Changes in Cell Wall Properties Coincide with Overexpression of Extensin Fusion Proteins in Suspension Cultured Tobacco Cells

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Abstract

Extensins are one subfamily of the cell wall hydroxyproline-rich glycoproteins, containing characteristic SerHyp4 glycosylation motifs and intermolecular cross-linking motifs such as the TyrXaaTyr sequence. Extensins are believed to form a cross-linked network in the plant cell wall through the tyrosine-derivatives isodityrosine, pulcherosine, and di-isodityrosine. Overexpression of three synthetic genes encoding different elastin-arabinogalactan protein-extensin hybrids in tobacco suspension cultured cells yielded novel cross-linking glycoproteins that shared features of the extensins, arabinogalactan proteins and elastin. The cell wall properties of the three transgenic cell lines were all changed, but in different ways. One transgenic cell line showed decreased cellulose crystallinity and increased wall xyloglucan content; the second transgenic cell line contained dramatically increased hydration capacity and notably increased cell wall biomass, increased di-isodityrosine, and increased protein content; the third transgenic cell line displayed wall phenotypes similar to wild type cells, except changed xyloglucan epitope extractability. These data indicate that overexpression of modified extensins may be a route to engineer plants for bioenergy and biomaterial production.
Introduction

Plants are the major source of food and chemicals on earth. With our intensifying desire for sustainable energy, the engineering of plants, in particular their cell walls, to achieve greater biofuel production is a major focus of current cell wall research. Genetic manipulation of plants using gene transformation or selective breeding allows the production plants possessing higher biomass and digestibility [1–3].

One approach is to modify the enzymes involved in wall polymer biosynthesis. Examples include the modification of caffeic acid O-methyltransferase (COMT) and caffeoyl CoA 3-O-methyltransferase (CCoAOMT) [4,5], two enzymes within the lignin biosynthesis pathway, to reduce lignin content and/or modify its structure for higher saccharification of biomass. Down-regulation of the COMT gene in switchgrass made the wall more amenable to degradation [4]. The modification resulted in low lignin, a low syringyl:guaiacyl lignin ratio in the transgenic plants, and a 8% increase in ethanol production during traditional fermentation. Similarly, suppression of CCoAOMT in transgenic poplar tree reduced cell wall lignin content as much as 10% and significantly increased glucose yield from mature poplar wood when subjected to enzymatic digestion [5].

To reduce the costs associated with chemical pretreatment of biomass, another approach involves expression of wall glycan degrading enzymes such as cellulases and hemicellulases in planta. Examples of this approach include expression of Clostridium thermocellum xylanase (xynZ) in the apoplast of transgenic tobacco [6] and expression of Acidothermus cellulolyticus endoglucanase E1 in tobacco chloroplasts [7]. Interestingly, E1 protein was biologically active in both fresh and dried leaves. Expression of a poplar cellulase (PaPopCel1) in Arabidopsis resulted in cell elongation and subsequent increased cell size, most likely because the poplar cellulase removed disordered glucose from the cellulose microfibrils, which probably reduced their cross-links with xyloglucans [8]. Transgenic maize expressing xylanase XynA, or endoglucanase, or both enzymes, yielded up to 141% higher glucose and 172% higher xylose compared to control plants. This resulted in a total of 55% increase in ethanol production [9].

Here we report an approach to changing the cell wall properties by overexpressing cross-linkable chimeric P3 extensin analogs that also contain repeats of an AGP glycosylation-motif and repeats of the human elastin peptide VPGVG in tobacco (Nicotiana tabacum) BY-2 (Bright Yellow) suspension cultured cells. Extensins belong to a superfamily of plant cell wall glycoproteins, the hydroxyproline-rich glycoproteins (HRGPs), which also include proline-rich proteins (PRPs) and arabinogalactan-proteins (AGPs) [10]. Unlike the AGPs that contain clustered AlaHyp, SerHyp, and ThrHyp glycosylation-motifs and variations thereof, allowing for arabinogalactan polysaccharide addition [11,12], extensins share a diagnostic repetitive SerHyp4 glycosylation-motif that directs Hyp-O-glycosylation with short oligoorabinosides [13]. Extensins often include Tyr-containing cross-linking motifs, such as ValTyrLys in P1-type extensins and...
TyrXTyrLys (X represents any amino acid except Pro) in P3-type extensins [10]. These motifs lead to the formation of the cross-linked tyrosine derivatives, including isodityrosine (Idt), pulcherosine, and di-Idt. Pulcherosine and di-Idt covalently cross-link extensin monomers to form protein scaffolds in the cell wall [14–17]. The cross-linking property of extensins provides a potential route to modify the plant cell wall.

Elastin is a cross-linked protein distributed throughout the animal extracellular matrix, such as arteries, lungs, elastic ligaments, bladder, and skin [18, 19], imparting elasticity to those tissues. Due to the number of cross-links formed through Lys residues, elastin is highly insoluble. However, its precursor, tropoelastin, is soluble and composed of two major domains, a hydrophilic cross-linkable domain rich in Lys and Ala residues, and a hydrophobic elastic domain rich in Val, Pro, Gly, and Ala residues. The elastic domain contains repetitive amino acid motifs, including the peptides VPGG, VPGVG, APGVGV, and VPGFGVGAG [20]. Among these sequences, the pentapeptide VPGVG is the most common element with more than 50 such repeats occurring in one elastin molecule. We aimed to change cell wall properties by introducing the exogenous elastin motifs, as well as AGP motifs, into the wall through the synthetic extensin analogs [17].

The cell wall properties of three different transgenic cell lines were all changed, but in different ways. One transgenic cell line was observed with a decreased cellulose crystallinity and increased wall xylolucan content; the second transgenic cell line contained dramatically increased hydration capacity and notably increased cell wall biomass, increased di-isodityrosine, and increased protein content; the third transgenic cell line displayed wall phenotypes similar to wild type cells, except changed extractability of xylolucan epitopes. These results suggest that wall extensin modification has potential as an approach to manipulate wall structure, in some instances resulting in greater sugar release for bioenergy production.

Materials and Methods

1. Construction of binary vectors
Two oligonucleotides, 5′-CCATCAGGAGTAGGTGTTCCAGGAGTTGGCGTTCC-AGGAGTTGGCCTCCAGCACCTGCACCAGCCCGGTGTGGAGATCTGGTG- TTGTTGTACCTGTTGTTGTTGTGGT-3′ and 5′-TACCTGGTTGTGGGTTCACTGGTGTT-GTGTGG-TCCTATGAGTTGGTGCCACGGTGACCCGACCACGACCTCGCCGGTTGGAC-GGACCTCGCGTTGAGGACCTTGATGGAGGAC-3′, were designed encoding an AGP motif (Ala-Pro-Ala-Pro-Ala-Pro) flanked by two repeats of human elastin motif Val-Pro-Gly-Val-Gly at each end (Fig. 1A). The oligonucleotide set was abbreviated as $E_2AE_2$ (E: elastin motif; A: AGP motif).

Fifty nanograms of each above oligonucleotide were annealed in 1X ligase buffer and ligated to a $pUC18$ vector between the BbsI and BsmF1 restriction sites. The resulting plasmid was named as $pUC-E_2AE_2$, which was digested by two sets
Fig. 1. Designed nucleotides (A), deduced protein sequences (B), and purification of 442 glycomodule (C).

A. Designed nucleotides encode four repeats of human elastin motifs separated by an AGP motif Ala-Pro-Ala-Pro-Ala-Pro-Ala, with two sticky ends for cloning.

B. Predicted polypeptide sequence of each glycomodule based on previous work on synthetic gene products. However, based on amino acid composition analysis, Pro residues in the elastin motifs were also hydroxylated.

C. Size exclusion chromatograms of the three Wall Modification by Over-Deposition of Extensin Analogs
of restriction enzymes BbsI/ScaI and BsmFI/ScaI, respectively [17]. The 1.1 kb BsmFI-ScaI fragment and the 1.8 kb BbsI-ScaI fragments were ligated to form a vector containing two repeats of \(E_2AE_2\). The corresponding plasmid was named \(pUC-E_2AE_4AE_2\).

By using the same strategy shown above, the BsmFI-ScaI fragment of \(pUC-YK_2\) or \(pUC-YK_n\) [17] containing the \(YK_2\) or \(YK_4\) gene was ligated with the BbsI-ScaI fragment of \(pUC-E_2AE_2\) and \(pUC-E_2AE_4AE_2\), respectively, which lead to the formation of \(pUC-YK_2-E_2AE_2\) and \(pUC-YK_2-E_2AE_4AE_2\), and \(pUC-YK_4-E_2AE_4AE_2\). Here, \(YK\) stands for gene encoding peptide Ser-Pro-Ser-Pro-Ser-Pro4-Tyr-Tyr-Tyr-Lys [17].

Similarly, \(pUC-YK_2-E_2AE_2\), \(pUC-YK_2-E_2AE_4AE_2\), and \(pUC-YK_4-E_2AE_4AE_2\) were dimerized with themselves, respectively. The desired gene sizes of \((YK_2-E_2AE_4AE_2)_2\) and \((YK_4-E_2AE_4AE_2)_2\) were about 600 and 800 bps. The corresponding plasmids were named \(pUC-(YK_2-E_2AE_4AE_2)_2\) and \(pUC-(YK_4-E_2AE_4AE_2)_2\). The obtained \(pUC-(YK_2-E_2AE_2)_2\) was further dimerized, formed \(pUC-(YK_2-E_2AE_2)_4\) with a gene size about 800 bps. From now on, the \((YK_2-E_2AE_2)_2\), \((YK_4-E_2AE_2)_2\), and \((YK_2-E_2AE_2)_4\) genes were abbreviated as 242, 442, and 224, respectively.

The XmaI-NcoI fragments containing the synthetic genes in \(pUC-224\), \(pUC-242\), and \(pUC-442\) were ligated into \(pUC-SS^{lob}-224-EGFP\), respectively, to replace the \((AP)_{51}\) fragment (\((AP)_{51}\) stands for a synthetic gene encoding 51 repeats of Ala-Pro [12], and \(SS^{lob}\) stands for the signal sequence of a tobacco extensin [11]). The three new plasmids were named as \(pUC-SS^{lob}-224-EGFP\), \(pUC-SS^{lob}-242-EGFP\), and \(pUC-SS^{lob}-442-EGFP\). The gene cassettes were further cloned into a \(pBI121\) vector between the BamHI and SacI restriction sites, formed plant transformation vectors \(pBI121-SS^{lob}-224-EGFP\), \(pBI121-SS^{lob}-242-EGFP\), and \(pBI121-SS^{lob}-442-EGFP\). All gene sequences were confirmed by DNA sequencing.

2. Transfer of the genes into tobacco cells

One hundred nanograms of above each constructed \(pBI121\) plasmid were transformed into \(Agrobacterium\) strain LBA4404 by the freeze-thaw method [21]. Positive colonies of \(Agrobacterium\) were selected via Kanamycin/Streptomycin resistance. The transformed \(Agrobacteria\) were used to co-culture with 4-day-old tobacco BY2 cells at 28°C for 2 days [12]. The infected tobacco cells were washed 4 times with Schenk and Hildebrandt (SH) culture media and were spread on SH solid plates with Kanamycin (100 µg/ml) and Timentin (200 µg/ml). The Kanamycin-selected cells were subcultured in SH culture media as described earlier [12].
3. Purification of fusion proteins and glycomodules from suspension culture media

The culture media of 20-day-old transgenic tobacco BY2 cells were filtered from the cultures, concentrated through rotary evaporation, and dialyzed against distilled deionized (d.d.) H₂O. The EGFP fusion proteins in the dialyzed medium were separated via a combination of chromatography, including hydrophobic interaction (HIC) (Phenyl Sepharose 6 Fast Flow, 16 × 700 mm, Amersham-Pharmacia Biotech) and C-18 reverse-phase (10 μm, PRP-1, 7 × 305 mm, Hamilton) as described [12].

Eight to ten milligrams of 224-EGFP, 242-EGFP, and 442-EGFP were dissolved in 500 μl of d.d. H₂O and denatured at 100 °C for 5 min. Five hundred μl of 4% (w/v) ammonium bicarbonate and 10 μl of 10 μg/μl trypsin were added to the solution, which were incubated at room temperature for 1 day. The resulted 224, 242, and 442 glycomodules were purified from the digestion mixture, respectively, through a semipreparative Superose-12 gel filtration (16 × 500 mm, Amersham-Pharmacia Biotech) using 50 mM sodium phosphate buffer (pH 7.0) and C-18 reverse-phase (5 μm, PRP-1, 4.1 × 150 mm, Hamilton) chromatography as reported [12].

4. Cell wall preparations of suspension cultured cells

Forty grams of above filtered fresh cells from each suspension culture were resuspended in 80 ml of suspension buffer (Tris-HCl, 50 mM, pH 7.0, 0.1% BSA (w/v), 1% sucrose, 0.1% sodium metabisulfite) and cooled on ice, followed by sonication for 12 minutes on ice (30 second sonication and 1 minute interval cycles). The cell debris was spin at 3000 × g for 5 minutes. The supernatant was discarded. The cell walls were washed with 600 ml of 1 M NaCl and then with 2 L of d.d. H₂O until the conductivity of the supernatant equal to that of d.d. H₂O. The resulted cell walls were lyophilized.

Cell walls used for glycosyl composition, IDT and di-IDT measurement, glycome profiling, sugar release, and solid state NMR analyses were prepared as follows [22]: Forty grams of filtered cells were washed with 100 mM potassium phosphate (pH 7.0) and 500 mM potassium phosphate (pH 7.0), broken in 500 mM potassium phosphate (pH 7.0). The cell walls were washed in order with 500 mM potassium phosphate (pH 7.0), d.d. H₂O, CHCl₃/MeOH (1:1), and acetone, and air-dried in a hood.

5. Coprecipitation with (β-D-Galactosyl)₃-Yariv reagent

Three hundred μl of each sample (100 μg/300 μl distilled water) were mixed with 300 μl of (β-D-Galactosyl)₃-Yariv reagent (1 mg/ml in 2% [w/v] NaCl aqueous solution). The mixture was incubated at room temperature for 1 hour followed by pelleting the precipitate in a microfuge. The pellets were washed with 2% NaCl solution, then redissolved in 0.1 M NaOH. The absorbance of the resulting
solution was measured at 420 nm [12]. The characterized glycomodule (AlaHyp)_{51} was used as a standard AGP [12].

6. Glycosyl composition analysis of purified glycomodules and cell wall preparations
For neutral sugar analysis, 60 µg of each glycomodule or 100 µg of each cell wall preparation were hydrolyzed in 200 µl of 2N TFA at 121°C for 1 hr. The neutral monosaccharides were analyzed as alditol acetate derivatives by gas chromatography as described [12]. Uronic acids were quantified by colorimetric method using m-hydroxydiphenyl [23].

7. Amino acid composition and Hyp content analysis of purified glycomodules and cell walls
The amino acid compositions of glycomodules were analyzed as reported [24]. Hyp contents in cell walls were measured as described earlier [25].

8. Tyr, IDT, and di-IDT measurement of cell wall preparations
The Tyr, IDT, and di-IDT amounts of each cell wall preparation were analyzed as described earlier [17]. One hundred µg of each cell wall preparation were hydrolyzed in 200 µl of 6 N HCl with 10 mM phenol at 110°C for 20 hr. Hydrolysates were dried under nitrogen gas. The residues were separated on a polyhydroxyethyl A (10 nm, 9.4 × 200 mm, Poly LC Inc) gel filtration column [17]. The IDT and di-IDT in each cell wall preparation were quantified by comparing with IDT and di-IDT standards (gift from professor Stephen Fry, University of Edinburgh).

9. In vitro cross-linking of EGFP fusion proteins and glycomodules
In vitro cross-linking reactions of the glycomodules or fusion glycoproteins were carried out using the tomato pI 4.6 extensin peroxidase fraction isolated from tomato cell suspension cultures as reported earlier [17, 26].

10. Confocal microscopy study
Suspension cultured cells were checked for green fluorescence under a Zeiss LSM 510 laser-scanning confocal microscope, with excitation wavelength at 488 nm and emission wavelength at 510 nm. The cells were inspected either in SH culture media or in 1 M mannitol for plasmolysis.

11. Glycome profiling of cell wall preparations
Extraction of cell walls with 50 mM ammonium oxalate, 50 mM sodium carbonate, 1 M KOH, 4 M KOH, sodium chlorite in acetic acid, and post chlorite
4 M KOH were carried out following the method described earlier [27, 28]. Glycome profiling of wall glycans in the extracts using ELISA analysis to monitor the glycan epitopes were carried out as reported [27, 28]. ELISA assays were done on an equal sugar basis. All ELISA wells were coated with 15 μl of a stock solution containing 20 μg/ml carbohydrate (glucose equivalents).

12. Analysis of glucose release from cell wall preparations
Cellulase from Trichoderma reesei ATCC 26921 and Novozyme 188 from Aspergillus niger were used. Enzymatic hydrolysis of cell walls was performed on a 1% (w/v) suspension in 50 mM citrate buffer (pH 4.8) with cellulase and β-glucosidase loadings of 20 FPU/g and 40 CBU/g, respectively. The mixture was incubated at 50°C under continuous agitation at 150 rpm with antibiotic antifungal solution (1%, v/v) added. Samples of hydrolysis liquid (0.5 mL) at time intervals of 1, 2, 4, 12, 24, and 48 hr were withdrawn and the hydrolysis was quenched by submersion for 10 min in a vigorously boiling water bath. The liquid samples were then immediately frozen at −20°C until analysis on an Agilent 1200 series HPLC system (Agilent Technologies) equipped with an autosampler and an Aminex HPX-87H column and pre-column (Bio-rad Laboratories). The analysis was carried out at 65°C using 10 mM nitric acid as eluent at a flow rate of 0.6 mL min⁻¹ and with refractive index detection. The calibration of the system was performed with glucose standards.

13. Solid state NMR analysis
The cell wall samples were characterized by solid state cross polarization/magic angle spinning (CP/MAS) ¹³C nuclear magnetic resonance (NMR) using a Bruker Avance III-400 spectrometer operating at 100.59 MHz for ¹³C nuclei as described [29]. The sample (~60% moisture) was packed in a 4 mm cylindrical ceramic MAS rotor and spun at 10 kHz. The CP/MAS NMR spectra were acquired using a 5 μs (90°) proton pulse, 1.5 ms contact time, 4.0-s recycle delay and 8–12 k scans. Bruker’s TopSpin 2.1 software and NUTS NMR software (Acorn NMR, Inc., Livermore, CA) were used for processing the NMR spectral data.

14. Measurement of the radius of gyration of glycomodules
The effect of temperature on glycomodule conformations was measured using a DynaPro Titan and ProteinSolutions Temperature controlled MicroSmapler (Wyatt Technology, Santa Barbara, CA). Five hundred μg of each glycomodule were dissolved in 1 ml of d.d. water. Radii of gyration were measured at 5°C, 8°C, 10°C, 14°C, 20°C, and 24°C. Five readings were taken at each temperature.
Results

1. Each transgenic tobacco cell line produced the corresponding fusion glycoprotein

The \( SS^{\text{tob}}\)-224-\( \text{EGFP} \), \( SS^{\text{tob}}\)-242-\( \text{EGFP} \), and \( SS^{\text{tob}}\)-442-\( \text{EGFP} \) genes cassettes were cloned into a \( \text{pBI121} \) binary vector and transformed into suspension cultured tobacco BY2 cells (Fig. 1A and 1B). After selection via kanamycin resistance, at least two cell lines of each construct were cultured independently in liquid culture media. The green fluorescent fusion glycoproteins were purified from the culture media as described earlier \[12\]. The corresponding fusion glycoproteins 224-\( \text{EGFP} \), 242-\( \text{EGFP} \), and 442-\( \text{EGFP} \) were treated with trypsin to cleave the \( \text{EGFP} \) tag. The resulting 224, 242, and 442 hybrid glycomodules (Fig. 1B) were further purified via size exclusion (SEC) and reversed phase chromatography as previously reported \[12, 17\]. The retention times of the glycomodules on SEC were consistent with the estimated molecular weights of 55.9 kDa, 68.7 kDa, and 68.7 kDa for 242, 224, and 442, respectively, calculated based on the masses of the polypeptides and the attached glycans (Fig. 1C) \[12, 17\].

Amino acid composition analyses revealed that the mole percent of each amino acid in each glycomodule was close to the predicted composition, except for the higher content of Hyp and lower amount of Pro (Table 1). This may have resulted from the hydroxylation of Pro residues in the elastin motifs (Fig. 1B). The amino acid compositions indicated the glycomodules were the expected gene products and were consistent with earlier results obtained from expression of other extensin and AGP analogs in tobacco cells \[11–13, 17\].

The 224, 242, and 442 glycomodules reacted with \( \beta \)-Gal Yariv regent, the dissolved precipitate giving the following absorbances at 420-nm: 0.432, 0.302, and 0.210, respectively, compared to 0.305 of a synthetic AGP glycomodule, (AlaHyp)\(_{51}\) \[12\]. This indicated that the AGP motifs in 224, 242 and 442 were glycosylated with type II arabinogalactans (AG). Glycosyl composition analyses demonstrated that 224, 242 and 442 all contained mainly arabinose, with lesser amounts of Gal, Rha, and uronic acid residues (Table 2). These results were consistent with the Hyp-contiguity hypothesis, which predicts the AG addition to the Hyp residues in the AlaHypAlaHypAlaHypAla motifs and attachment of oligoarabinosides to Hyp residues in the SerHyp\(_{4}\) extensin motifs \[30\].

Earlier work demonstrated that tomato pI 4.6 extensin peroxidase can catalyze in vitro cross-linking of P3 extensin analog YK\(_{8}\) and YK\(_{20}\) \[17\]. To test the cross-linking possibilities of 224, 242, and 442, each glycomodule was incubated with hydrogen peroxide and the pI 4.6 peroxidase isolated via DEAE-anion exchange chromatography as reported by Held et al. \[17\]. The reaction mixture was fractionated on an analytical Superose-6 size exclusion column to monitor the time-dependent decrease of 224, 242 and 442 monomers and the concomitant increase of multimers. The cross-linking rate was determined from calculating the loss of monomer during the reactions \[17\]. In the presence of 1 \( \mu \)g of peroxidase, the 224 and 242 glycomodules shared a cross-linking reaction rate of about 2,000 \( \mu \)g glycomodule/min, while that of 442 was nearly 4,000 \( \mu \)g/min, a rate
comparable to that of the YK₈ P₃ extensin analog reported earlier. This result was consistent with our earlier results that larger extensin analogs cross-linked faster than smaller ones [17].

Although elastins share an extended 3-dimensional structure, elastins fold to an ordered beta-spiral structure when the environmental temperature is higher than a certain point, the transition temperature [19]. If the fused elastin motifs in 224, 242, and 442 glycomodules would inherit this phase transition feature, the sizes of these glycomodules hence their possible stereo-structure in the walls would be temperature-dependent. We measured the radius of gyration of each glycomodule at different temperatures using dynamic light scattering [31]. When the temperature was increased from 20°C to 24°C, the average radii of 224 and 242 increased from 25 nm to 180 nm and from 40 nm to 80 nm, respectively, while the average size of 442 decreased to 60 nm (Fig. 2). The control extensin analog, APYK₂₀ (hybrid glycoprotein (AlaHyp)₄(SerHyp₄SerHyp₄TyrTyrTyrLys)₂₀) [32], exhibited the same behavior as 224 and 242 after treatment. However, the standard deviations of the 224 and 242 radii measured at 24°C were much greater than that of the 442 radius, indicating that the 224 and 242 glycomodules may be responding to temperature in a broader size range at 24°C than at 20°C.

### Table 1. Amino acid composition of three glycomodules isolated from the culture media of individual transgenic cell line.

<table>
<thead>
<tr>
<th>Amino acids</th>
<th>224</th>
<th>242</th>
<th>442</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Measured</td>
<td>Putative</td>
<td>Measured</td>
</tr>
<tr>
<td>O</td>
<td>37.0</td>
<td>32.8</td>
<td>30.1</td>
</tr>
<tr>
<td>S</td>
<td>11.6</td>
<td>10.4</td>
<td>8.9</td>
</tr>
<tr>
<td>G</td>
<td>18.2</td>
<td>17.9</td>
<td>23.6</td>
</tr>
<tr>
<td>A</td>
<td>8.1</td>
<td>6.0</td>
<td>9.9</td>
</tr>
<tr>
<td>P</td>
<td>1.6</td>
<td>6.0</td>
<td>2.6</td>
</tr>
<tr>
<td>Y</td>
<td>4.8</td>
<td>9.0</td>
<td>3.8</td>
</tr>
<tr>
<td>V</td>
<td>15.8</td>
<td>14.9</td>
<td>19.6</td>
</tr>
<tr>
<td>K</td>
<td>2.9</td>
<td>3.0</td>
<td>1.6</td>
</tr>
</tbody>
</table>

The data were shown as molar percentage and were average of two data sets. Amino acid residues in total weight percentage (w/w): 224, 13.5%; 242, 11.6%; 442, 11.7%. The low Y contents were due to the easy degradation of Y residues during hydrolysis.

doi:10.1371/journal.pone.0115906.t001

### Table 2. Sugar composition of three glycomodules isolated from the culture media of individual transgenic cell line.

<table>
<thead>
<tr>
<th>Sugar residues</th>
<th>224</th>
<th>242</th>
<th>442</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rha</td>
<td>10.2</td>
<td>10.0</td>
<td>9.9</td>
</tr>
<tr>
<td>Ara</td>
<td>57.5</td>
<td>47.6</td>
<td>54.8</td>
</tr>
<tr>
<td>Gal</td>
<td>24.7</td>
<td>33.7</td>
<td>26.7</td>
</tr>
<tr>
<td>Uronic acid</td>
<td>7.6</td>
<td>8.7</td>
<td>8.6</td>
</tr>
</tbody>
</table>

The data were shown as molar percentage. The presented numbers were average of two analyses.

doi:10.1371/journal.pone.0115906.t002
the 442 glycomodule might exhibit an elastin-like transition in response to temperature change.

2. Physical changes of the cell walls of transgenic tobacco cells

Cells expressing 224-EGFP and 242-EGFP yielded up to 15 mg of fusion glycoproteins per liter of culture media. In contrast, the 442-EGFP transgenic tobacco cells produced 442-EGFP glycoprotein in a yield of more than 70 mg/L of culture media. Indeed, the culture medium of 20-day-old 442-EGFP cells was dark green due to the accumulation of secreted EGFP fusion protein.

The incorporation of EGFP fusion proteins into the cell walls was monitored by confocal microscopy and EGFP labeling of cell walls was evident (Fig. 3). It was also confirmed by the green fluorescence retained by the isolated 442-EGFP cell walls (Fig. 3G and 3H). Furthermore, the high production of 442-EGFP led to partial separation of the plasma membrane from the walls of the 442-EGFP cells when those cells were checked under normal culture conditions (Fig. 3C and 3D). After plasmolysis in 1M mannitol, the 442-EGFP fluorescence was distributed between the plasma membrane and the 442-EGFP cell wall (Fig. 3E). The Hechtian threads were evident under plasmolyzing conditions (Fig. 3F) but absent under normal culture conditions (Fig. 3D), suggesting the green

![Fig. 2. Temperature and molecular radius relationship of the glycomodules.](image)
fluorescence distribution in Fig. 3C resulted from accumulation of the EGFP fusion protein in that area.

The deposition of the fusion proteins into the cell walls prompted us to further check the wall physical properties and chemical composition. One striking feature was the change of cell wall volume of 442-EGFP cells, compared to that of wild type tobacco cells and walls isolated from the other transgenic lines, that arose when wall preparations were washed with NaCl solution and water (see methods). Specifically, after breaking 40 grams of filtered fresh cells by sonication, each preparation was centrifuged to pellet the crude walls. Initially, the packed cell wall volume (PCV) after centrifugation did not vary significantly between the screened lines.
cell lines (about 15 ml PCV for WT, 27 ml for 224-EGFP, and 19 ml for 242-EGFP cell walls), and nor were differences in PCV observed after washes with 1M NaCl (Table 3). However, the 442-EGFP cell walls swelled dramatically in response to the water washes, the PCV increasing to 66 ml from 19 ml. Furthermore, the washed and pelleted 442-EGFP walls were not ‘sticky’ as they could be poured easily, while the other walls after centrifugation formed pellets that were clumped and would not pour.

### 3. Cell wall glycan changes

To further check the effect of extensin fusion protein deposition on the wall, glycome-profiling of cell wall extracts was used to compare the wall glycan epitopes between the transgenic versus wild type cells [27, 28]. Each cell wall preparation was extracted sequentially with 50 mM oxalate, 50 mM carbonate, 1M KOH, 4M KOH, acidic chlorite, and 4M KOH, which mainly extracts pectins (oxalate and carbonate), xyloglucans and xylans with varying amounts of strongly bound pectin (1 M and 4 M KOH), lignin associated glycans (chlorite), and remaining tightly bound wall glycans (post chlorite 4 M KOH). The amount of material (measured as dry weight) released from the cell walls of each cell line by each extractant varied significantly (Fig. 4, top bars), suggesting that the wall structures of all the transgenic lines were modified compared with WT walls. These extracts were ELISA-screened using 155 plant cell wall glycan-directed monoclonal antibodies. The results, presented in a heatmap, showed that the epitope compositions and extractabilities of xylan, glucan, homogalacturonan (HG) backbone, and rhamnogalacturonan-I (RG-I) backbone did not change significantly across all the samples. However, two obvious altered patterns were observed after comparing the heatmaps of the transgenic cell lines versus WT. The first altered pattern is that the oxalate and carbonate extracts from walls of the 442-EGFP cells contained more arabinogalactan (AG) epitopes, including epitopes in the antibody groups AG1 to AG4 (dotted white box, Fig. 4) [27, 28], than WT, 224-EGFP, and 242-EGFP cell walls. The oxalate and carbonate extracts of 442-EGFP cells also had increased AG/RG-I epitopes (dotted white box, Fig. 4).

Because the total weight of the oxalate and carbonate extracts from 442-EGFP walls was comparable to that of wild type, these data suggest that the extracts from 442-EGFP cell walls contained more easily extractable arabinogalactan-containing
polymers, consistent with the incorporation/deposition of 442-EGFP fusion proteins into the walls which were released, at least partially, from the walls with oxalate and carbonate extractions.

The second altered pattern is the dramatically increased presence of non-fucosylated xyloglucan epitopes in the chlorite extracts prepared from the 224-EGFP or 242-EGFP cell walls (dotted green box, Fig. 4) compared to the chlorite extracts from WT and 442-EGFP walls. These data suggest that the degree of xyloglucan cross-linking in 224-EGFP and 242-EGFP cell walls was modified as a consequence of overexpression of the 224-EGFP and 242-EGFP extensin fusion proteins.

4. Chemical changes in the transgenic tobacco cell walls

The modification of cell walls through extensin fusion protein overexpression was further corroborated by analyses of Hyp and protein contents in the walls. Colorimetric Hyp assays showed that the 442-EGFP cell walls contained significantly more Hyp (4.7 μg/mg of wall) than did the wild type walls (0.9 μg/mg of wall) (Table 4). The amount of Hyp in the 242-EGFP walls also increased significantly. However that of the 224-EGFP cell walls remained comparable to wild type walls (Table 4). Although amino acid composition analyses showed similar compositions for all walls analyzed (Table 5), the absolute amount of each amino acid was significantly higher in the 442-EGFP walls with the total amino acid/protein content of 442-EGFP walls accounting for 16.2% (w/w) of the dry wall weight, comparing to 2.9% (w/w) protein in wild type cell walls. This indicates that overexpression of 442-EGFP significantly increased the protein content of the wall, which might increase the hydration capacity of these cell walls and hence cause the PCV increase.

Pulcherosine and di-IDT are likely intermolecular cross-linking amino acids in extensins [14, 15]. The separation and quantification of Tyr and Tyr derivatives from the wall hydrolyzates showed a significant increase of di-IDT/IDT content in the 442-EGFP walls. Wild type walls contained 1.92 nmol di-IDT/2.19 nmol IDT per mg of walls compared 37.17 nmol di-IDT/7.38 nmol IDT per mg of 442-EGFP cell wall (Table 4). Interestingly, no significant amount of pulcherosine was detected in these samples. The 242-EGFP cell walls also exhibited increased amounts of Tyr and Tyr derivatives (Table 4). However, the amount of Tyr derivatives in 224-EGFP walls decreased with respect to wild type cell walls. These results suggest
that the four tandem repeats of SerHyp₄SerHypSerHyp₄TyrTyrTyrLys P₃ extensin motifs of 442-EGFP might allow more efficient cross-linking than 224-EGFP and 242-EGFP which each contain only two P₃ extensin motifs in tandem (Fig. 1). These results are consistent with our speculation that a small portion of the 442-EGFP fusion glycoproteins that saturated the walls might have been cross-linked via the di-IDT cross-linking amino acids.

To further check the chemical basis of wall property changes, we analyzed the glycosyl residue compositions of the transgenic cell walls. Analyses of neutral sugars hydrolyzed with 2 N TFA and uronic acid residues hydrolyzed with concentrated H₂SO₄ showed that the 442-EGFP cell walls contained similar amount of total monosaccharides as wild type (Table 6). Given the protein

<table>
<thead>
<tr>
<th>Cell wall type</th>
<th>Hyp (µg)</th>
<th>di-Idt (nmol)</th>
<th>Idt (nmol)</th>
<th>Tyr (nmol)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wild Type</td>
<td>0.91</td>
<td>1.92</td>
<td>2.19</td>
<td>5.11</td>
</tr>
<tr>
<td>442-EGFP</td>
<td>4.70</td>
<td>37.17</td>
<td>7.38</td>
<td>25.89</td>
</tr>
<tr>
<td>242-EGFP</td>
<td>2.93</td>
<td>3.62</td>
<td>2.37</td>
<td>7.57</td>
</tr>
<tr>
<td>224-EGFP</td>
<td>1.10</td>
<td>0.81</td>
<td>1.32</td>
<td>2.63</td>
</tr>
</tbody>
</table>

One mg of each sample was used for analysis. The presented numbers were average of two analyses.

doi:10.1371/journal.pone.0115906.t004

Table 5. Amino acid composition of each cell wall (CW) preparation.

<table>
<thead>
<tr>
<th>Amino acid Residue</th>
<th>Wild type CW</th>
<th>442-EGFP CW</th>
<th>242-EGFP CW</th>
<th>224-EGFP CW</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>nmol</td>
<td>mol%</td>
<td>nmol</td>
<td>mol%</td>
</tr>
<tr>
<td>D</td>
<td>1.05</td>
<td>9.8</td>
<td>6.47</td>
<td>10.8</td>
</tr>
<tr>
<td>E</td>
<td>1.01</td>
<td>9.4</td>
<td>6.14</td>
<td>10.2</td>
</tr>
<tr>
<td>O</td>
<td>0.69</td>
<td>6.4</td>
<td>2.10</td>
<td>3.5</td>
</tr>
<tr>
<td>S</td>
<td>0.84</td>
<td>7.8</td>
<td>4.47</td>
<td>7.4</td>
</tr>
<tr>
<td>G</td>
<td>0.79</td>
<td>7.3</td>
<td>5.08</td>
<td>8.4</td>
</tr>
<tr>
<td>H</td>
<td>0.21</td>
<td>2.0</td>
<td>1.13</td>
<td>1.9</td>
</tr>
<tr>
<td>R</td>
<td>0.34</td>
<td>3.2</td>
<td>2.42</td>
<td>4.0</td>
</tr>
<tr>
<td>T</td>
<td>0.43</td>
<td>1.0</td>
<td>3.11</td>
<td>5.2</td>
</tr>
<tr>
<td>A</td>
<td>0.89</td>
<td>8.3</td>
<td>5.41</td>
<td>9.0</td>
</tr>
<tr>
<td>P</td>
<td>0.78</td>
<td>7.3</td>
<td>3.80</td>
<td>6.3</td>
</tr>
<tr>
<td>Y</td>
<td>0.29</td>
<td>2.7</td>
<td>1.69</td>
<td>2.8</td>
</tr>
<tr>
<td>V</td>
<td>0.70</td>
<td>6.5</td>
<td>3.86</td>
<td>6.4</td>
</tr>
<tr>
<td>M</td>
<td>0.13</td>
<td>1.2</td>
<td>0.16</td>
<td>0.3</td>
</tr>
<tr>
<td>I</td>
<td>0.58</td>
<td>5.4</td>
<td>2.63</td>
<td>4.4</td>
</tr>
<tr>
<td>L</td>
<td>0.92</td>
<td>8.6</td>
<td>5.28</td>
<td>8.8</td>
</tr>
<tr>
<td>F</td>
<td>0.46</td>
<td>4.3</td>
<td>2.51</td>
<td>4.2</td>
</tr>
<tr>
<td>K</td>
<td>0.64</td>
<td>5.9</td>
<td>3.90</td>
<td>6.5</td>
</tr>
<tr>
<td>% (w/w) dry CW</td>
<td>2.9%</td>
<td>16.2%</td>
<td>4.3%</td>
<td>1.2%</td>
</tr>
</tbody>
</table>

Each data set was collected from 40 µg of sample. O stands for hydroxyproline. The presented numbers were average of two analyses.

doi:10.1371/journal.pone.0115906.t005
content in each wall preparation, we calculated that the total acid-hydrolysable carbohydrates account for 50.7% (w/w) of the non-protein portion of the 442-EGFP walls, comparable to the 50.9% (w/w) for wild type walls. However, compared to wild type walls, the 442-EGFP walls generally had less individual hydrolysable monosaccharides except for glucosyl residues, even after correction for the protein contents. Interestingly, the 224-EGFP and 242-EGFP walls contained significantly more glucosyl residues than wild type, with other monosaccharide compositions comparable to those of wild type and/or 442-EGFP walls.

Because TFA cannot hydrolyze crystalline cellulose, the different cell wall preparations were treated directly with cellulase and β-glucosidase to monitor the release of cellulosic glucose, which was quantified via HPLC cation exchange chromatography. Over the first five-hours of digestion, slightly more glucose was released from the walls of the 242-EGFP cells and both 442-EGFP cell lines than from WT walls (Fig. 5). When the digestions were extended to more than 12 hours, the cell walls of two 442-EGFP lines and the 242-EGFP cell walls yielded less glucose than the wild type walls (Fig. 5). However, the 224-EGFP walls released much greater amounts of glucose than wild type walls at all incubation times, with a 70% (w/w) glucose yield after a one-day digestion. To further check if the crystallinity of each transgenic cell line was changed, one-dimensional 13C solid state NMR spectra were collected with the walls of transgenic cells versus WT (Fig. 6). The chemical shifts and relative signal intensities of anomeric carbons of cellulosic glucosyl residues in the 242-EGFP and 442-EGFP cell walls were comparable to those of wild type cell walls. However, the 224-EGFP cell walls showed weaker C-4 signals of crystalline cellulose than the other walls (Fig. 6, boxes), suggesting lower cellulose crystallinity in the 224-EGFP walls.

### Table 6. Acid-hydrolysable sugar composition of cell walls.

<table>
<thead>
<tr>
<th>Sugar residue</th>
<th>Wild type cell wall</th>
<th>442-EGFP cell wall</th>
<th>242-EGFP cell wall</th>
<th>224-EGFP cell wall</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>µg</td>
<td>nmol</td>
<td>µg</td>
<td>nmol</td>
</tr>
<tr>
<td>Rha</td>
<td>80.6</td>
<td>491</td>
<td>41.3</td>
<td>252</td>
</tr>
<tr>
<td>Fuc</td>
<td>17.3</td>
<td>105</td>
<td>3.0</td>
<td>18.0</td>
</tr>
<tr>
<td>Ara</td>
<td>140</td>
<td>933</td>
<td>99.2</td>
<td>661</td>
</tr>
<tr>
<td>Xyl</td>
<td>66.5</td>
<td>443</td>
<td>50.1</td>
<td>334</td>
</tr>
<tr>
<td>Man</td>
<td>21.8</td>
<td>121</td>
<td>15.5</td>
<td>86.0</td>
</tr>
<tr>
<td>Glc</td>
<td>63.1</td>
<td>351</td>
<td>153</td>
<td>852</td>
</tr>
<tr>
<td>Gal</td>
<td>105</td>
<td>582</td>
<td>63.2</td>
<td>351</td>
</tr>
<tr>
<td>Uronic acid</td>
<td>274</td>
<td>1398</td>
<td>134</td>
<td>684</td>
</tr>
<tr>
<td>% (w/w)</td>
<td>49.4%</td>
<td>42.5%</td>
<td>49.2%</td>
<td>65.1%</td>
</tr>
</tbody>
</table>

Each set of data was collected from 1 mg of sample. The presented numbers were average of two analyses.

doi:10.1371/journal.pone.0115906.t006
Fig. 5. Sugar release of wild type versus 224-EGFP, 242-EGFP, and 442-EGFP cell walls. Cell walls were digested with cellulase and glucosidase. The glucose contents in hydrolysis solution withdraw at time intervals of 1, 2, 4, 12, 24, and 48 hour were analyzed. The calibration of the system was performed with glucose standards. The 224-EGFP cell walls showed a better sugar release performance than 242-EGFP, 442-EGFP, and wild type cell walls.

doi:10.1371/journal.pone.0115906.g005

Fig. 6. Solid state one dimensional $^{13}$C NMR spectra of walls of transgenic cells versus WT walls. The chemical shifts and relative signal intensities of anomic carbons (i.e., $C_1$) of glucosyl residues of cellulose in 242-EGFP and 442-EGFP cell walls were comparable to those of wild type cell walls. However, 224-EGFP cell walls displayed much stronger signals of anomic $^{13}$C of xylosyl residues (oval) and lower signal of crystalline cellulose (boxes).

doi:10.1371/journal.pone.0115906.g006
Discussion

Phenotype-chemotype changes of the three transgenic cell lines

Overexpression of the three extensin analogs in tobacco BY2 cells resulted in changes of primary cell wall properties. The wall phenotype-chemotype changes can be classified into three types based on the results of our physical and chemical analyses. One type of change was observed in the walls of 224-EGFP cells, which exhibited decreased cellulose crystallinity (Fig. 6). The decreased cellulose crystallinity appears to be the primary factor leading to the reduced recalcitrance of the 224-EGFP walls as measured by the significantly greater amounts of cellulosic glucose released from the transgenic walls after cellulase treatment compared with WT walls (Fig. 5). In addition, unlike the WT walls, the 224-EGFP walls contained more xyloglucans in the chlorite wall extracts (Fig. 4), indicating that the expression of the 224-EGFP fusion protein led to cross-linking of xyloglucans to chlorite-susceptible polymers or substituents, which has never been observed in WT Arabidopsis walls. In addition, there appears to be an increase in the amount of xyloglucan in the walls of this transgenic cell line. The latter is consistent with both the solid state NMR data showing an increase of the Xyl anomeric carbon signal (Fig. 6, oval) and the sugar composition analysis documenting that the 224-EGFP walls contained five-times more non-cellulosic glucose than WT (Table 6).

A second type of change was observed in the 242-EGFP walls. Like the 224-EGFP walls above, the 242-EGFP walls contained chlorite-extractable non-fucosylated xyloglucan epitopes (Fig. 4) that are not found in chlorite extract of WT walls. However, the 242-EGFP walls did not exhibit any changes in cellulose crystallinity (Fig. 6), nor did they exhibit reduced recalcitrance (as determined by cellulase-releasable glucose) (Fig. 5). Thus, the cross-linking of some xyloglucans to chlorite-susceptible wall components appears not to be a factor in recalcitrance.

A third type of wall change was exhibited by the 442-EGFP cell walls. Walls from the 442-EGFP cells showed a WT-like pattern of non-fucosylated xyloglucan epitopes in their glycome-profiles, but significantly more AG and AG/RG-I epitopes in the oxalate and carbonate extracts (Fig. 4). More dramatically, the 442-EGFP cell lines secreted substantially more of the extensin fusion protein into the culture supernatant than did either of the other two transgenic cell lines. Furthermore, 442-EGFP walls exhibited a dramatically increased packed cell wall volume (PCV) after water washes, which was not exhibited by either WT or the other transgenic cell walls. The 442-EGFP walls also showed no reduced recalcitrance (Fig. 5), suggesting that the increased PCV phenotype did not affect any wall characteristics associated with recalcitrance.

Possible mechanisms underlying the observed changes in phenotype-chemotype

The high production and secretion of the 442-EGFP fusion protein driven by the CaMV S35 promoter led to extensive accumulation of the EGFP-extensin fusion
protein in the culture medium; the medium turned dark green after two-weeks of culturing. This result is consistent with our earlier reports that AGP glycosylation facilitates secretion of exogenous fusion proteins into culture media \[33, 34\]. We also observed the localization of green fluorescence between the plasma membranes and the cell walls of these cells (Fig. 3C and 3D), suggesting the overexpression of the 442-EGFP gene construct results in saturation of the culture medium, the interspace between plasma membrane and cell wall, and the walls themselves with the 442-EGFP fusion protein. In addition, saturation of the walls with 442-EGFP, as reflected in higher (16\%) protein content of the walls, might also saturate the extensin peroxidases that are responsible for cross-linking of functional endogenous extensins. The lack of endogenous wall extensin network consequently resulted in loosen cell walls, contributing to the increased PCV or hydration capacity of the 442-EGFP walls.

In contrast, overexpression of the 224-EGFP and 242-EGFP gene constructs resulted in much lower accumulation of the corresponding fusion proteins in the culture media than was observed for the 442-EGFP cells. We did not observe EGFP fluorescence saturated 224-EGFP and 242-EGFP cells as we documented in Fig. 3C. In addition, the 224 and 242 modules are less cross-linkable based on cross-linking rates measured in vitro for 224, 242, and 442, suggesting that 224-EGFP and 242-EGFP might not be ideal substrates for the extensin peroxidases thought to be responsible for cross-linking extensins in the wall. Thus, the endogenous extensins in these two transgenic cell lines could form regular extensin network for wall integrity, which might be why the 224-EGFP and 242-EGFP cells showed similar PCV as WT cells.

Furthermore, although the 224 and 442 glycomodules shared the same molecular mass and the same total content of each motif, including 8 YK cross-linking motifs, 4 AOAOAOA AGP motifs, and 16 VPGVP elastin motifs, the faster in vitro cross-linking rate of 442 suggests the two (YK)\(_4\) units in 442 were more cross-linkable than the four (YK)\(_2\) units in 224. This might be a reason that the 442-EGFP cell walls contained more di-IDT content than WT or the other transgenic cell lines. However, why did the 242-EGFP cell walls contain more di-IDT than the 224-EGFP cell walls as 242 had only two (YK)\(_2\) units? One possible reason is that the more clustered AGs on 242 glycomodules might be a favorite factor for in vivo cross-linking given the adhesion property of AGs favoring intermolecular adhesion and alignment \[35\]. Moreover, why did overexpression of 224-EGFP in the 224-EGFP cells result in less di-IDT content than WT? The elastin motifs in 224-EGFP might play a central role in this case, most likely due to the observed wide-range size change of the 224 glycomodule at 24°C attributed to the stretch-fold conformations of the 16 VPGVG motifs in each 224-EGFP. The conformations of elastin motifs in each extensin fusion protein might introduce some steric space into the walls, which would explain why all the transgenic cell walls released more cellulose glucose than WT during the first five-hour cellulase digestion, as the cellulases could more easily access cellulose microfibrils near the steric spaces of those transgenic walls (Fig. 5). Overall, the total effect of each
extensin fusion protein on its transgenic cell walls was a collective effect of all the three motifs.

Besides the change in hydration properties, the biomass of 442-EGFP walls increased to thrice that of wild type walls (Table 3). This may result from either the deposition of 442-EGFP fusion glycoproteins, or increased endogenous wall polysaccharides in the walls, or both. The significantly higher Hyp and Tyr contents in the 442-EGFP walls (Table 4), as well as greater than five-fold increase in protein content compared to that of wild type walls is consistent with the biomass increase being due, at least partially, to the over-deposition of 442-EGFP. However, we also observed that the biomass of 224-EGFP and 242-EGFP cell walls almost doubled compared to that of wild type cell walls (Table 3). Although the Hyp and protein contents of 242-EGFP walls were slightly higher than that of wild type, the amounts of Hyp and protein in 224-EGFP walls were even less than those of wild type. Therefore, the biomass increase in these two cases might mainly result from the increase of endogenous wall polysaccharides.

Finally, identifying peptide fragments of the extensin analogs from the walls of transgenic cells would provide direct evidence for their wall deposition. However, LC-MS/MS analysis of the total tryptic peptides generated from hydrogen fluoride (HF)-deglycosylated 442-EGFP cell walls did not yield any peptide sequence attributable to 442-EGFP. It may be due to either the cross-linking of 442-EGFP in the walls (evident in high content of di-IDT) or the large size of the tryptic 442 peptides (Fig. 1). Nevertheless, labeling of the walls with green fluorescence, changes of amounts of wall di-IDT and protein, and changes of wall glycans in the walls of transgenic cells are consistent with deposition/incorporation of extensin fusion proteins into the walls.

Conclusion

Overexpression of extensin analogs in tobacco primary cell walls resulted in changes of wall properties that were distinct for each transgenic cell line. The arrangement and sizes of the extensin motifs, AGP motifs, and elastin motifs on the polypeptide affected the effects on overall wall structure for each line. The 442-EGFP walls showed no change in wall cellulose crystallinity; however, wall saturated with 442-EGFP may have prevented the formation of endogenous extensin network, which significantly changed wall physical and chemical properties. In contrast, the 224-EGFP walls exhibited a higher ratio of xyloglucans to celluloses and lower cellulose crystallinity, most likely due to the spatial alterations caused by deposition of the more extended 224-EGFP fusion glycoproteins. However, the 242-EGFP walls only displayed changes in xyloglucan extractability. These results indicate that regulated expression of modified extensins in planta may be a way to manipulate wall architecture to achieve higher biomass and higher biofuel production.
Author Contributions
Conceived and designed the experiments: LT YP UA LC MGH AJR MJK. Performed the experiments: LT YP SP UA JQ AA LC. Analyzed the data: LT YP SP UA JQ AA LC MGH AJR MJK. Contributed reagents/materials/analysis tools: LT YP SP UA JQ AA LC MGH AJR MJK. Wrote the paper: LT. Manuscript revision: LT YP SP UA JQ AA LC MGH AJR MJK.

References


