Abaxinolysis Plays a Crucial Role in Extensin Cross-linking In Vitro

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ABSTRACT: Extensins (EXTs) are hydroxyproline-rich glycoproteins (HRGPs) that are structural components of the plant primary cell wall. They are basic proteins and are highly glycosylated with carbohydrate accounting for >50% of their dry weight. Carbohydrate occurs as monogalactosyl serine and arabinosyl hydroxyproline, with arabinosides ranging in size from ~1 to 4 or 5 residues. Proposed functions of EXT arabinosylation include stabilizing the polypeptide II helix structure and facilitating EXT cross-linking. Here, the involvement of arabinosylation in EXT cross-linking was investigated by assaying the initial cross-linking rate and degree of cross-linking of partially or fully de-arabinosylated EXTs using an in vitro cross-linking assay followed by gel permeation chromatography. Our results indicate that EXT arabinosylation is required for EXT cross-linking in vitro and the fourth arabinosyl residue in the tetraarabinoside chain, which is uniquely α-linked, may determine the initial cross-linking rate. Our results also confirm the conserved structure of the oligoarabinosides across species, indicating an evolutionary significance for EXT arabinosylation.

KEYWORDS: HRGP, extensin, arabinosylation, in vitro cross-linking, RSH, tomato P1, peroxidase

Introduction

Plant cells are surrounded by a thin primary cell wall comprised of interpenetrating networks of polysaccharide (cellulose, hemicellulose, and pectin) and structural glycoproteins (~10% dicot cell wall dry weight).¹ Structural proteins are often rich in hydroxyproline (Hyp) and are hence named hydroxyproline-rich glycoprotein (HRGP).² HRGPs are categorized into three major types: the extensins (EXTs), the arabinogalactan proteins (AGPs), and the proline-rich proteins (PRPs) based on their repetitive peptide motifs and the type and extent of glycosylation, which occurs mainly on the Hyp residues. Despite being a relatively minor component in the primary cell wall compared with the matrix polysaccharides, HRGPs play an important role in wall architecture,³-⁴ plant development,⁵-⁹ embryogenesis,¹⁰,¹¹ stress responses,¹²-¹⁴ and defense against pathogen attacks.¹⁵-¹⁸

EXTs are a major class of HRGP. Hyp accounts for >30 mol% of an EXT’s amino acids¹⁹,²⁰ and occurs in short peptide repeats that alternate with Hyp-poor repeats containing hydrophobic and basic residues. Hydrophobic motifs often contain Tyr residues that participate in intra and intermolecular cross-linking. Ser-(Hyp), is the “signature” repeat motif in EXTs with Ser being monogalactosylated²¹ and all the Hyp residues O-arabinosylated (Hyp–O–Ara)²²-²⁴ EXTs are highly basic²⁵ which cleaves glycoconjugates.³³ EXT cross-linking leads to the formation of a covalently linked protein network that is somehow associated peroxidases.³³-³⁴ The cross-linking motifs contain Tyr residues, commonly in the sequence of Tyr–X–Tyr (X usually = Lys, Tyr, Leu, or Val)³² and possibly Val–Tyr–Lys.³³ The Tyr–X–Tyr motif gives rise to intramolecular isodityrosine (Idt),³⁴,³⁵ which can itself undergo further cross-linking to produce intermolecular di-isodityrosine (Di-Idt),³⁶,³⁷ or pulcherosine³⁸ cross-links. The cross-linking of EXTs (with other EXTs or possibly other wall structural proteins) is likely catalyzed by wall-associated peroxidases.³³,³⁹ Cross-linking leads to the formation of a covalently linked protein network that is somehow independent of wall polysaccharide networks, since this protein network remains intact after treatment of the wall with anhydrous hydrogen fluoride (HF),⁴⁰-⁴² which cleaves glycosidic bonds but not peptide bonds.⁴³

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EXTs are a major class of HRGP. Hyp accounts for >30 mol% of an EXT’s amino acids¹⁹,²⁰ and occurs in short peptide repeats that alternate with Hyp-poor repeats containing hydrophobic and basic residues. Hydrophobic motifs often contain Tyr residues that participate in intra and intermolecular cross-linking. Ser-(Hyp)
Previously, Schnabelrauch et al.\textsuperscript{[33]} reported that deglycosylated EXT monomers were not cross-linked \textit{in vitro} by EXT peroxidase. This observation suggested a role for arabinosylation in EXT self-assembly and cross-linking, one of the most fundamental functions of EXTs. In this study, we investigated the involvement of arabinosylation in EXT cross-linking and provide evidence suggesting that arabinosylation controls the initial rate and the extent of EXT cross-linking \textit{in vitro}. We also found that \textit{Arabidopsis} oligoarabinosides share a common structure with arabinosides from other species, indicating that these glycosides play a significant role in the evolutionary progression of EXT molecular function.

**Material and Methods**

**Chemicals and reagents.** All general chemical reagents were purchased from Sigma-Aldrich, and all general lab equipment were from VWR unless otherwise indicated.

**Isolation of EXT precursors from suspension culture cells.** Monomeric EXTs from tomato (TOMP1, \textasciitilde 400 aa protein, MW \textasciitilde 110 kDa) or \textit{Arabidopsis} (RSH, 404 aa protein, MW \textasciitilde 100 kDa) were isolated from suspension culture cells using the intact cell elution method as described earlier by Smith et al.\textsuperscript{[38]} Briefly, cells at 25\% packed cell volume (\(V_{	ext{cell}}/V_{	ext{culture}}\)) were harvested by filtration on a course-sintered funnel and eluted with 80 mM AlCl\(_3\) for 15 minutes. The AlCl\(_3\) eluate was collected by filtration, lyophilized, and re-dissolved in 90 mL of water, and 10 mL cold 100\% (w/v) trichloroethanoic acid (TCA) was added to the eluate. The mixture was incubated overnight at 4°C and then centrifuged at 30,000 \(\times\) g. The supernatant was collected, dialyzed using 3500-Da cut-off dialysis tubing (Spectrum Laboratories) against distilled water for 48 hours, and then lyophilized. The resulting TCA-free eluate powder was dissolved at 5 mg/mL in 30 mM sodium phosphate buffer (pH 7.6), and EXTs were purified using a SP Sepharose Fast Flow (GE Healthcare) strong cation exchange chromatography followed by reverse-phase HPLC chromatography using a PRP-1 column (Hamilton) as described by Terneus.\textsuperscript{[20]}

**Amino acid composition of purified EXTs.** The amino acid compositions of proteins were determined by phenylisothiocyanate (PITC) derivatization of acid hydrolyzed proteins as described by Janssen et al.\textsuperscript{[43]} Complete deglycosylation using anhydrous HF. One hour HF deglycosylations were performed as described by Mort and Lamport.\textsuperscript{[40]} After quenching, the HF was removed by blowing down the aqueous mixtures under a stream of N\(_2\) gas twice and then lyophilized. The deglycosylated EXTs were isolated by reverse-phase HPLC fractionation followed by lyophilization. The products were designated as RSHHF and P1HF, respectively.

**Colorimetric arabinose/galactose assay.** The remaining arabinose and galactose contents on EXTs after deglycosylation were measured colorimetrically as described by Fry.\textsuperscript{[45]}

**Hyp-arabinoside profiling.** Hyp-arabinoside profiling was carried out using methods described by Lamport and Miller.\textsuperscript{[21]} Native and partially dearabinosylated (\(\alpha-L\)-arabinofuranosidase or pH 2.0 HCl treated, 2 mg) EXTs were dissolved (5 mg/mL) in saturated (\(\sim 0.2 \) M) Ba(OH)\(_2\) in sealed tubes and hydrolyzed at 105°C for 18 hours. The reactions were then cooled on ice, and the Ba(OH)\(_2\) was neutralized with ice-cold 1 M H\(_2\)SO\(_4\) to a pH of 7.0 (monitored by pH paper). BaSO\(_4\) generated by neutralization was removed from the reactions by centrifugation, and the neutralized reaction mixtures were then loaded onto a strong cation exchange Chromobeads C-2 column (Technicon) equilibrated with water. The column was then washed with a linear gradient of HCl generated by mixing 100 mL of water with 100 mL of 0.5 M HCl at 0.5 mL/minute. Fractions eluted from the C-2 column were collected at 4 minutes/tube, and 0.5 mL of each fraction was assayed for Hyp using the colorimetric Hyp assay as described previously.\textsuperscript{[23]}

**Purification of EXT peroxidase from tomato cell suspension culture.** A pH 4.6 peroxidase was isolated from tomato suspension cell culture using a combination of diethylaminoethyl anion-exchange chromatography and isoelectric focusing as described by Dong et al.\textsuperscript{[46]} The resulting protein fraction was analyzed by SDS–PAGE. Peroxidase activity was determined by reaction with 2,2′-azino-bis-(3-ethylbenzothiazoline-6-sulphonic acid) (ABTS) as described by Everdeen et al.\textsuperscript{[47]}

**In vitro cross-linking of deglycosylated EXTs.** Partially or fully dearabinosylated and fully deglycosylated RSH
and TOMP1 samples were cross-linked in vitro with a crude preparation of the pl 4.6-extensin peroxidase from tomato suspension culture medium. All protein samples were first made into 10 mg/mL stock solutions. Stock solutions (60 μL) were added to 40 μL of 0.1 M citrate buffer (pH 6.0) to form working solutions. The cross-linking reactions contained 10 μL of extensin working solution (60 μg of EXT), 5 μL diluted crude peroxidase containing 0.05 ng of peroxidase based on the ABTS assay, and 5 μL 0.24 mM H₂O₂ (the final concentration of H₂O₂ in reaction system was 60 μM) in a total reaction volume of 20 μL. Native RSH and TOMP1 were cross-linked in the same way as controls. Timing of the reactions was started upon the addition of H₂O₂ and terminated by the addition of the stop reagents: 10 μL of 50 mM β-mercaptoethanol. The initial cross-linking rates were determined using the first-order rate equation \( v_0 = kC_0 \) presented in terms of microgram EXT cross-linked in 1 mL reaction solution per minute as described by Everdeen et al.\(^\text{47} \) where \( v_0 \) is the initial rate, \( k \) is the first-order reaction rate constant, and \( C_0 \) is the initial EXT concentration at zero minute of cross-linking. The rate constant \( k \) was determined by equation \( k = \text{ln}(A_o/A)/(t) \), where \( A_o \) is the peak area of monomeric EXTs cross-linked for zero minute, and \( A \) is the peak area of monomeric EXTs after certain reaction time. The monomeric EXT peaks before and after the cross-linking reactions were assayed by gel filtration chromatography using a Superose 6 10/300 GL column (GE Healthcare), monitored at 220 nm absorbance. Three parallel reactions were run for each sample to determine the initial cross-linking rate (cross-linking for five minutes) and the data averaged. For zero-minute cross-linking reactions, the stop reagents were added before the addition of H₂O₂. Each EXT sample was also cross-linked for 30 minutes for complete reaction. Completely dearabinosylated (treated with pH 1.0 HCl or HF) samples had very slow cross-linking rates, and thus only the 30-minute cross-linking experiment was performed. Additionally, these reactions were succinylated before injecting onto the column by adding 1 mg succinic anhydride into the reaction mixture and incubated for 60 minutes. Succinylation reverses the positive charges on the proteins, which interact with the Superose column matrix and retard protein migration.

**Hyp-arabinosyl linkage analysis of RSH.** Hyp-arabinosyl linkage analysis was performed on RSH at the Center for Plant and Microbial Complex Carbohydrates at the Complex Carbohydrate Research Center (CCRC), University of Georgia. RSH was permethylated, depolymerized, reduced, and acetylated. The resulting partially methylated alditol acetates (PMAAs) were analyzed by gas chromatography-mass spectrometry (GC-MS) as described by York et al.\(^\text{48} \)

**Isolation of Hyp-oligoarabinosides from RSH.** RSH was hydrolyzed with 0.2 M Ba(OH)\(_2\) at 105°C for 18 hours, and the reaction was neutralized with H\(_2\)SO\(_4\) before the Hyp-arabinosides were isolated by cation exchange Chromobeads C-2 column chromatography as described earlier. However, gradient elution was performed by mixing 100 mL of water (elucent A) with 110 mL of 0.7 M trifluoroacetic acid (TFA, eluent B) for >300 minutes. The C-2 column was washed after TFA gradient with 0.7 M TFA to elute Hyp-diarabinoside (Hyp-Ara\(_2\)) for 30 minutes, and TFA was replaced with 0.5 M HCl to elute Hyp-monoarabinoside and non-glycosylated Hyp. The column eluates were collected in four-minute fractions, and 0.1 mL of each fraction was assayed for Hyp using the colorimetric Hyp assay. The fractions containing Hyp-tetraarabinosides (Hyp-Ara\(_4\)) and Hyp-triarabinosides (Hyp-Ara\(_3\)) were lyophilized for three times with water to remove the excess TFA before sending for NMR structural analysis.

**Structure elucidation of Hyp-oligoarabinosides by NMR spectroscopy.** Hyp-Ara\(_4\) (1.7 mg) and Hyp-Ara\(_3\) (2 mg) were each dissolved in 180 mL of D\(_2\)O. Each solution was transferred into a 3 mm NMR tube. NMR data were collected at 25°C on a VNMR 800 instrument equipped with a 3 mm cold probe at the CCRC Facilities. The 1D H NMR and 2D H homonuclear COSY, TOCSY (DIPSI2 mixing time 60 milliseconds), and NOESY (NOE mixing time 100 milliseconds), 2D heteronuclear H\(^1\)-C\(^{13}\) HSQC, and HMBC experiments were performed as described earlier.\(^\text{49} \)

**Results**

**Monomeric EXT isolation.** SP Sepharose fractionation of tomato eluates yielded three peaks (Supplementary Fig. 1A). The void peak and peak 2 (Supplementary Fig. 1A) were relatively poor in Hyp judging by colorimetric Hyp assay (Hyp\(_{\text{void}}\) = 2.4 μg Hyp/mg protein and Hyp\(_{\text{peak2}}\) = 1.6 μg Hyp/mg protein, respectively). Peak 1 contained 38.2 μg Hyp/mg protein and was further fractionated by reverse-phase HPLC, yielding a single peak (Supplementary Fig. 1B). The amino acid composition identified the protein in this peak as EXT precursor 1 (TOMPI) as described earlier by Smith et al.\(^\text{49} \) (Supplementary Table 1).

Similarly, SP Sepharose fractionation of Arabidopsis eluates (Supplementary Fig. 1C) resulted in one major peak, peak 1, containing 33.7 μg Hyp/mg protein, in addition to the void peak (8.3 μg Hyp/mg protein). Peak 1 was further fractionated by reverse-phase HPLC to produce one major peak (Supplementary Fig. 1D). Through the analysis of amino acid composition (Supplementary Table 1), the protein in this peak was identified as RSH, as described earlier.\(^\text{30} \) All minor Sepharose peaks (the small peak group and peak 2, Supplementary Fig. 1C) in the fractionated eluate of Arabidopsis suspension culture were relatively rich in Hyp (Hyp\(_{\text{null peak group}}\) = 23.7 μg Hyp/mg protein and Hyp\(_{\text{peak2}}\) = 11.7 μg Hyp/mg protein, respectively). The protein components in those Sepharose peaks formed precipitate with β-Yariv reagent, indicating the presence of AGP-type structural proteins that may potentially contribute to the high Hyp content; thus, they were not fractionated or analyzed further.

**Arabinosylation affects extensin cross-linking in vitro**
Characterization of deglycosylated EXTs. The removal of Ara residues by each deglycosylation was first assayed using colorimetric arabinose assays. Treatment with pH 1.0 HCl and HF deglycosylation removed almost all arabinosyl residues on the EXTs (Table 1), which was judged by comparison with native RSH and TOMP1. Arabinofuranosidases removed about 30% of Ara residues from RSH and TOMP1, while the HCl treatment at pH 2.0 removed about half the Ara residues from both EXTs. These data are also supported by the decrease in the weight percentage of Hyp-oligoarabinosyl residues in Hyp-Arabinoside (Hyp-Ara) profiles (Table 2).

EXT galactosyl residues are much less susceptible to acid hydrolysis than arabinosyl residues. A previous study has showed that Ser-Gal remained intact after hydrolysis for one hour at 100°C by pH 1.0 HCl. The partial dearabinosylation approaches used here were milder and thus unlikely to remove EXT Gal residues. Thus only the residual Gal content of the pH 1.0 HCl and HF-treated EXTs were determined (Table 1). HF removed almost all the Gal on RSH and TOMP1, while galactose was still present in pH 1.0 HCl-treated EXTs, as expected.

Hyp-Ara profiles of EXTs treated with arabinofuranosidase and pH 2.0 HCl (native RSH and TOMP1 as controls, Fig. 1 and Table 2) were determined, and the results revealed that no Hyp-tetraarabinosides were observed after treatment with arabinofuranosidases, while some Hyp-Ara remained after pH 2.0 treatments (Fig. 1).

In vitro cross-linking of deglycosylated EXTs. The initial cross-linking reaction rates of partially dearabinosylated RSH and TOMP1 were determined by gel filtration chromatography using the first-order rate equation employed earlier by Everdeen et al47: $\nu = kC_0$ with native RSH and TOMP1 as controls. The results are summarized in Table 3.

Samples that essentially lacked Ara did not cross-link at all (pH1 or HF treated), even after 30 minutes of incubation with the pH 4.6 extensin peroxidase (Supplementary Fig. 2). Thus, the initial cross-linking rates of these samples were documented as zero (Table 3). Initial cross-linking rates of the partially dearabinosylated samples were lower than that of the native EXTs. For partially dearabinosylated RSH samples, initial rate decreases of 17% and 68% were observed for $\text{RSH}_{\text{pHH}}$ and $\text{RSH}_{\text{Ara}}$, respectively, compared with fully arabinosylated RSH.

Notes: Hyp-Ara, the number of arabinoses attached to Hyp, $n = 0$ means non-glycosylated Hyp. Ratio of different arabinino-oligosaccharides was expressed as percentage of Hyp bearing those oligosaccharides to total Hyp.

### Table 1. Remaining arabinose and galactose on deglycosylated RSH and TOMP1.

| SAMPLE (µg) | ARABINOSE (µg) | ARABINOSE CONTENT (W/W) |
|------------|----------------|-------------------------|
| RSH        | 30             | 13                      | 43.3%                   |
| RSHArA     | 30             | 8.7                     | 29.0%                   |
| RSHpH2     | 30             | 6.6                     | 21.7%                   |
| RSHpH1     | 30             | 0.4                     | 1.3%                    |
| RSHHF      | 30             | 0.2                     | 0.6%                    |
| TOMP1      | 30             | 13.8                    | 46.0%                   |
| P1Ara      | 30             | 9.9                     | 33.0%                   |
| P1pH2      | 30             | 7.2                     | 24.0%                   |
| P1pH1      | 30             | 0.4                     | 1.3%                    |
| P1HF       | 30             | 0.2                     | 0.6%                    |

| SAMPLE (µg) | GALACTOSE (µg) | GALACTOSE CONTENT (W/W) |
|-------------|----------------|-------------------------|
| RSHpH1      | 100            | 5.1                     | 5.1%                    |
| RSHHF       | 100            | 0.2                     | 0.2%                    |
| P1pH1       | 100            | 3.5                     | 3.5%                    |
| P1HF        | 100            | 0.3                     | 0.3%                    |

### Table 2. Hyp-Ara profiles of partially deglycosylated EXTs.

| Hyp-Ara | RSH | RSHArA | RSHpH2 | TOMP1 | P1Ara | P1pH2 |
|---------|-----|--------|--------|-------|-------|-------|
| n = 4 (%) | 21.4 | 0.0    | 5.8    | 25.4  | 0.0   | 9.7   |
| n = 3 (%) | 20.7 | 16.7   | 4.9    | 10.2  | 25.2  | 5.7   |
| n = 2 (%) | 13.2 | 13.3   | 8.0    | 10.7  | 15.5  | 8.2   |
| n = 1 (%) | 17.3 | 23.2   | 23.0   | 28.1  | 25.7  | 30.2  |
| n = 0 (%) | 27.0 | 46.8   | 58.3   | 25.6  | 33.6  | 46.2  |

Notes: Hyp-Ara, the number of arabinoses attached to Hyp, $n = 0$ means non-glycosylated Hyp. Ratio of different arabinino-oligosaccharides was expressed as percentage of Hyp bearing those oligosaccharides to total Hyp.
the extent of EXT cross-linking after deglycosylation. With a higher initial cross-linking rate, more monomers were converted to higher molecular weight oligomers (Figs. 2 and 3) after 30 minutes. The results are also summarized in Table 3.

RSH Hyp-oligoarabinosides structure determination.
In order to determine the structures of RSH Hyp-Araγ and Hyp-Araγ, we first analyzed the glycosyl linkages of the RSH glycans (Table 4). RSH showed a simple glycan profile that contained mainly terminally linked Gal residues (presumably attached to the serine residues) and twice as much 2-linked as 3-linked arabinofuranosyl residues (2-Araf and 3-Araf). There was also a trace of 2, 4-linked Ara (0.3%) and a small amount of ribose; undoubtably, a contamination as ribose is not a component of extracellular molecules. Further, an earlier neutral sugar composition of RSH indicated it contained mainly Ara and Gal (Table 4), but clearly no ribose.

Figure 1. Hyp-Ara profiling of partially deglycosylated EXTs. (A) Top to bottom: RSH, RSHAr, and RSHpH2; (B) top to bottom: TOMP1, P1Ara, and P1pH2. Each Hyp-arabinosides were present in two peaks due to two configurations of Hyp (cis or trans) generated during base hydrolysis. The Hyp-Araγ peaks disappeared after α-arabinofuranosidase treatment, while some Hyp-Araγ glycans were still present after pH 2.0 HCl deglycosylation. See Table 2 for Hyp-oligoarabinan content quantification in each sample.

Table 3. Initial cross-linking (XL) reaction rates of deglycosylated EXTs.

| EXT | INITIAL CROSS-LINKING RATE (µg/mL-min) | PEAK AREA LOSS (5–30 min XL, %) |
|-----|--------------------------------------|---------------------------------|
| Native RSH | 268.8 ± 9.4 | 80.3 |
| RSHAr | 85.0 ± 3.3 | 40.1 |
| RSHpH2 | 223.2 ± 12.4 | 74.2 |
| RSHpH1 | 0 | 0 |
| RSHHF | 0 | 0 |
| Native TOMP1 | 213.3 ± 6.3 | 73.9 |
| P1Ara | 28.5 ± 2.5 | 24.9 |
| P1pH2 | 95.3 ± 1.7 | 54.2 |
| P1pH1 | 0 | 0 |
| P1HF | 0 | 0 |

Notes: Three technical replicas averaged to obtain the rate of native extensins and extensins deglycosylated with arabinofuranosidases and pH 2.0 HCl. Area loss is presented as (monomer area five minutes–monomer area 30 minutes)/monomer area five minutes. Area losses were calculated once from profiles in Figure 2 (RSH) and Figure 3 (TOMP1).

Discussion
In EXTs, glycans comprise ~55% of the glycoprotein on a dry weight basis. In the cell, these glycans are attached to EXTs via a series of complex posttranslational modifications prior to the delivery of “mature” EXTs to the cell wall by Golgi vesicles, where they carry out their molecular function by self-assembling into positively charged scaffolds that guide
Figure 2. Superose 6 size exclusion chromatography profiles of the cross-linking of deglycosylated RSH. Left to right: native RSH (control), RSH treated with arabinofuranosidase (RSHAra) and pH 2.0 HCl (RSHpH2), respectively. A conversion of monomers to oligomers with the increase of cross-linking time is observed. The two peaks after 100 minutes were contributed by buffer and 2-mercaptoethanol.
Figure 3. Superose 6 size exclusion chromatography profiles of the cross-linking of deglycosylated TOMP1. Left to right: native TOMP1 (control), TOMP1 treated with arabinofuranosidase (P1Ara) and pH 2.0 HCl (P1pH2), respectively. A conversion of monomers to oligomers with the increase of cross-linking time is observed. The two peaks after 100 minutes were contributed by buffer and 2-mercaptoethanol.
the correct deposition of wall polysaccharides (mainly pectin) during growth and mitosis.\textsuperscript{54} Several glycosyltransferases are responsible for EXT glycosylation after hydroxylation of certain Pro residues to Hyp (eg, in Ser-(Hyp) motifs) by prolyl hydroxylase.\textsuperscript{55,56} One galactosyltransferase has been identified in Chlamydomonas reinhardtii (C. reinhardtii)\textsuperscript{57} that is responsible for adding the Gal to some EXT Ser residues, while several genes in Arabidopsis genome (RRA1—AT1G75120 and RRA2—AT1G75110;\textsuperscript{58} RRA3—AT1G19360;\textsuperscript{56} XEG113—AT2G35610)\textsuperscript{59} have been identified to potentially involve in EXT arabinosylation. XEG113 was proposed to add the third \(\beta\)-Ara onto the Hyp-Ara and Hyp-Ara chains,\textsuperscript{59} while RRA3 adds the second \(\beta\)-Ara to EXT oligoarabinosyl chains.\textsuperscript{56} Despite their high proportion in the protein, especially the major arabinoglycosidase structures that contribute ~95% (w/w) of the total EXT carbohydrate,\textsuperscript{60,61} the function of EXT glycans remains unclear. Earlier evidence has indicated that EXT arabinosylation may play a role in the protein–glycan network in the primary cell walls.\textsuperscript{29,44} Arabinosylation was also proposed to be important for conferring resistance to proteolysis\textsuperscript{44} and stabilizing the extended EXT polyproline-II (ppII) conformation.\textsuperscript{60–62} A loss of ppII conformation was observed in deglycosylated EXT monomers by circular dichroism spectroscopy\textsuperscript{64} and electron microscopy.\textsuperscript{61} The role of arabinosylation in EXT cross-linking was first suggested by the observation that completely dearabinosylated TOMP1 could not be cross-linked by the extensin peroxidase from tomato cell culture medium.\textsuperscript{33} Later, by gene knockout experiments, it was found that mutants-lacked EXT arabinosyltransferase genes showed a decreased EXT content in their cell walls.\textsuperscript{56,59} To test the effect of arabinosylation on EXT in vitro cross-linking, we partially dearabinosylated RSH and TOMP1 with \(\alpha\)-l-arabinofuranosidase (ideally removes only the fourth Ara on Hyp-Ara due to its unique \(\alpha\) configuration) and pH 2.0 HCl (less selective and removes Ara on all arabinoglycosidase chains). RSH and TOMP1 were also completely dearabinosylated by pH 1.0 HCl (preserved the Gal residues) and completely deglycosylated with anhydrous HF (removed all the sugar residues). EXT samples after each treatment were cross-linked in vitro using tomato extensin peroxidase, and their initial cross-linking rates were determined.

Treatment of RSH with \(\alpha\)-l-arabinofuranosidase removed all the fourth Ara on Hyp-Ara, as expected (Table 2). However, this treatment did not increase the percentage of the Hyp-Ara, but instead a twofold increase in non-glycosylated Hyp was observed according to the Hyp-Arabinoside profile (Table 2). This observation indicated that either the Hyp-Ara oligosaccharides in RSH had more \(\alpha\)-linked arabinofuranosides than only the fourth Ara in Hyp-Ara as that of tobacco\textsuperscript{60,51} or the enzyme used was contaminated with \(\beta\)-l-arabinofuranosidase activity. The fact that Arabidopsis and tobacco are only distantly related raises the possibility that the structure of the arabinosides may not be monolithic. Thus Hyp-Ara and Hyp-Ara were isolated from RSH and their structures determined by NMR. The structures of RSH Hyp-Ara and Hyp-Ara are identical to those from tobacco (Figs. 4–6, Table 5). Thus, it is likely that the old \(\alpha\)-l-arabinofuranosidase preparation might contain \(\beta\)-l-arabinofuranosidase activity (also confirmed by a communication from Megazyme Company).

Besides the structural conservation of Hyp-Ara and Hyp-Ara between Arabidopsis and tobacco, a recent study of the Hyp-Ara and Hyp-Ara on EXTs of Boswellia serrata and B. carteri also showed similar structures.\textsuperscript{64} These findings indicate a conserved glycosylation pattern among these dicot species. Interestingly, Waissfich et al’s group isolated Hyp-oligosaccharides from green algae C. reinhardtii and determined the glycosidic linkages and configurations in those glycans. Their proposed C. reinhardtii glycan structure were \(\alpha\)-D-Galf-(1→2)-\(\beta\)-l-Araf-(1→2)-\(\beta\)-l-Araf-(1→4)-Hyp and 3-methyl-\(\beta\)-l-Araf-(1→5)-\(\alpha\)-D-Galf-(1→2)-\(\beta\)-l-Araf-(1→2)-\(\beta\)-l-Araf-(1→4)-Hyp.\textsuperscript{64} The algae Hyp-oligosaccharides showed similarity in the inner-core glycan structure (\(\beta\)-l-Araf-(1→2)-\(\beta\)-l-Araf-(1→4)-Hyp) with those from dicots,\textsuperscript{51,64} which are undoubtedly due to some common steps in their biosynthetic evolution.\textsuperscript{64} Such conserved arabinoglycosidase structures across the evolutionary progression imply an important functional role for EXT oligoarabinosides. Data from this study support the notion that this structural conservation could play a role in EXT cross-linking.

First of all, the completely dearabinosylated RSH and TOMP1 (pH 1.0 HCl and HF treatment) could not

| MONOSUGAR | COMPOSITION (mole%) | GLYCOSYL LINKAGE | % PEAK AREA |
|-----------|---------------------|-----------------|-------------|
| Arabinose (Ara) | 79 | Terminal linked arabinofuranosyl residue (t-Araf) | 8 |
| | | 3-linked arabinofuranosyl residue (3-Araf) | 21 |
| | | 2-linked arabinofuranosyl residue (2-Araf) | 54 |
| Galactose (Gal) | 15 | Terminal linked galactopyranosyl residue (t-Gal) | 15 |
| Rhamnose (Rha) | 2 | | |
| Xylose (Xyl) | 2 | | |
| Glucose (Glc) | 2 | 2-linked ribofuranosyl residue (2-Ribf) | 2 |

Notes: Monosugar analysis results were obtained from Terneus.\textsuperscript{20} Trace amount (0.3%) of 2,4-linked Ara was also observed.
Figure 4. Structure elucidation of RSH Hyp-Ara₄. (A) HSQC spectrum: cross-peaks identified the chemical shifts of each carbon atom and its corresponding hydrogen atom(s) in each arabinose ring system. A₄ is the fourth Ara at the non-reducing end of the Hyp-Ara₄ chain, while A₁ occupies the reducing end and is attached to Hyp (first Ara in the chain). A₂ and A₃ are the second and third Ara of the chain. The A₁ C₁/H₁ label indicates the cross-peak arising from the chemical shifts of the anomeric carbon (C₁) and its corresponding hydrogen (H₁) on the A₁ residue. The cross-peaks for the other carbon atoms and their corresponding hydrogens of A₁ and the cross-peaks for A₂ to A₄ are similarly labeled. Two cross-peaks are observed for the fifth carbon atoms on each ring system due to their possession of two corresponding hydrogen atoms. (B) HMBC spectrum: the cross-peaks arising from A₄H₁(5.2 ppm) + A₃C₃(84.8 ppm) and A₄C₁(112.1 ppm) + A₃H₃(4.2 ppm), highlighted by red circles, established the α-Ara-(1→3)-β-Araf linkage between A₄ and A₃; cross-peaks arising from A₃H₁(5.1 ppm) + A₂C₂(84.5 ppm) and A₃C₁(103.8 ppm) + A₂H₂(4.3 ppm), highlighted by blue circles, established the β-Araf-(1→2)-β-Araf linkage between A₃ and A₂; cross-peaks arising from A₂H₁(5.2 ppm) + A₁C₂(84.5 ppm) and A₂C₁(102.7 ppm) + A₁H₂(4.3 ppm), highlighted by orange circles, established the β-Araf-(1→2)-β-Araf linkage between A₂ and A₁, and cross-peak arising from A₁H₁(5.3 ppm) + HypC₄ (81.1 ppm), highlighted by the green circle, established the β-Araf-(1→4)-Hyp linkage between A₁ and Hyp. The chemical shifts of the cross-peaks are summarized in Table 5.
Figure 5. Structure elucidation of RSH Hyp-Ara₃. (A) HSQC spectrum: cross-peaks identified the chemical shifts of each carbon atom and its corresponding hydrogen atom(s) in each arabinose ring system. A₃ is the third Ara at the nonreducing end of the Hyp-Ara₃ chain, while A₁ occupies the reducing end and is attached to Hyp (first Ara in the chain). A₂ is the second Ara of the chain. The A₁ C₁/H₁ label indicates the cross-peak arising from the chemical shifts of the anomeric carbon (C₁) and its corresponding hydrogen (H₁) on the A₁ residue. The cross-peaks for the other carbon atoms and their corresponding hydrogens of A₁ and the cross-peaks for A₂ to A₃ are similarly labeled. Two cross-peaks are observed for the fifth carbon atoms on each ring system due to their possession of two corresponding hydrogen atoms. There are overlapping signals between the cross-peaks from A₁ C₂/H₂ and A₂ C₂/H₂ and those from A₂ C₅/H₅ and A₃ C₅/H₅ due to their identical chemical shifts (Table 5). (B) HMBC spectrum: cross-peaks arising from A₃H₁(5.1 ppm) + A₂C₂(84.5 ppm) and A₃C₁(103.9 ppm) + A₂H₂(4.3 ppm), highlighted by blue circles, established the β-Ara-(1→2)-β-Ara linkage between A₃ and A₂; cross-peaks arising from A₂H₁(5.2 ppm) + A₁C₂(84.5 ppm) and A₂C₁(102.7 ppm) + A₁H₂(4.3 ppm), highlighted by orange circles, established the β-Ara-(1→2)-β-Ara linkage between A₂ and A₁, and cross-peak arising from A₁H₁(5.3 ppm) + HypC₄(81.0 ppm), highlighted by the green circle, established the β-Ara-(1→4)-Hyp linkage between A₁ and Hyp. The chemical shifts are summarized in Table 5.
Arabinosylation affects extensin cross-linking in vitro

Table 5. Chemical shifts (ppm) of RSH arabinose C/H atoms in Hyp-Ara4 and Hyp-Ara3.

| RESIDUES    | C1/H1 | C2/H2 | C3/H3 | C4/H4 | C5/H5 |
|-------------|-------|-------|-------|-------|-------|
| Hyp-Ara4    |       |       |       |       |       |
| α-Araf (A4) | 112.1 | 85.3  | 80.7  | 88.3  | 65.4  |
| β-Araf (A3) | 103.9 | 80.2  | 84.6  | 85.6  | 66.6  |
| β-Araf (A2) | 102.7 | 84.5  | 76.3  | 85.9  | 66.5  |
| β-Araf (A1) | 102.9 | 84.5  | 76.5  | 85.6  | 66.9  |
| Hyp         | 175.4 | 61.1  | 38.7  | 81.1  | 54.2  |
| Hyp-Ara3    |       |       |       |       |       |
| β-Araf (A3) | 103.9 | 77.9  | 80.5  | 86.1  | 66.4  |
| β-Araf (A2) | 102.7 | 84.5  | 76.3  | 85.8  | 66.4  |
| β-Araf (A1) | 102.8 | 84.5  | 76.5  | 85.5  | 66.9  |

Notes: A4 in Hyp-Ara4 and A3 in Hyp-Ara3 are the terminal arabinose residues at the non-reducing end. A1 in both Hyp-Ara4 and Hyp-Ara3 is the reducing end arabinose that attached to Hyp. Values corresponding to the cross-peaks in Figures 4 and 5. Anomeric C chemical shift obtained from Akiyama et al51 identified A4 to be α-Ara while the remaining β-Ara in Hyp-Ara4 and all the Ara in Hyp-Ara3 as β-Ara. Two H shift values for the fifth carbons due to their possession of two corresponding hydrogen atoms. The difference in chemical shifts between the fourth C of Hyp in Hyp-oligoarabinan and that of free Hyp (81 ppm vs 71.3 ppm, Akiyama et al51) identified a linkage on the fourth C of Hyp in Hyp-oligoarabinan.

be cross-linked by the tomato extensin peroxidase (initial rate = 0, Table 3). This result agreed with that from a previous study33 and indicates that arabinosylation is indeed required for EXT cross-linking.

Second, EXTs partially dearabinosylated by α-arabinofuranosidase (RSHAra and P1Ara) and pH 2.0 HCl (RSHpH2, and P1pH2) all have lower Ara content (Table 1) and slower initial cross-linking rates than the native EXTs (Table 3). This observation indicates a positive correlation between the cross-linking rate and the Ara content.

Finally, RSHpH2 and P1pH2 showed a much faster initial cross-linking rate (Table 3), and at the end of a 30-minute cross-linking, a much greater loss in the monomeric EXT content (Figs. 2 and 3, Table 3) compared with RSHAra and P1Ara was observed. The cross-linking profiles also revealed that monomeric RSHpH2 were converted to a larger

Figure 6. Characterized primary structure of RSH Hyp-Ara4 and Hyp-Ara3. Each arabinose residue is labeled corresponding to those in Table 5 with their anomeric configurations. The glycosidic linkages between arabinose residues and with Hyp are labeled according to the HMBC data (Figs. 4 and 5) with carbon numbers indicating the C atoms involved in the linkages.
of RSHpH2. In cross-linking sites may account for the better cross-linking of EXTs in the tomato genomic database—Sol Genomic Network potentially form pulcherosine (deduced from TOMP1 protein sequence). Notably, TOM1 has only three potential Idt motifs at the protein terminus that can cross-link with its abundant Val-Tyr-Lys motifs to potentially form pulcherosine (deduced from TOM1 protein sequence in the tomato genomic database—Sol Genomic Network, accession SGN-U315189). Thus, greater abundance in cross-linking sites may account for the better cross-linking of RSHpH2.

**Conclusion**

The cross-linking of partially or fully deglycosylated EXTs led to the first conclusion that EXT arabinosylation, especially the α-configured fourth Ara on the Hyp-Araα chains, is crucial for the initial rate and extent of EXT cross-linking in vitro by peroxidase. Determination of glycan structures of RSH Hyp-Araα and Hyp-Araβ showed structural conservancy of those glycans between different dicot species and even with green algae, indicating the importance of the Hyp-arabinogalactosaccharides throughout the evolutionary progression of different plant species.

**Author Contributions**

Designed the research: YC, MJK. Performed the experiments: YC, WD, LT. Analyzed the data: YC, MAH, MJK. Wrote the paper: YC, LT, MAH, MJK. All authors read and approved the final manuscript.

**Supplementary Materials**

**Supplementary figure 1.** EXTs isolation by cation exchange and reverse-phase chromatography. Dialyzed tomato culture eluate was fractionated by SP Sepharose (A) chromatography and Peak 1 (collected between markers) was purified by reverse-phase (B) that yielded TOMP1. Similarly, dialyzed Arabidopsis culture eluate was fractionated by SP Sepharose (C) chromatography and Peak 1 was purified by reverse-phase (D) that yielded RSH.

**Supplementary figure 2.** Cross-linking of deglycosylated extensins. Deglycosylated TOM1 (A) and RSH (B) cross-linked for 30 minutes by peroxidase. No change in monomer areas were observed in all cases. pH 1.0 HCl deglycosylated sample runs were recorded for a shorter time (120 minutes); thus, the succinic acid peaks were not included in those profiles. Monomer peak shoulders could be due to the uneven succinylation of the monomers.

**Supplementary table 1.** Amino acid composition (mol%) of extensin precursors TOMP1 and RSH.

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