declined in granulated juice sacs of pummelo fruits, including pectin methylesterase, polygalacturonase, and cellulase [12]. These findings indicate that sugar and organic acid degradation may be connected with cell wall material synthesis in granulated juice sacs.

It has been demonstrated that various critical factors may be involved in the regulation of juice sac granulation: hydrogen peroxide (H$_2$O$_2$) and abscisic acid are positively associated with granulation of pummelo fruits, while auxin is negatively linked [16–18]. Three MYB transcription factors (CsMYB330, CsMYB308, and CsMYB85) regulate the expression level of Cs4CL1, a gene encoding for a 4-coumaric acid coenzyme A ligase. CsMYB330 and CsMYB308 can recognize and bind AC elements in the Cs4CL1 promoter and act as a transcription activator and a transcription suppressor, respectively. Additionally, CsMYB85 binds the CsMYB330 promoter and thereby regulates CsMYB330 expression levels. Also, CsMYB85 overexpression leads to higher Cs4CL1 expression levels and higher lignin accumulation in juice sacs of sweet orange [19, 20]. These findings are beneficial for explaining the juice sac granulation mechanism; however, the organization of these factors in signaling networks is still poorly understood.

Most metabolic pathways are closely associated with specific organelles in plant cells. For instance, proteins involved in the tricarboxylic acid cycle, oxidative phosphorylation, and biosynthesis of secondary metabolites can be isolated from mitochondria in the pulp of ripening citrus fruits from satsuma mandarin, ponkan mandarin, sweet orange, and pummelo [21]. Also, a high concentration of cinnamate-4-hydroxylase (an enzyme of lignin synthesis) was found in Golgi bodies in differentiating hypocotyls of French bean (Phaseolus vulgaris L.) [22], and cellulose synthases were shown to be anchored on the plasma membrane in tobacco (Nicotiana tabacum L.) plants [23]. UDP-xyllose synthase, an enzyme that catalyzes the conversion of UDP-glucuronic acid to UDP-xyllose, is localized in the cytoplasm and at the Golgi apparatus in different cell types of Arabidopsis thaliana [24]. Before UDP-xyllose synthase complexes are accumulated at the Golgi apparatus, their subunits are assembled at particular areas of the endoplasmic reticulum in etiolated wheat (Triticum aestivum) seedlings [25]. Besides, plasma membrane and mitochondria are two important sites for ROS generation [26, 27]. Studying this targeting of sugar metabolism components, cell wall substances, and H$_2$O$_2$ specific organelles or cell structures may provide further evidence to reveal the mechanism of citrus fruit granulation.

A previous report showed that the middle and stylar-end sections of the fruit segment are most sensitive to granulation in “Magallanes” pummelo [Citrus maxima (Burm. ex Rumph.) Merr.] [28]. In this study, we combined this information with our previous investigation of the granulation sequences of juice sacs of “Guanxi” pummel fruit. We classified juice sacs into juice sacs isolated from the dorsal vascular bundles (light granulation degree) and juice sacs isolated from the septal vascular bundles (heavy granulation degree). Based on the classification of juice sacs, the aim of this study is to reveal the important connection between ultrastructure dynamics and carbohydrate fluctuation in juice sacs of “Guanxi” pummel fruits during postharvest storage at room temperature.

2. Materials and Methods

2.1. Fruit Samples. We harvested 180 ripe fruits from ten “Guanxi” pummelo (Citrus grandis L. Osbeck) trees (15-year-old) from a commercial orchard located in Zhangzhou City, China, on October 30, 2016. Eighteen medium-sized fruits were selected from the outside canopy (fruits grown under similar light and air conditions) of each tree. Fruits were wrapped with polyethylene to prevent water loss, divided into three biological replicates, and stored for 60 days at room temperature.

Three fruits were randomly selected from each biological replicate at 0, 10, 20, 30, 40, 50, and 60 days. Juice sacs isolated from the dorsal vascular bundles (dorsal juice sacs) and juice sacs isolated from the septal vascular bundles (septal juice sacs) were collected separately (Figure 1). Juice sac samples collected from each biological replicate were mixed and stored at −80°C for the determination of cell wall materials, soluble sugar, and organic acid contents. Additionally, two types of flesh juice sacs were separately isolated from fruits stored for 0, 10, 20, 40, and 60 days. Flesh juice sacs were selected from one fruit that perfectly represented the granulation degree of fruits from three biological replicates at each stage. Approximately 2 mm long sections were dissected from juice sacs isolated from the dorsal or the septal vascular bundles and immediately fixed in 3.5% glutaraldehyde in 0.2% phosphate buffer (pH 7.2) for two days at 4°C.

2.2. Transmission Electron Microscopy (TEM). The fixed samples were rinsed with 0.2% phosphate buffer three times for 15 min each. Samples were then postfixed with a solution containing 1% osmic acid and 1.5% potassium ferrocyanide for 1.5 h at 4°C. Following the postfixation, samples were again rinsed with 0.2% phosphate buffer five times for 15 min each. The rinsed samples were dehydrated in a graded series of ethanol and acetone: 50% ethanol for 15 min, 70% ethanol overnight, 90% ethanol for 15 min, a mixed solution of 90% ethanol and acetone (v/v, 1:1) for 15 min, 90% acetone for 15 mins, and anhydrous acetone for 15 min. Dehydrated samples were infiltrated in a mixture of anhydrous acetone and epoxy resin 618 (v/v, 1:1) for 1.5 h and in pure epoxy resin 618 for 3 h at 35°C. The samples were embedded in pure epoxy resin 618 and allowed to polymerize at 35°C for 12 h, at 45°C for 12 h, and at 60°C for 2-3 h. The embedded tissues were cut into ultrathin sections (90 nm) with a glass knife on an ultramicrotome (Leica EM
2.3. Lignin Determination. The detection of lignin was
Carried out according to the method described by Bomal
Et al. [29] with some modifications. Frozen juice sacs were
Ground in liquid nitrogen, and 1.5 g powder was placed in
A 10 mL tube and extracted with 6 mL 80% ethanol at 80°C for
30 min to remove sugar, acid, and pigment and then
centrifuged at 15,000 g for 10 min at 4°C. The residues were
Suspected in 80% ethanol, and the extraction was repeated
twice. The pellet was resuspended with 6 mL 80% methanol
And kept at room temperature for 30 min, followed by
centrifugation at 15,000 g for 10 min at 4°C. The supernatant
Was removed, and the residues were extracted with methanol
twice. Next, the pellet was dissolved in 6 mL n-hexane and
And kept at room temperature for 30 min to further remove
Pigment and lipid. This extraction was repeated twice. After
centrifuging at 15,000 g for 10 min at 4°C, the final pellet was
dried in oven at 80°C.

10 mg of the final pellet was transferred into 5 mL glass
tube, and 1 mL 25% acetyl bromide (dissolved in glacial
acetic acid) was added. The sample solution was incubated at
50°C for 2 h. 2 mL of 2 M NaOH solution and 5 mL glacial
acetic acid was added to a 25 mL volumetric flask. The
Extracted solution was added, followed by 0.35 mL 0.5 M
Hydroxylamine. Finally, the mixed solution was filled up to
25 mL with glacial acetic acid. The absorbance of lignin was
detected using an ultraviolet spectrophotometer at 280 nm.

2.4. Cellulose and Hemicellulose Determination. The analyses
Of cellulose and hemicellulose content were performed using
The method described by Ookawa et al. [30] with slight
Modifications. Flesh samples were weighed and recorded as
W0. Removal of sugar, acid, pigmented, and lipid from the
Sample was performed as described for the lignin deter-
mination. The dried pellet was weighed and recorded as W1.
20 mg of the dried pellet was digested in 1 mL 15 U·mL⁻¹ α-
amylase solution at 85°C for 30 min to remove starch, fol-
dowed by centrifugation at 15,000 g for 10 min at 4°C. The
Supernatant was discarded, and the residue was dried at
80°C. The dry pellet was mixed with 650 μL ultrapure water,
6 mg CINaO₃, and 5 μL glacial acetyl acid and subsequently
placed in water for 1 h at 80°C to remove lignin. These steps
were repeated twice. Next, the pellet was washed with ul-
papse:ure water and dried at 80°C. The delignified pellet was
resuspended with 1 mL neutral detergent solution (18 mM
Sodium tetaborate decahydrate, 66 mM ethyl-
enediaminetetraacetic acid, 10.4 mM sodium dodecyl sul-
fate, 32 mM dibasic sodium phosphate, and 1% (v/v)
Triethylene glycol) and boiled in water for 1 h to remove
Soluble substance and pectin. After centrifugation, the
remaining pellet was washed with ultrapure water and dried
at 80°C. The pellet (holocellulose) weight was recorded as
W2. The dried pellet was digested in 1 mL 1 M H₂SO₄ so-
lution for 1 h at 100°C. Then, 1 mL 2 M H₂SO₄ solution was
added, and the sample was boiled for 1 h to remove
Hemicellulose. Ultimately, the pellet was washed in ultrapure
water and dried at 80°C. The final pellet (cellulose) was
weighed (W3). The contents of cellulose and hemicellulose
In flesh samples were calculated with the following formulas:
Cellulose (mg·g⁻¹ FW) = W3 × (W1/20)/W0. Hemicellulose
(mg·g⁻¹ FW) = (W2 − W3) × (W1/20)/W0, FW means flesh
Weight of sample, and 20 is the initial weight (20 mg) of dry
pellet used for cellulose and hemicellulose determination.

2.5. Extraction and Quantification of Soluble Sugars and
Organic Acids. The samples were prepared according to the
Method described by Niu et al. [31] with some modifications.
Frozen juice sacs (about 5 g fresh weight) were ground in
Liquid nitrogen. Then 1.2 g of fine powder was transferred
Into a 10 mL tube and extracted in 6 mL 80% ethanol for
30 min at 37°C. Samples were centrifuged at 15,000 g for
6 min at 4°C. The residues were reextracted twice. The
Combined supernatants were placed in a volumetric flask
And diluted to 25 mL with 80% ethanol. 3 mL of homoge-
nized solution was dried at 60°C using a vacuum rotary
evaporator. The dried extracts were dissolved in 3 mL ul-
trapure water, filtered through a 0.45-μm hydroponic
membrane, stored at −80°C, and used to determine soluble sugar. Meanwhile, 5 mL of homogenized solution was dried using the same method, dissolved in 1 mL ultrapure water, and then passed through a 0.45 μm hydroponic membrane. This filtered solution was used for detecting the content of organic acid.

Determination of soluble sugar content was performed using ultra-high-performance liquid chromatography (UltiMate 3000, Dionex, California, USA), equipped with an ODS-NH2 column (4.6 × 250 mm, 5 μm) and CAD detector. 10 μL of filtered solution of each sample was separated on a column at a temperature of 35°C and eluted with 70% acetonitrile at a flow rate of 1 mL min⁻¹. D (+)-Sucrose, D-Fructose, and D (+)-Glucose were used as standard sugars (Solarbio, Beijing, China).

Quantification of organic acid was carried out using the same instrument. The chromatographic conditions are as follows [32]. 10 μL of filtered solution of each sample was loaded on RD-C18 columns (4.6 × 250 mm, 5 μm) at 35°C and separated with mobile solution (1.8 mM KH2PO4, pH adjusted to 2.25 with phosphoric acid) at 0.8 mL min⁻¹ flow rate. The absorbance of organic acids was recorded with an ultraviolet detector at 210 nm wavelength. Citric acid and malic acid standards were purchased from Solarbio, Beijing, China.

2.6. Statistical Analysis. Data were expressed as mean ± standard error from three biological replicates. Differences between the two types of juice sacs were compared with Student’s t-test at p < 0.05. Multiple comparisons between stages were performed with one-way ANOVA based on Duncan’s multiple range test at p < 0.05. Statistical analyses were carried out with the SPSS 16.0 software.

3. Results

3.1. Changes in the Ultrastructure of Juice Sacs. In order to provide anatomic proof for juice sac granulation mechanisms, the subcellular structures of juice sacs were investigated during postharvest storage at room temperature. At the beginning of the storage period, the cytoplasm was uniformly distributed in the intracellular spaces of dorsal juice sacs (Figure 2(a)). The plasma membranes showed no damage. Particularly, a bright region was observed in the space between the plasma membrane and cell wall (Figure 2(b)). In septal juice sacs, the cytoplasm was distributed along the cell walls (Figure 2(c)), while the plasma membranes partially ruptured but vesicles were empty (Figure 2(d)).

After ten days of storage, organelles were distributed along the cell walls (Figure 3(a)) in dorsal juice sacs. Dense substances were scattered in the intracellular spaces (Figure 3(b), arrowhead) and also attached to the surface of vesicles adjacent to mitochondria. Particularly, the plasma membrane partially ruptured and the bright spaces between the plasma membrane and the cell walls disappeared (Figure 3(c)). In septal juice sacs, cytoplasm more closely attached to cell walls after ten days compared to the previous stage (Figure 3(d)). The plasma membranes were partially pulled away from the cell walls (Figure 3(d), open arrow). Some vesicles began accumulating dense substances; meanwhile, sparse dense substances were distributed in the intracellular spaces (Figures 3(e) and 3(f)).

At 20 days of storage, striking differences in structural characteristics were observed between the two types of juice sacs. In dorsal juice sacs, vesicles of various sizes appeared in the intracellular spaces; also, mitochondrion fission caused a slight increase in the number of mitochondria (Figures 4(a), 4(b), and 4(d)). The damage to the plasma membranes became severe, but the amount of cell wall substances was not obviously increased (Figure 4(c)). In septal juice sacs, numerous mitochondria and vesicles appeared in the intracellular spaces. Besides, a large number of small, filled vesicles were secreted from the Golgi apparatus adjacent to the mitochondria (Figures 4(e) and 4(f)). The plasma membranes ruptured and degraded (Figure 4(g)). At this stage, cell wall materials were distributed throughout the intracellular spaces and intracellular spaces (Figures 4(e), 4(f), 4(g) black arrowed head).

At 40 days of storage, mitochondria were prominent in the intracellular spaces of dorsal juice sacs (Figure 5(a)). Many small vesicles were observed concurrent with the mitochondria, but the Golgi apparatus did not appear in this type of juice sac (Figure 5(b)). The plasma membranes ruptured and degraded continuously (Figure 5(c)). It can be found that cell wall materials were uniformly distributed in the intracellular spaces (Figures 5(b) and 5(c)). In contrast, the number of mitochondria decreased, and they moved towards the cell walls in septal juice sacs (Figure 5(c)). Vesicles almost disappeared but long microfibrils accumulated in the intracellular spaces (Figure 5(d)). Cell wall materials were deposited on the surface of the cell walls (Figure 5(f), black arrowhead).

At the end of the storage period, mitochondria and vesicles disappeared in cells of both types of juice sacs (Figures 6(a) and 6(c)), and cell wall materials were continuously deposited on the surface of the cell walls (Figures 6(b) and 6(d)). Small black particles in the intracellular spaces in cells of dorsal juice sacs demonstrated that the cell wall components in this type of juice sac were different from those in septal juice sacs (Figure 6(b), black arrowhead).

Additionally, in both types of juice sacs, cell wall substances accumulated in the intracellular spaces throughout the whole period of storage.

3.2. Dynamics of Cell Wall Material Contents in Juice Sacs. In dorsal juice sacs, lignin contents showed no significant changes from 0 to 30 days of storage, peaked at 40 days (p < 0.05), and then decreased to similar levels as detected before day 30. Similarly, lignin contents were relatively steady in septal juice sacs from 0 to 30 days; however, they rapidly increased from 30 to 50 days, leading to significantly increased lignin contents at 40 and 50 days. Until day 60, lignin levels decreased, as seen before in dorsal juice sacs (Figure 7(a)).
The above results show a similar trend in the fluctuation of lignin contents in both types of juice sacs during post-harvest storage. However, lignin contents in septal juice sacs were 4-fold, 2-fold, and 7-fold higher than those in dorsal juice sacs at 30, 40, and 60 days, respectively.

Cellulose contents in dorsal juice sacs remained relatively steady during postharvest storage. In contrast, in septal juice sacs, cellulose levels sharply increased at 40 days and remained high at 50 and 60 days of storage. Cellulose contents in septal juice sacs were significantly higher than those in dorsal juice sacs at 30 to 60 days (Figure 7(b)).

In dorsal juice sacs, hemicellulose was significantly accumulated at 10 and 60 days and has no apparent changes in the rest of stages. Hemicellulose in septal juice sacs remained relatively steady during postharvest storage but was significantly higher (1.6-fold) than that in dorsal juice sacs at 30 days (Figure 7(c)).

During the whole storage period, fructose contents significantly increased in dorsal juice sacs but showed no significant changes in septal juice sacs. Statistical differences in fructose contents between the two types of juice sacs were investigated at 0, 10, and 30 days (Figure 8(b)).

Glucose levels showed no distinct changes for either type of juice sac during the postharvest storage; also they showed no apparent differences between the two types of juice sacs at each stage (Figure 8(c)).

3.4. Dynamics of Organic Acid Contents in Juice Sacs. Neither citric acid contents nor malic acid contents showed significant changes in either type of juice sac during storage. However, citric acid contents in dorsal juice sacs were much higher than those in septal juice sacs at 10 and 60 days, while malic acid contents in dorsal juice sacs were significantly lower compared to those in septal juice sacs at 40 and 60 days (Figures 9(a) and 9(b)).

4. Discussion

4.1. Two Types of Juice Sacs with Different Degrees of Granulation. In this study, the results showed that cell wall components accumulate differently in two types of juice sacs. Lignin and hemicellulose transiently accumulated in dorsal juice sacs at certain stages of postharvest storage. In
contrast, lignin and cellulose contents significantly increased in septal juice sacs from day 40 of postharvest storage. These results indicate that lignin and hemicellulose might be degraded in dorsal juice sacs. Cellulose contents in dorsal juice sacs and hemicellulose contents in septal juice sacs showed no significant differences during postharvest storage. Lignin, cellulose, and hemicellulose contents in septal juice sacs began to significantly exceed those of dorsal juice sacs at 30 days of storage. However, in contrast to cellulose and lignin, the significant differences in hemicellulose levels between the two types of juice sacs disappeared after 30 days. Taken together, these findings suggest that lignin and cellulose are the main contributors to the difference in the granulation degree of the two types of juice sacs.

4.2. Changes in Soluble Sugar Contents Are Connected with Cell Wall Material Accumulation. In septal juice sacs, sucrose contents significantly decreased from 30 to 60 days, while lignin and cellulose were primarily accumulated after 30 days. This suggests that sucrose degradation starts slightly before cell wall material accumulation. Sucrose can provide the primary components for cellulose and lignin synthesis. Invertase and sucrose synthase are two distinct enzymes that

Figure 3: Transmission electron micrographs of cross sections of dorsal juice sac (a–c) and septal juice sac (d–f) of pummelo fruit stored for ten days at room temperature. (a) Overview of the cell structure of a dorsal juice sac. (b, c) Magnified micrographs of the regions marked by rectangles in (a). Note the distribution pattern of organelles and the occurrence of cell wall materials in the intracellular spaces. (d) Overview of the cell structure of a septal juice sac. (e, f) Higher magnification micrographs of the regions marked by rectangles in (d). Note cell wall materials distributed in the intracellular space and filled vesicles. Cw: cell wall, M: mitochondrion, V: vesicle, Is: intercellular space, open arrow: plasma membrane, and black arrowhead: dense substances.
catalyze the conversion of sucrose into UDP-glucose. UDP-glucose is a direct substance for cellulose synthesis [33]. Invertase activity is positively correlated with UDP-glucose and cellulose contents [34–36]. Sucrose synthase is an enzyme that catalyzes the reversible conversion of sucrose and uridine diphosphate (UDP) into fructose and UDP-glucose [37]. In cotton fiber cells, sucrose synthase was found to play a critical role in fiber initiation and early elongation and to be involved in rapid cellulose synthesis at the later development stages [38]. Also, the glycolysis pathway is linked with the polysaccharide and lignin synthesis pathways during protoxylem vessel development in tobacco [39]. Therefore, it is assumable that the carbon source used for lignin synthesis is partially derived from sucrose degradation. In this study, the cellulose and lignin contents increased immediately just after the decline of sucrose contents in septal juice sacs, indicating that sucrose degradation is activated earlier than cellulose synthesis.

In dorsal juice sacs, fructose contents significantly increased from 10 to 60 days, while sucrose and glucose levels did not change. In rice, fructose availability can reverse the sucrose degradation reaction into the sucrose synthesis reaction mediated by sucrose synthase [40]. In this study, increased fructose contents may provide a feedback effect on suppressing sucrose degradation reaction and consequently any significant changes in the sucrose contents can be

Figure 4: Transmission electron micrographs of cross sections of dorsal juice sac (a–d) and septal juice sac (e–g) of pummelo fruit stored for 20 days at room temperature. (a) Overview of the cell structure of a dorsal juice sac. (b–d) Higher magnification micrographs of the regions marked by rectangles in a. Note mitochondrial fission in d. (e) Overview of the cell structure of a septal juice sac. (f and g) Higher magnification micrographs of the regions marked by rectangles in e. Golgi apparatus appeared and the number of mitochondria and vesicles sharply increased in the intracellular space. Cw: cell wall, G: Golgi apparatus, M: mitochondrion, Is: intercellular space, V: vesicle, open arrow: plasma membrane, black arrowhead: dense substances, and white arrowhead: microfibril.
detected in dorsal juice sacs. However, further experiment needs to be carried out to determine the reason for increased fructose in dorsal juice sacs.

4.3. The Possible Roles of Organic Acids Involved in Regulating Juice Sac Granulation. Organic acid contents detected in both types of juice sacs remained relatively steady during the whole storage period. It is well known that malic acid and citric acid are metabolites in tricarboxylic acid cycle [41]. Malate dehydrogenase catalyzes the conversion of malic acid to oxaloacetate [42]. Subsequently, oxaloacetate is used as a substance to produce citric acid [43]. In pummelo fruits, enzymes (H+-ATPase, Ca2+-ATPase, Mg2+-ATPase, cytochrome C oxidase, succinate dehydrogenase, and malate dehydrogenase) that participated in energy metabolism were markedly maintained by chitosan coating, which was accompanied by decreased cellulose content in pulp, alleviated postharvest senescence, and reduced energy depletion of fruits [44]. Based on findings reported by Chen et al. [44], it can be deduced that markedly maintained malate dehydrogenase in pummelo fruits may act as a mediator for regulation of malic acid and citric acid contents. The findings in this study combined with the reports suggest that malic acid and citric acid contents remain constant as much as possible in response to juice sac granulation.
4.4. Plasma Membrane Rupture Induces Cell Wall Material Synthesis. In this study, the damage to plasma membrane is accompanied by cell wall material accumulation in juice sacs. Lipid peroxidation leads to decreased plasma membrane integrity and can be detected by malonaldehyde and H$_2$O$_2$ production [45, 46]. Considering the finding that H$_2$O$_2$ is accumulated in granulated juice sacs [16], combined with the ultrastructure obtained in this study, it appears that the loss of plasma membrane integrity may act as an early signal for juice sac granulation.

4.5. An Increase in the Number of Mitochondria Accelerates Cell Wall Material Accumulation. After ten days of storage, plasma membrane ruptures continuously increased for the rest of the storage period. However, this is not the key factor inducing considerable accumulation of cell wall materials and noticeable decreases in soluble sugar contents. This conclusion is supported by the findings that TEM images of dorsal juice sacs showed an increased number of vesicles and damage of the plasma membranes at day 20, which did not lead to significant increases in cell wall material contents at this stage. Until the number of mitochondria sharply increased at 40 days, cell wall substance was accumulated mainly in dorsal juice sacs. Furthermore, in septal juice sacs, dramatical increase in the number of mitochondria was accompanied by distinct distribution of cell wall materials. Additionally, lignin, cellulose, and hemicellulose contents in septal juice sacs began to significantly exceed those of dorsal juice sacs at 30 days of storage (just after the number of mitochondria greatly increased at 20 days), providing further proof of the key role of mitochondria in regulating carbon source partitioning to cell wall material synthesis.

In this study, the Golgi apparatus and/or numerous vesicles occurred simultaneously with the number of mitochondria in juice sacs at specific stages. In line with this finding, many reports have shown that cell wall material accumulation, at least in part, depends on the Golgi apparatus and vesicles. Suppressing vesicle secretion from the Golgi apparatus, cellulose contents decrease in whole plants of rice [47]. Moreover, enzymes or proteins involved in cell wall material synthesis, such as xylan synthase complexes [27], xylan deacetylases [48], STELLO1, 2 (proteins regulating the secretion and activity of cellulose synthase) [49], and cell wall materials like p-glucoumaryl alcohol [50], xylanogluca [51], and xylan [52], have been detected in Golgi apparatus and/or vesicles in various plant tissues (e.g., root, internode, differentiating xylem). In this study, the occurrence of Golgi apparatus and/or numerous vesicles was associated with a high accumulation of cell wall materials in juice sacs, especially lignin and cellulose, which indicates that Golgi and vesicles are also essential organelles involved in the development of juice sac granulation, although their roles link to the occurrence of numerous mitochondria.

Figure 6: Transmission electron micrographs of cross sections of dorsal juice sac (a, b) and septal juice sac (c, d) of pummelo fruit stored for 60 days at room temperature. (a) Overview of the cell structure of a dorsal juice sac. (b) Higher magnification micrograph of the region marked by a rectangle in (a). (c) Overview of the cell structure of a septal juice sac. (d) Higher magnification micrograph of the region marked by a rectangle in (c). Cw: cell wall, Is: intercellular space, white arrowhead: microfibril, and black arrowhead: particles of cell wall substances.
Mitochondria are versatile organelles in plants. It is well known that the tricarboxylic acid cycle, a pathway for ultimate degradation of sugar, acid, and lipid (providing intermediate production and energy), is located in mitochondria [53]. Also, transcriptome profiles revealed that energy metabolism mediated by mitochondria plays a vital role in cotton fiber elongation [54]. Furthermore, it was found that invertases are located at mitochondria in Jerusalem artichoke (Helianthus tuberosus L.) tubers [55]. Knockdown of the genes encoding for alkaline/neutral invertases (A/N-Invs) localized in mitochondria led to reduced oxygen consumption in Arabidopsis Invs mutants [56].

Additionally, oxygen uptake in granulated juice sacs of freshly harvested “Lee” tangelos, stored “Dancy” tangerine, and stored “Marsh” grapefruits was 2- to 3-fold higher when compared with that in normal juice sacs [57]. Combined with our findings that mitochondria are closely connected with evident cell wall material accumulation during juice sac granulation, it can be deduced that mitochondria may play a key role in accelerating sucrose degradation in granulated juice sacs.

On the other hand, mitochondria are also an important site for reactive oxygen species (ROS) synthesis [58–60]. H$_2$O$_2$ has been associated with juice sac granulation and
positively correlated with xylem lignification [22] and cellulose accumulation in plants growing in salt stress conditions [61]. Although the prominent role of H$_2$O$_2$ in cell wall material accumulation is clear, it is uncertain whether H$_2$O$_2$ is generated from mitochondria during juice sac granulation. Therefore, the detailed roles of mitochondria in regulating juice sac granulation need further research. It has been shown that mitochondrial fission and fusion can be imbalanced due to the cell environment, which leads to fluctuations in mitochondrial numbers [62]. Some evidence also demonstrated that ROS accumulation promotes mitochondrial fission in animal cells [63–65]; however, whether ROS participates in regulating mitochondrial dynamics in plant cells is elusive. In the future, suppressing the increase in mitochondria could be a promising approach for controlling juice sac granulation of pummelo fruits.

Figure 8: Soluble sugar contents in dorsal juice sacs and septal juice sacs during postharvest storage. (a) Sucrose contents in juice sacs. (b) Fructose contents in juice sacs. (c) Glucose contents in juice sacs. The x-axis represents the storage time of fruits; the y-axis represents soluble sugar content. Error bars represent the standard errors derived from three biological replicates. The lowercase letters represent the significant differences among stages. Asterisks represent the significant differences between the two types of juice sacs at each stage.
4.6. Cell Wall Material Accumulation in the Intracellular and the Intercellular Space of Juice Sac Cells. In both types of juice sacs, cell wall material accumulation in the intercellular space was earlier compared to that in the intracellular space. However, this did not lead to immediate significant increases in cell wall material contents in juice sacs. These results suggest that the abundance of cell wall materials synthesized in the intracellular space of juice sac cells is closely connected with the degree of juice sac granulation.

5. Conclusions

Cell wall material accumulation in dorsal juice sacs differs from that in septal juice sacs. Lignin and hemicellulose are the major cell wall components synthesized in dorsal juice sacs, and the transient increases in their levels indicate that their degradation reaction should not be ignored. In contrast, lignin and cellulose are the two kinds of cell wall compounds contributing to cell wall substance accumulation in septal juice sacs. The significant increases in their levels begin to be observed at 40 days and prolong to the end of storage. The significant decreases in sucrose contents (at 30 days) prior to the obvious increases in lignin and cellulose levels (at 40 days) in septal juice sacs suggest that sucrose degradation is the key step for providing enough carbon for cell wall component synthesis. In contrast, sucrose contents in dorsal juice sacs remain constant during the postharvest storage. However, fructose contents are significantly elevated from 10 to 60 days. Therefore, it could be deduced that the increased fructose contents may have a feedback effect on suppressing sucrose degradation. Accordingly, cell wall material is less accumulated.

The comprehensive analysis of TEM images of the two types of juice sacs found that, until the number of mitochondria largely increases, cell wall materials are significantly accumulated, although plasma membrane rupture and accumulation of vesicles occur in earlier stages. Taken together, plasma membrane rupture may provide a signal for induction of juice sac granulation, while mitochondria play a critical role in regulating sugar carbon flux partitioning for cell wall material synthesis.

Data Availability

All data used to support the findings of this study are included within the paper.

Conflicts of Interest

The authors declare that they have no conflicts of interest.
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