Comparison of Cell Wall Components in Normal and Disordered Juice Vesicles of Grapefruit

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Abstract. Cell wall composition and structure were examined in visually normal (N), granulated (G), and collapsed (VC) juice vesicles of ‘Marsh Seedless’ grapefruit (Citrus paradisi Macf.). According to gel-filtration data, VC appeared to be associated with a modification of water-soluble (WSP) and chelate-soluble (CSP) pectin molecular weight (\(M_r\)); small-\(M_r\) pectins increased, whereas large-\(M_r\) pectins decreased. The difference in \(M_r\) of pectins did not appear to be mediated by polygalacturonases. Molecular weight of hemicelluloses did not differ. Granulated vesicles contained about two times more structural polysaccharides (pectins, hemicelldose, and cellulose) than N vesicles, although hemicelldose and pectin \(M_r\) modification were absent. Ion-exchange profiles of WSP, CSP, and hemicelldose fractions of VC and G vesicles were not different from those of N vesicles. Individual cells in vesicles with G and these vesicles themselves were much larger than those of N vesicles, whereas cells in VC were partially or completely collapsed.

According to Bartholomew et al. (1941), juice-vesicle disorders in citrus were recognized in the early 1900s. Two basic types of disorders have been distinguished in the literature: a hardening of the vesicles followed by partial dehydration (granulation; G) and the complete collapse of vesicles (vesicle collapse; VC) (Bartholomew et al., 1941; Albrigo et al., 1980). Depending on the fruit and location within the fruit where VC has been found, several names have been used in the literature to describe VC: dry juice sac (Bartholomew et al., 1941), core dryness (Noort, 1969), blossom-end G and stylar-end G (Bartholomew et al., 1941), and VC (Albrigo et al., 1980). We found no detailed studies of these VC disorders, although G has been studied extensively.

Hardening and enlarged juice vesicles are characteristic of G (Bartholomew et al., 1941). Granulation reduces fruit quality due to a decrease in sugar and acid content (Awasthi and Nau-
riyal, 1972; Bartholomew et al., 1941; Chakravar and Singh, 1977a, 1977b; El-Zeftawi, 1973, 1978; Gilfillan and Stevenson, 1977; Matsumoto, 1964; Sinclair and Jolliffe, 1961). Granulated vesicles contain more ethanol-insoluble solids (EIS) (Sinclair and Jolliffe, 1961), which are largely composed of cell walls (Bartholomew et al., 1941), than do N vesicles. Among the cell-wall components, pectins in particular have been proposed to be a potential factor associated with the hardening of G vesicles through gelation (Bartholomew et al., 1941; Sinclair and Jolliffe, 1961). However, no studies have been conducted to compare detailed changes in these and other cell-wall components of G vesicles.

Both G and VC occur in grapefruit, but usually VC occurs more commonly than G (Hwang et al., 1988). Since visible symptoms of VC differ from G, we thought that VC in grapefruit might occur via a mechanism unlike that responsible for G. The present study focused on comparing the biochemical differences characteristic of G and VC in grapefruit, especially regarding the structural and compositional features of cell-wall polysaccharides, to see if similarities existed between the two vesicle disorders.

Materials and Methods

Plant materials. Visually normal and disordered juice vesicles were isolated from disordered 'Marsh Seedless' grapefruit at harvest or from fruit stored at 21°C and 90% to 95% RH for 40 days. Adjacent juice vesicles that showed no visible symptoms of G or VC were regarded as N vesicles for comparison. A seasonal study from August to March (unpublished data) indicated that no major EIS or wall structural component changes occurred in juice vesicles in healthy nondisordered fruit. Vesicles were placed in an ice bath as they were isolated and were quickly frozen at –40°C and then stored in polyethylene bags at –20°C. Each replication, usually three, of vesicles for a given disorder and its adjacent apparent normal vesicles was collected from fruit of the same harvest date and grove source. Standard deviations are presented for the values in these paired comparisons.

Preparation of ethanol insoluble solids. Ethanol-insoluble solids were prepared basically as described by Huber (1984), with slight modification. Vesicle tissue (45 g) was homogenized in 190 ml 95% ethanol for 3 min. The homogenate was reflushed for 20 min in a boiling water bath and then stored overnight at –20°C. The suspension was filtered through Miracloth (Biochemical Corp., La Jolla, Calif.) and washed with 250 ml of 80% ethanol (v/v), followed by 250 ml of 80% acetone (v/v) and 400 ml of 100% acetone. After evaporating the excess acetone, EIS were dried overnight at 35°C and stored in a desiccator at room temperature.

Determination of total pectins and cellulose. Total polyuronides in EIS were measured by the method of Ahmed and Labavitch (1977). Chilled sulfuric acid (2.5 ml) was added to 5 to 6 mg EIS in an ice bath. After each of two 5-rein intervals, 0.7-ml aliquots of deionized (DI) water were added and the suspensions were stirred. After an additional 5 rein, 35 ml of DI water were added and aliquots (0.5 ml) analyzed for total uronic acid content (Blumenkrantz and Asboe-Hansen, 1973). Cellulose content in EIS was determined as described by Updegraff (1969). Twenty milligrams of EIS were placed in 5 ml of 4 acetic acid :1 nitric acid :2 H₂O (by volume) and placed in a boiling water bath for 30 min. The suspension was centrifuged in a clinical centrifuge for 10 min at 490x g. The pellet was washed with 10 ml of DI water (2 ×) and centrifuged. The pellet material was then transferred to 67% sulfuric acid and maintained at room temperature for 1 hr. The hydrolysate was diluted by transferring to 25 ml of distilled water. Total hexoses were measured by the anthrone procedure (Hedge and Hofreiter, 1962).

Isolation and determination of soluble polyuronides and hemicelluloses. Water-soluble pectins (WSP) were isolated by suspending EIS (100 mg) in 35 ml of DI water. The suspension was placed on a shaker for 6 hr at room temperature. After 6 hr, the suspension was filtered through Miracloth and washed with 10 ml of DI water. The combined filtrates were further filtered through glass-fiber filters (Whatman G/FC) under aspiration. The filtrates were stored at –20°C until used. After extraction of water-soluble polyuronides, chelator-soluble pectins (CSP) were extracted by placing the recovered powder in 35 ml of Na-acetate (40 mM pH 5.0) containing 20 mM NaEDTA. The subsequent methods were as described for WSP.

Before extraction of hemicelluloses, polyuronides were removed from the EIS by placing 100 mg of EIS in 150 ml of Na-phosphate (100 mM, pH 6.5) containing 20 mM NaEDTA. The suspension was then placed in a boiling water bath for 30 min. Afterwards, the suspension was filtered through Miracloth, washed with 150 ml of extraction buffer, followed by 500 ml of DI water. Hemicelluloses were extracted by suspending the residual powder in 15 ml of 4 N NaOH containing 6 mg NaBH₄/ml. After two successive 4-hr extractions, the alkali-soluble fractions were filtered through Miracloth. The combined filtrates were neutralized with concentrated acetic acid and dialyzed against running tap water (12 hr, room temperature), 4 liters of 10% methanol (2 × 12 hr, 4°C), and finally with 4 liters of DI water (2 × 12 hr, 4°C). The dialyzed samples were filtered through GF/C filter paper. The filtrates were stored at –20°C. Sugar content of the alkali extracts was measured by the phenol-sulfuric acid method (Hedge and Hofreiter, 1962) using D-galactose as the standard.

Gel-filtration chromatography of soluble polyuronides and hemicelluloses. Gel-filtration of polyuronides was conducted on a bed (60 cm long × 1.5 cm wide) of Ultrogel AcA 22 (Bio-Rad, Richmond, Calif.) packed in Na-acetate buffer (25 mM, pH 5.0) containing 3 mM NaEDTA and 100 mM NaCl. About 2 mg polyuronide in 2 ml of extraction medium (water or buffer) were applied to the column and eluted at a flow rate of 17.5 ml·cm⁻²·hr⁻¹. Aliquots (0.5 ml) of the 2-ml fractions were analyzed for total uronic acid content (Blumenkrantz and Asboe-Hanson, 1973).

Hemicelluloses (≈4 mg) in 2 ml of the 30 mM acetate buffer were applied to a bed (60 cm long x 1.5 cm wide) of Ultrogel AcA 34 operated in the same buffer system. Hemicelluloses were eluted at a flow rate of 63 ml·cm⁻²·hr⁻¹. Fractions of 2 ml were collected and 0.5-ml aliquots were assayed for hexose by the phenol–sulfuric acid procedure.

Ion-exchange chromatography. Samples were subjected to ion-exchange chromatography before neutral sugar analysis of polyuronides. Ion-exchange chromatography was carried out on a bed (16 cm long × 1.5 cm wide) of DEAE-Sephadex (Sigma, St. Louis) that was equilibrated and packed in Na-phosphate buffer (15 mM, pH 6.8) containing 20 mM NaCl and 5 mM NaEDTA. Following dialysis (Spectrapor 2000) against the column buffer, 8 mg of polyuronides were applied. After 100 ml of starting buffer was collected, the column was step-eluted with the same buffer containing 0.5 mM NaCl. Polyuronides eluted by 0.5 M NaCl were recovered and then dialyzed against DI water for 24 hr at 4°C. Higher NaCl concentrations eluted no additional pectin.

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Neutral sugar analysis. Polyuronide (1 mg) was air-dried in Reacti-Vials (Pierce Chemical Co., Rockford, Ill.) and hydrolyzed and acetylated as described by Loescher and Nevins (1972). For compositional analyses of hemicelluloses, ≈ 8 mg was applied to the DEAE-Sepharose column operated as described above. Polymers recovered in the column wash (unbound) were collected and subjected to hydrolysis and acetylation. The derivatized neutral sugars were examined by gas chromatography on a glass column packed with 3% SP 2340 (Supelco, Bellefonte, Pa.).

Polygalacturonase extraction and assay. Partially thawed vesicle tissue (15 g) was homogenized for 3 min in 15 ml Na-phosphate buffer (200 mm, pH 6.0) containing 2.4 m NaCl. After centrifugation (20 min at 32,000× g), proteins were precipitated from the supernatant by adding solid ammonium sulfate to 75% saturation. The precipitated protein was centrifuged for 20 min at 32,000× g. The pellet was dissolved in 2.5 ml Na-acetate buffer (50 mM, pH 5.0) and desalted by passing through a PD-10 column packed with Sephadex G-25M (Sigma). Protein content was determined calorimetrically as described by Bradford (1976) using bovine serum albumin as the standard.

Scanning electron microscopy. Some of the disordered and N juice vesicles isolated from fruit stored at 21°C for 40 days were prefixed in 2% glutaraldehyde in Na-phosphate (100 mM, pH 7.2) for 2 hr and postfixed in 2% osmium tetroxide in the same buffer. Specimens were washed with DI water for 2 hr, dehydrated using an ethanol series, and CO2 in a critical-point dryer (Ladd Industry, Burlington, Vt.). Specimens were razor blade-fractured, mounted, sputter-coated with Au, and examined with a Hitachi scanning electron microscope (S-530) at 20 kV of accelerating voltage.

Results and Discussion

Fresh weight, dry weight, and ethanol-soluble solids of grapefruit juice vesicles. The fresh weight (FW) of G vesicles was 60% greater than that of N vesicles (Table 1), which is in agreement with previous reports (Bartholomew et al., 1941; Sinclair and Jolliffe, 1961). Bartholomew et al. (1941) proposed that the size increase in G resulted from new growth of the affected vesicles. Whether the size of G vesicles is due to inherently large vesicles developing G or new growth during G development is not clear. G vesicles contained more dry matter (dry weight, DW) per vesicle than the N but DW on a per-gram FW basis was less. This observation indicates a reduced tissue density of G vesicles compared to N vesicles, as was also observed in G vesicles from oranges (Chakrawar and Singh, 1977a). The FW and DW, on a per-vesicle basis, were lower in VC than in N vesicles (Table 1). VC vesicles were severely shriveled at later stages. These symptoms are similar to those observed in vesicles affected by dry juice sac or core dryness (Bartholomew et al., 1941; Matsumoto, 1964; Noort, 1969).

Ethanol-soluble solids were higher in G and VC than in N vesicles on a per-gram FW basis. However, on a per-vesicle basis, a higher EIS content was observed for G vesicles (163%), whereas VC vesicles had only 57% more EIS than N vesicles, which was less than the standard deviations (Table 1). Sinclair and Jolliffe (1961) suggested that vesicles might produce more cell-wall material during the development of G. That EIS content did not change significantly in VC per vesicle indicates that the physiological events occurring in G vesicles did not occur in VC vesicles.

Polysaccharide composition of EIS from grapefruit tissue. Analysis of structural polysaccharides recovered from EIS revealed marked differences between the disorder types (Table 2). On a per-milligram EIS basis, there were slight or no differences in total pectin content between N and the disordered vesicles; however, on a per-vesicle basis, G vesicles had 2.5 times more total pectin content than N vesicles. For pectins, only WSP differed in VC vesicles compared to N. On either an EIS or vesicle basis, WSP decreased 45%.

In G compared to N vesicles, both WSP and CSP were higher on a per-vesicle basis, but lower on an EIS basis. Note that considerable difference in WSP and some difference in CSP content occurred between visually N vesicles isolated from fruit containing the respective disordered vesicles. Visually normal vesicles from G and VC fruit contained 27.8 µg and 17.6 µg WSP per vesicle, respectively. This observation and the slight difference in CSP indicate that the visually N vesicles were probably beginning to develop symptoms of the vesicle disorders. More soluble pectins may be the first wall components to change in disordered vesicles. Even so, WSP constituted a major component (1% to 3%) of the EIS and the contents of other polysaccharides examined in the visually N vesicles from VC and G fruit were similar, providing evidence that the normally appearing vesicles were still more like completely healthy vesicles than disordered vesicles, and, therefore, served as reasonable comparisons. Our recent work (unpublished data) indicates that the wall component changes occur very rapidly when they start.

Hemicellulose content on an EIS basis was lower in G than

| Table 1. Juice-vesicle fresh weight (FW), dry weight (DW), and ethanol-insoluble solids content (EIS) of normal and disordered vesicles of 'Marsh Seedless' grapefruit. |
|---------------------------------------------------------------|
| **Component** | **FW** | **DW** | **EIS** | **FW** | **DW** | **EIS** | **FW** | **DW** | **EIS** | **% change** | **FW** | **DW** | **EIS** | **% change** |
|----------------|-------|-------|--------|-------|-------|--------|-------|-------|--------|----------------|-------|-------|--------|----------------|
| Normal | 107 ± 8.9 | 11.9 | 11.9 | 83.6 ± 3.0 | 14.5 | 21.8 | 119 ± 3.1 | 12.4 | 113 ± 5.6 | 5.2 | -58 |
| Granulation | 174 ± 13.5 | 14.5 | 13.5 ± 0.8 | 2.34 | 163 | 8.8 ± 0.3 | 0.92 | 20.9 ± 0.6 | 0.97 | 5 |
| Granulated | 104 ± 7.2 | 21.8 | 13.5 ± 0.8 | 2.34 | 163 | 8.8 ± 0.3 | 0.92 | 20.9 ± 0.6 | 0.97 | 5 |
| Vesicle collapse | -55.6 | -58 | -58 | -58 | -58 | -58 | -58 | -58 | -58 |

*Values are the average of three replications ± SD (50 vesicles per replication for DW and FW). Values per vesicle are calculated from fresh-weight content and fresh weight per vesicle.

**Percentage change from apparently normal vesicle value.**
in N vesicles; however, on a per-vesicle basis, hemicellulose content increased 2-fold (Table 2). The hemicellulose contents for N and VC vesicles were similar.

Cellulose content did not change in either disorder type on an EIS basis, but was nearly three times higher in G than in N vesicles (Table 2). On a per-vesicle basis, G vesicles contained 1.8 to 2.5 times more structural polysaccharides (pectins, hemicellulose, and cellulose) than did N vesicles (Table 2). Although cell wall components were not always greater in G vesicles on an EIS basis, they were significantly greater on a FW basis than in N vesicles. These results suggest that G is associated with an anomalous increase in cell-wall synthesis and/or changes

Table 2. Comparisons of juice-vesicle cell wall components between normal and disordered tissue of ‘Marsh Seedless’ grapefruit.

| Disorder       | Normal | Granulated | % change | Normal | Vesicle collapse |
|----------------|--------|------------|----------|--------|-----------------|
|                | µg sugars per mg EIS | µg sugars per vesicle | % | µg sugars per mg EIS | µg sugars per vesicle | % |
| Pectins<sup>2</sup> |        |            |         |        |                 |       |
| TP             | 375 ± 12.1 | 334        | 54       | 349 ± 3.9 | 321        | 6      |
| WSP            | 31.2 ± 1.7 | 27.8       | 14.0     | 19.1 ± 0.8 | 17.6       | -45    |
| CSP            | 116 ± 8.7  | 103        | 13.7     | 130 ± 1.4 | 120        | 9      |
| Hemicellulose  | 111 ± 3.9  | 98.7       | 106      | 103 ± 3.2 | 95.0       | -1     |
| Cellulose      | 134 ± 11.6 | 119        | 179      | 121 ± 11.2 | 111        | -3     |

<sup>2</sup>Percentage change from normal vesicle value.

<sup>2</sup>TP, total pectins; WSP, water-soluble pectins; CSP, chelator-soluble pectins. Numbers are the average of three replications ± sd (50 vesicles per replication for DW and FW). Values per vesicle are calculated from content per EIS and EIS per vesicle.

Fig. 1. Gel-filtration profiles of water-soluble pectins from normal-appearing and disordered vesicles of ‘Marsh Seedless’ grapefruit. Decreasing molecular weight occurs with increasing fraction number. (A) Pectins from normal (N) vesicles from granulated (G) fruit; (B) pectins from G vesicles; (C) pectins from N vesicles from fruit with collapsed vesicle (VC); (D) pectins from VC.
in cell-wall turnover rate at the latter stages of fruit development. Differences in structural polysaccharide content were not observed in VC, although some qualitative changes were observed, particularly the decrease in WSP.

**Chromatography of pectins and hemicelluloses.** The gel-filtration elution patterns of WSP from N and G vesicles were similar (Fig. 1A and B); whereas the profile of the VC WSP exhibited a markedly lower void component and a relatively greater quantity of smaller Mr pectins fractionating on the gel (Fig. 1D) when compared to the N vesicle profile (Fig. 1C). The N vesicle isolates were nearly identical (Fig. 1A and C). These results are further evidence for WSP modification in VC.

The elution behavior of CSP was essentially similar to that of WSP, with the exception that the void peaks were more pronounced (Fig. 2). As with the WSP, disorder-associated changes were most pronounced in VC vesicles that yielded a more heterogeneous (in terms of Mr) population of CSP (Fig. 2D). Since the combined pectins in WSP and CSP comprised 40% of the total pectins, the data obtained from gel chromatography indicate that 1) G vesicles contained pectins unaltered in terms of Mr, and 2) pectin degradation might be a potential factor involved in the collapse of vesicles in VC or a secondary result from cellular breakdown.

Polygalacturonase (endo-PG) is temporally associated with softening and pectin changes in many fruit types (Brady, 1987; Huber, 1983a). In citrus, only very low levels of exo-PG have been reported in the fruit (Riov, 1974). In the present study, endo-PG activity was not detected in either N or disordered vesicles (data not shown). Although exo-PG may be present, exo-PG alone is unable to degrade polymeric pectins (Reese, 1977). The alternative explanation for these pectin changes in VC vesicles is non-PG-mediated modification. Low-molecular weight pectins appear during fruit ripening, at least in muskmelons, which also contain no endo-PG (McCullum, 1987). We suggest the possibility that there is release of pectins during turnover of hemicelluloses, but the factors causing this change are unknown.

Hemicelluloses extracted from G and VC vesicles and their respective, adjacent N control vesicles exhibited similar elution patterns on Ultrogel 34 (data not shown). The proportion of large-Mr, hemicelluloses was slightly higher in G than in N vesicles. The hemicellulose profiles indicate that slight modification of Mr may occur; however, if cell-wall degradation is involved in VC, pectin degradation is probably of more importance. Although the hemicelluloses may be of little importance in these disorders, the Mr distributions of all samples examined were remarkably similar to those observed for ripe tomato (Huber, 1983b), strawberry (Huber, 1984), and hot pepper (Grosset al., 1986) fruit. Ion exchange fractionation against a NaCl gradient did not indicate any differences in WSP, CSP, or hemicelluloses between VC, G, or N vesicles (data not shown).

**Neutral sugar composition of pectins and hemicelluloses.** Total neutral-sugar content of WSP was 87% higher in G and 16% higher in VC than in N vesicles (Table 3). There were higher levels of all neutral sugars associated with WSP from G, particularly arabinose and galactose, the two major neutral sugars

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Fig. 2. Gel-filtration profiles of chelator-soluble pectins from normal-appearing and disordered vesicles of ‘Marsh Seedless’ grapefruit. Decreasing molecular weight occurs with increasing fraction number. (A) Pectins from normal (N) vesicles from granulated (G) fruit; (B) pectins from G vesicles; (C) pectins from N vesicles from fruit with collapsed vesicles (VC); (D) pectins from VC.
in the WSP. Collapsed vesicles showed similar trends, but arabinose and galactose were present at still higher levels, even in apparently N vesicles, than in corresponding N and disordered vesicles from G fruit. This result again suggests that the normal-appearing vesicles in VC fruit were beginning to change and that pectin modification may be the first wall component affected. Pectins derived from vesicles in VC fruit contained about two times more neutral sugar than vesicles from G fruit. Chelator-soluble pectins from both N and disordered vesicles were extremely low in neutral sugars, 0.4 to 1.3 mol percent total neutral sugars (data not shown). By what mechanism neutral sugars in WSP increased in both disordered vesicles (VC and G) is not known. Gross (1986) reported that some water-soluble, ethanol-insoluble neutral sugars increased along with galacturonic acid during ripening of tomatoes. Thus, the increase in neutral sugars in disordered tissue may be an expression of tissue senescence. In N vesicles, the major sugars in hemicelluloses were xylose, glucose, and arabinose, in decreasing order (Table 4). Mannose was the third most prevalent sugar in disordered “vesicles. Major compositional differences in hemicellulose sugars were not detected between N and disordered vesicles, although arabinose levels appeared to be significantly lower in disordered vesicles compared to the N vesicles.

Scanning electron microscopy (SEM) disclosed significant differences between N, G, and VC vesicles. The micrographs were taken of the inner parenchymatous juice cells (Fig. 3). Healthy vesicles (Fig. 3A) had noticeably smaller juice cells than in G vesicles (Fig. 3B). In VC, most of the cells were partially or completely collapsed, resulting in stacked layers of cell walls (Fig. 3C). The lack of major changes in cell wall components of VC vs. N vesicles, along with the cell collapse, suggests that VC is a relatively rapid cell death phenomenon.

The greater size of cells in G has been proposed to result from new growth of the affected vesicles (Bartholomew et al., 1941; Matsumoto, 1964). Whether the size increase in G juice cells is due to originally large vesicles developing this disorder or new growth of cells during granulation is unknown. Juice vesicles at the stem end and stylar end are generally larger than

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### Table 3. Neutral sugars in water-soluble pectins from normal and disordered juice vesicles of ‘Marsh Seedless’ grapefruit.

| Neutral sugars | Granulation | Vesicle collapse | Normal | Granulated | Normal | Collapsed |
|----------------|-------------|------------------|--------|------------|--------|-----------|
| Rhamnose       | 0.3         | 1.5              | 2.0    | 1.0        |
| Fucose         | 0.4         | 0.7              | 1.6    | tr         |
| Arabinose      | 10.3 ± 0.2  | 13.9 ± 0.2       | 16.9 ± 0.4 | 23.6 ± 1.2 |
| Xylose         | tr          | 1.4              | 4.0    | 2.1        |
| Mannose        | tr          | 0.7              | 1.2    | 0.4        |
| Galactose      | 4.2 ± 0.1   | 10.0 ± 0.3       | 9.0 ± 1.0 | 13.3 ± 0.8 |
| Glucose        | nd          | 0.3              | tr     | tr         |
| Total          | 15.2        | 28.5             | 34.7   | 40.4       |

*Values are the average of two replications with two chromatographic injections per replication ± sd.

*nd and tr indicate nondetectable and trace, respectively.

### Table 4. Sugars composition of hemicelluloses from normal and disordered juice vesicles of ‘Marsh Seedless’ grapefruit.

| Neutral sugars | Granulation | Vesicle collapse | Normal | Granulated | Normal | Collapsed |
|----------------|-------------|------------------|--------|------------|--------|-----------|
| Rhamnose       | tr          | tr               | 2.4    | 0.9        | 1.6    | 0.8       |
| Fucose         | 1.7 ± 0.3   | 2.9 ± 0.6        | 11.5 ± 0.4 | 7.9 ± 0.2 |
| Arabinose      | 10.2 ± 1.4  | 7.3 ± 0.7        | 56.5 ± 4.2 | 60.4 ± 1.2 |
| Xylose         | 59.9 ± 0.4  | 55.7 ± 2.9       | 9.0 ± 1.3 | 11.1 ± 0.8 |
| Mannose        | 8.0 ± 0.1   | 9.9 ± 1.3        | 9.0 ± 1.2 | 6.8 ± 1.1 |
| Galactose      | 7.5 ± 1.0   | 8.3 ± 0.9        | 11.5 ± 0.6 | 12.3 ± 0.5 |
| Glucose        | 12.5 ± 0.6  | 15.9 ± 1.4       | 11.5 ± 0.6 | 12.3 ± 0.5 |

*Values are the average of two replications with two injections per replication ± sd.

*tr and nd indicate trace and nondetectable, respectively.
those in the central portion (Matsumoto, 1964). In the present study, fresh and dry weights increased only 60% and 22%, respectively, in G vesicles, whereas EIS increased > 150% on a per-G vesicle basis. This result can be interpreted as evidence of new synthesis in G vesicles, especially pectins, cellulose, and possibly lignin (Shomer et al., 1989), all of which may contribute to the increase of juice-cell size, as new wall synthesis, if wall structural component bonding release mechanisms also operate (Bolwell, 1988). Lignin was readily detected histochemically by phloroglucinol and toluidine blue in G vesicles, whereas N vesicles seldom showed positive staining for lignins (Burns and Achor, 1989). We also treated EIS with phloroglucinol, and positive lignin staining occurred only in G vesicles. Uneven cell-wall thickening, especially in hyperdermal cells, in G vesicles has been found in oranges (Bartholomew et al., 1941; Matsumoto, 1964), in partially collapsed grapefruit vesicles (Burns and Achor, 1989) and in pummelo (Shomer et al., 1989), but the marked changes in cell-wall thickness were not observed in epidermal or hypodermal cells in our study using SEM. About 40% of the EIS is unaccounted for by the structural polysaccharides measured. Proteins and pectic neutral sugars may account for most of this additional EIS in N and VC, since starch is not present in citrus vesicles (Gebhardt et al., 1982). Lignin is implicated in the large increase (> 150%) of EIS per-G vesicle (Shomer et al., 1989). Protein was not elevated in pummelo (Shomer et al., 1989). In future work, lignin and protein contents of the EIS should be quantified.

The differences in wall components measured in this study indicate that, in grapefruit, G and VC of juice vesicles are different phenomena and not stages of the same disorder or senescence process, even though G vesicles partially collapse in the final stages of this disorder (Bartholomew et al., 1941).

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