Mobility Limitations of Bound Polygalacturonase in Isolated Cell Wall from Tomato Pericarp Tissue

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Abstract. Enzymically active cell wall isolated from mature-green and ripening tomato (Lycopersicon esculentum Mill cv. ‘Rutgers’) fruit was employed to investigate the mobility of the enzyme polygalacturonase (PG, EC 3.2.1.15). Cell walls from mature-green ‘Rutgers’ fruit or from the ripening mutant rin, which alone exhibits little or no release of pectin, were unaffected by the addition of enzymically active cell wall from ripening ‘Rutgers’ fruit, indicating that PG is either not transferred at all or is not transferred to sites of pectin hydrolysis. The quantity of pectin released by the addition of soluble PG to enzymically active wall depended on the quantity of enzyme added. Similar data were obtained using purified PG2. Pectin solubilization from all walls exhibiting enzymically mediated pectin release diminished with time; however, transfer to fresh buffer initiated a resumption of autolytic activity, indicating that an inhibitor is released during the course of pectin hydrolysis.

Depolymerization of cell wall polyuronides by the endopolygalacturonases (EC 3.2.1.15) is believed to be the dominant feature of softening metabolism in tomato and several other fruit types (Hobson, 1964; Huber, 1983). Purified polygalacturonases (PGs) and crude PG extracts readily hydrolyze pectic polymers in solution and also cause solubilization of uronic acids from isolated cell walls of normal tomato and from the mutant rin (Ali and Brady, 1982; Themmen et al., 1982). Little is known about the action of these enzymes in the wall environment.

In previous work (Rushing and Huber, 1984), we reported on the use of isolated cell wall to investigate details of PG-mediated wall hydrolysis in ripening tomato fruit. Evidence of the enzymic nature of cell wall autolysis included the dependency of activity on ripening and the inhibitory and stimulator effects of Ca++ and NaCl, respectively, which are known to influence the activity of purified PG (Ali and Brady, 1982; Pressey and Avants, 1982). Additional studies revealed the tenacity of PG binding to isolated cell walls. Treatments with high ionic strength, which have been used to facilitate extraction of PG (Hobson, 1964; Ali and Brady, 1982; Pressey, 1986), did not reduce autolytic activity in wall isolated from ripening tomato fruit (Rushing and Huber, 1987). Other results suggested that PG maybe regulated through the interaction of pH, soluble Ca++ and ionic strength of the apoplast (Rushing and Huber, 1987). These factors may regulate PG activity by affecting the mobility of the enzyme within the cell wall (Rushing and Huber, 1985). In this paper, we report on the use of autolytically active cell wall to investigate the mobility characteristics of PG bound to cell wall.

Materials and Methods

Plant material. Normal ‘Rutgers’ tomato and the ripening mutant rin plants were field-grown on stakes and black plastic mulch at the Univ. of Florida Horticultural Unit near Gainesville. Fruits were harvested at the mature-green, pink, red (ripe), and overripe developmental stages based on a USDA visual aid for color classification of tomatoes (The John Henry Co., 1975). Fruit were surface-sterilized with 100 ppm NaOCl, rinsed, dried, and sectioned; seed and placental tissue were removed, and the pericarp was stored at −20°C in sealed polyethylene bags.

Cell wall preparation. Frozen pericarp (∼100 g) was peeled and homogenized in 400 mL of distilled H2O (4°C) for 1 min in a blender set at maximum speed. The homogenate was transferred onto Miracloth and washed with 3 liters of distilled H2O (4°C) and 1 liter of acetone (1°C) with continuous stirring. Cell wall was recovered from the Miracloth and suspended in 100 mL of cold acetone for 1 hr, then filtered through Whatman GF/C filters and washed with cold acetone (∼500 mL) until pigments were no longer visible. Wall was transferred to a beaker and dried for 24 hr at 30°C, with occasional stirring to reduce formation of aggregates. Dry cell wall was stored in sealed containers at 22°C and was used for autolysis experiments within 2 weeks after preparation. For some autolysis experiments, freshly prepared wall was used immediately after removal from the Miracloth without exposure to acetone. All wall preparations were analyzed for total uronic acid content (Ahmed and Labavitch, 1977).

Preparation of PG extract and of purified PG2. About 100 g of frozen pericarp from peeled, ripe ‘Rutgers’ tomatoes were added to 100 mL (IC) buffer (50 mM Na-acetate, pH 4.5) containing 0.02% (w/v) Thimerosal (Sigma, St. Louis). After the addition of NaCl to a concentration of 1 M, the mixture was homogenized for 1 min in a blender set at maximum speed. The homogenate was held at IC for 2 hr, then stirred, centrifuged (10 min at 7000 × g), and the pellet discarded. The supernatant was adjusted to 80% saturation with (NH4)2SO4, held at IC for 24 hr, and then centrifuged (20 min at 7000 × g). The supernatant was discarded and the pellet was resuspended in 10 mL of buffer (100 mM Na-acetate, pH 4.5) containing 100 mM NaCl and 0.02% Thimerosal. This solution was dialyzed against 4 liters of the same buffer with three changes during 48 hr at 4°C. The final change of bathing buffer was analyzed for the presence of carbohydrate by the phenol–sulfuric assay for total sugar (Dubois et al., 1956). Protein content of the dialyzed extract was defrayed in part by the payment of page charges. Under postal regulations, this paper therefore must be hereby marked advertisement solely to indicate this fact.

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was determined by the method of Bradford (1976). PG2 was isolated and purified using the procedure of Ali and Brady (1982), as modified by DellaPenna et al. (1986).

**Autolysis experiments.** Cell wall (100 mg dry weight) from pericarp of either mature *rin* or ‘Rutgers’ tomatoes at mature-green, pink, red, or overripe stage of development was placed in a 50-ml centrifuge tube in an ice bath. Wall was dehydrated by dropwise addition of 15 ml of cold (1°C) reaction buffer (50 mM Na-acetate, pH 4.5) containing 150 mM NaCl and 0.02% Thimerosal. In other experiments, samples were composed of 50 mg of wall from either mature-green ‘Rutgers’ or *rin* tissue, along with 50 mg of wall from either pink, red, or overripe ‘Rutgers’ tissue. After dehydration with buffer, samples were held at IC for 30 rein, with occasional stirring, then centrifuged (10 min at 7000 × g) using a Beckman Model J2-21 centrifuge equipped with a JA-20 rotor. The supernatant was filtered through Whatman GF/C filters to obtain a time-zero sample and the wall was resuspended in 15 ml of fresh reaction buffer. Reaction mixtures were incubated in a shaking bath at 34°C. Centrifugation, supernatant removal, and pellet resuspension steps were repeated at selected intervals during incubation. All supernatants were analyzed for total uronic acids (Blumenkrantz and Asboe-Hansen, 1973) and uronic acid-reducing sugars (Milner and Avigad, 1967). After the 7 hr of incubation, filtered wall was suspended in 30 ml of acetone, filtered through tared Whatman GF/C filter paper, and dry weights obtained after storage for 48 hr in a desiccator *in vacuo*.

Other autolysis studies used freshly extracted, hydrated wall from tissues of the same stages of development as described above. Autolysis experiments employing various mixtures of cell wall preparations were also performed. Since preautolysis dry weights were not obtainable, walls were dispersed by volume using a spatula. After transferring walls to centrifuge tubes, the total volume was adjusted to 15 ml with reaction buffer. All subsequent steps were as described for experiments with dry cell wall preparations. Postautolysis dry weights were added to the weight (µg uronic acid equivalents) of solubilized uronic acids to obtain an estimate of the preautolysis wall dry weight.

In an additional autolysis study using dried cell wall, two methods for sampling soluble products were compared for their effect on release of polyuronides from walls of either mature-green or red ‘Rutgers’ pericarp. The first sampling method was as described earlier (i.e., centrifugation and supernatant removal), after which walls were provided with fresh reaction buffer. This method was compared to that of removing an aliquot (0.5 ml of buffer) from the reaction mixture at intervals throughout the experiment. Aliquots were filtered through Whatman GF/C filter papers and analyzed for total uronic acids and uronic acid-reducing groups. Postautolysis recovery of cell wall was performed as described previously.

**PG activity in supernatants from autolytic reactions.** Supernatants were recovered by filtration from autolysis reaction mixtures in which cell wall from either pink or red pericarp had been incubated. To 6 ml of the supernatant were added either 50 mg of wall isolated from mature-green pericarp or 12 mg polygalacturonic acid (Sigma). Boiled supernatant controls were included. An aliquot (0.5 ml) was taken immediately for assay of acid-reducing sugars (Milner and Avigad, 1967), and additional aliquots were removed after 4 and 12 hr incubation at 34°C in a shaking water bath.

**Cell wall treated with multiple additions of PG.** Autolytically inactive cell walls (100 mg dry weight) from ‘Rutgers’ mature-green tomatoes were suspended in 15 ml of cold buffer (50 mM Na-acetate, pH 4.5) containing 150 mM NaCl and 0.02% (w/v) Thimerosal. To the suspension was added 200 µl of either active or boiled enzyme (≈75 µg protein), or buffer alone. Reaction mixtures were incubated at 34°C in a shaking water bath. Initially, and at selected intervals, aliquots (0.5 ml) were removed, filtered through Whatman GF/C filters, and analyzed for uronic acid content. When the rate of polyuronide solubilization had declined to near zero, reaction mixture volumes were readjusted to 15 ml with fresh buffer and the enzyme addition repeated. A total of four aliquots of enzyme were added during the 32 hr of incubation. Cell wall recovery for dry weight determination was performed as described for autolysis experiments.

Similar experiments were performed using purified PG2. Cell walls (25 mg) from mature-green tomato fruit were dehydrated by addition of 7.5 ml cold buffer (50 mM Na-acetate) containing 150 mM NaCl and 0.02% (w/v) Thimerosal. To the suspension was added 200 µl of the PG2 preparation (≈18 µg protein). Other details were as described above.

In other experiments, cell wall was exposed to high pH to demethylate pectins before addition of PG (Babbit et al., 1973). Wall was dehydrated in 15 ml of buffer (50 mM NaHCO3-NaOH, pH 11.0) and allowed to stand for 30 min at 20°C. After this exposure to high pH, samples were adjusted to 80% acetone (v/v) to precipitate the cell wall, then centrifuged (10 min at 7000 × g). The supernatant was removed, and the cell wall was washed once with 20 ml of reaction buffer, and then resuspended in 15 ml of reaction buffer. An aliquot of enzyme was added and the procedure was performed as described above, except that the incubation time was reduced to 10 hr.

In another multiple addition experiment, either active or boiled PG was added to autolytically inactive cell was as described. When the rate of polyuronide solubilization declined to near zero, the reaction mixture was centrifuged (10 min at 7000 × g), the supernatant removed, and the wall provided with 15 ml of fresh buffer. No additional PG was added to the mixture, allowing us to determine whether further polyuronide solubilization would occur in the absence of additional enzyme.

**Results**

**Autolysis experiments.** Polyuronides were released from enzymically active cell wall (Fig. 1) at rates comparable to those observed in earlier studies, in which autolytic activity was first noted in wall from fruit at the onset of ripening and increased as ripening proceeded, then decreased slightly during senescence (Rushing and Huber, 1984). The only exception here is that uronic acid solubilization from wall of pink fruit was considerably lower here than in previous work. Reaction mixtures composed of cell wall from *rin* tissue along with enzymically active walls from ‘Rutgers’ pink, ripe, or over-ripe pericarp released quantities of uronic acids equal to the yield expected from the enzymically active wall (Fig. 1). Similar results were obtained when *rin* cell wall was used in place of wall from ‘Rutgers’ mature-green fruit (data not shown). In experiments using freshly prepared, hydrated cell wall, results were identical to those obtained with cell wall that had been dried following isolation (data not shown). In all experiments reported here, exposure of cell wall to boiling 80% EtOH before incubation at 34°C reduced autolytic activity to near zero, regardless of developmental stage of the tissue from which walls were isolated (data not shown).

Periodically transferring autolytically active wall to fresh buffer was found to have a dramatic effect on the quantity of polyuronides released. When cell wall was maintained in the initial
amounting to 6% of the initial wall dry weight was released during incubation, at which time 80 µg uronic acid/mg of cell wall had been solubilized (Fig. 2). In contrast, reaction mixtures that were resuspended in fresh buffer at each measurement released >100 µg polyuronide/mg of wall dry weight during the first 3 hr, and continued to release uronic acids throughout incubation (Fig. 2). Wall periodically provided with fresh buffer during 7 hr release nearly 150 µg uronic acid/mg of wall dry weight, almost a 95% increase over the amount released when fresh buffer was not provided. Cell wall from mature-green tissue did not release acid sugars during incubation, regardless of buffer conditions (data not shown).

PG activity in supernatants from autolytic reactions. Protein amounting to ≈16% of the initial wall dry weight was released from wall of both pink and red pericarp during autolysis at pH 4.5. Boiled controls released <1% of the initial weight (data not shown). Pressey (1986) reported that PG is not effectively extracted from tomato pericarp at this pH; however, we observed the release of protein from our enzymically active wall. The protein apparently did not contain more than a trace of PG activity; i.e., it did not cause the solubilization of uronic acid sugar-reducing groups when autolysis supernatants were mixed with polygalacturonic acid or with wall isolated from mature-green pericarp (data not shown).

Cell wall treated with multiple additions of PG. Each addition of the unpurified PG preparation (≈75 µg) to the reaction mixture resulted in the solubilization of 10 µg uronic acid/mg of wall dry wt (Fig. 3). The rate of release declined to near zero within 2 hr after each addition of active enzyme (Fig. 3). Boiled enzyme did not cause release of polyuronides (Fig. 3), nor did buffer alone (data not shown). Pretreating wall in pH 11.0 buffer before addition of enzyme did not influence the rate or extent of polyuronide solubilization compared to wall with no such exposure (Fig. 4).

As in the autolytic reactions, replacing the buffer caused further release of polyuronide in the absence of additional enzyme in quantities that amounted to ≈60% of those released by the initial aliquot of enzyme (Fig. 5). In contrast, walls that were centrifuged but resuspended in the same buffer did not release additional uronic acids (Fig. 5).

Figure 6 illustrates that the kinetics of pectin release exhibited by the crude PG preparation were also observed with purified buffer, the rate of release declined to near zero after 3 hr incubation, at which time ≈80 µg uronic acid/mg of cell wall had been solubilized (Fig. 2). In contrast, reaction mixtures that were resuspended in fresh buffer at each measurement released >100 µg polyuronide/mg of wall dry weight during the first 3 hr, and continued to release uronic acids throughout incubation (Fig. 2). Wall periodically provided with fresh buffer during 7 hr release nearly 150 µg uronic acid/mg of wall dry weight, almost a 95% increase over the amount released when fresh buffer was not provided. Cell wall from mature-green tissue did not release acid sugars during incubation, regardless of buffer conditions (data not shown).

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polygalacturonase (PG2). Since cell wall was not treated to inactivate-bound enzymes, the possibility remains that even the purified enzyme was acting in concert with other proteins.

Discussion

The tenacity of binding of PG to cell wall is exemplified by the fact that high concentrations of ionic strength are required to effect removal of the enzyme (Pressey, 1986). Heterogeneity of binding is also indicated by the observation that wall subjected to high concentrations of salt shows little alteration in the quantitative features of cell wall autolysis (Rushing and Huber, 1987).

On the basis of the inability of high salt and other agents to achieve complete removal of wall-associated enzymes, some workers have suggested that, for bacterial autolytic enzymes (Brown et al., 1970; Shockman et al., 1967) and for plant invertases (Parr and Edelman, 1975) and peroxidases (Dashek et al., 1979), a proportion of the enzyme population is covalently bound. Although there is no direct evidence for covalent association, it is quite clear that the binding between the proteins of both endogenous and foreign origin and cell wall is quite strong (Jansen et al., 1960; Nagahashi, 1987). Strong ionic interactions are often implicated, but steric occlusion, lectin interactions, or binding to substrates may also be involved. The tenacity of binding of PG to cell wall raises interesting questions regarding the manner by which PG functions in autolytic reactions. Within the duration of the autolytic experiments reported here, only trace quantities of PG activity were detected in the bathing solution, although considerable protein was present. We did not investigate the possibility that this protein might represent, in part, inactivated PG. Additional evidence for the failure of cell wall to yield significant quantities of active PG during autolytic reactions was the observation that PG was not transferred in reactions employing mixtures of active and inactive wall. This fact, in addition to the dependency of autolytic yields on the quantity of bound PG (Rushing and Huber, 1984), indicated that individual PG molecules function within a limited microenvironment. The failure of cell wall to release PG during autolytic reactions may seem expected, in view of reports that PG is not released from cell wall at pH 4.5 (Pressey, 1986). It must be borne in mind that in the latter study, extractions were performed under low temperature. Thus, these data do not address the question of whether PG is released under conditions of autolysis.

Inhibitors accumulating in the incubation buffer may be an important factor in this phenomenon, since providing fresh buffer significantly increased the quantity of uronic acids released during autolysis (Fig. 2). Evidence that the inhibitor may be solubilized by PG action is found in the results of the multiple-
addition experiments. The capacity of added PG to solubilize cell wall uronic acids was, in time, diminished; however, transferring the cell wall to fresh buffer caused a resumption of polyuronide solubilization without the addition of more enzyme (Fig. 5). This result indicates that, in a closed static system, individual PG molecules have a limited hydrolytic capacity. If one assumes that the inhibitor is released from the wall by PG action, there are numerous possibilities. Products released as a consequence of pectin hydrolysis in cell wall include uronic acid (Hobson, 1964), calcium (Rigney and Wills, 1981), neutral sugars (Gross and Wanner, 1979; Huber, 1983; Gross and Sams, 1984), protein (Hobson et al., 1983), and possibly other unidentified components of the cell wall. Calcium exhibits the characteristics capable of causing the observed suppression of autolysis in a static system. Calcium could explain the decrease in hydrolytic capacity via its ability to rigidify polymeric pectin and consequently restrict the capacity of PG to migrate within the wall. The addition of exogenous calcium has been shown to greatly diminish autolytic activity (Rushing and Huber, 1987). The reversibility of the inhibition is also consistent with a role for calcium.

Alkaline desterification of wall pectin did not appear to influence the extent of pectin hydrolysis in isolated cell wall. It is possible, however, that the cell wall preparations contained bound pectin methylesterase (PME, EC 3.1.1.11), which, if functioning under the reaction conditions, would serve to enzymically demethylate pectin. Seymour et al. (1987) demonstrated a promotive effect of PME on PG-mediated wall hydrolysis.

A final point worth noting is the variability observed in the amount of uronic acid released from cell wall from pink pericarp compared to the yields from walls of tissues at other stages of development (Rushing and Huber, 1984; Fig. 1). Pink tomatoes are not uniformly pink, but are composed of zones of green and light-red tissues (The John Henry Co., 1975). When selecting samples of pericarp for cell wall extraction, it is difficult to ensure that each sample is composed of the same proportion of the chronologically distinct tissue types. Since PG apparently does not move freely through mixtures of cell wall from different developmental stages in vitro (Fig. 1), considerable variability could result from using pink tissue. The consistency observed in autolytic yields from red fruit is probably due to a more-uniform distribution of PG within the pericarp tissue. We conclude from these data that the binding and mobility characteristics of PG are of considerable importance in regulating pectin hydrolysis in isolated cell wall and, possibly, in vivo. The inhibition observed in autolysis reactions performed under static conditions may in fact be more accentuated in vivo, particularly if the soluble inhibitor remains sequestered in the apoplast. This situation would explain why pectin hydrolysis in vitro is of a significantly greater magnitude than that occurring in vivo.

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