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Multi-omics analysis identifies genes mediating the extension of cell walls in the Arabidopsis thaliana root elongation zone

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INTRODUCTION

The plant kingdom displays an enormous diversity in shapes and sizes, varying from unicellular algae with a simple rather spherical morphology to very complex multicellular organisms that can reach more than 100 m in height. Growth of plants is the sum of two processes, namely the increase in cell number by repeated cycles of cell division and the subsequent—sometimes major—increase in volume of these newly formed cells by expansion. Both processes are controlled by the action of plant hormones among which auxin plays a major role (Perrot-Rechenmann, 2010). In roots, cells pass sequentially through different developmental stages along the root axis. Growth occurs through rapid elongation of cells in a zone shootward to the root apical meristem, which is the site of cell division, and before further cell type differentiation, for example root hair emergence and lateral organ initiation (Verbelen et al., 2006).

Plant cell walls are rigid yet deformable materials, and growth is seen as the irreversible increase in surface area of cell walls. This process requires an internal turgor pressure, which arises from water uptake into the cell. Turgor exerts a force against surrounding cell walls, but is otherwise mediated by changes in the mechanical properties of the walls, resulting in stress relaxation (Ray et al., 1972; Cosgrove, 1986, 1993, 2005; Guerriero et al., 2014). Cell walls are made up of a fibrillar component, the cellulose microfibrils, that is embedded in a highly hydrated matrix of pectins, which principally comprise

Keywords: root growth, plant cell walls, multiomics, transcriptomics, localisomics, epitomics, cell-wall polysaccharides, cell elongation

Plant cell wall composition is important for regulating growth rates, especially in roots. However, neither analyses of cell wall composition nor transcriptomes on their own can comprehensively reveal which genes and processes are mediating growth and cell elongation rates. This study reveals the benefits of carrying out multiple analyses in combination. Sections of roots from five anatomically and functionally defined zones in Arabidopsis thaliana were prepared and divided into three biological replicates. We used glycan microarrays and antibodies to identify the major classes of glycans and glycoproteins present in the cell walls of these sections, and identified the expected decrease in pectin and increase in xylan from the meristematic zone (MS), through the rapid and late elongation zones (REZ, LEZ) to the maturation zone and the rest of the root, including the emerging lateral roots. Other compositional changes included extensin and xyloglucan levels peaking in the REZ and increasing levels of arabinogalactan-proteins (AGP) epitopes from the MS to the LEZ, which remained high through the subsequent mature zones. Immuno-staining using the same antibodies identified the tissue and (sub)cellular localization of many epitopes. Extensins were localized in epidermal and cortex cell walls, while AGP glycans were specific to different tissues from root-hair cells to the stele. The transcriptome analysis found several gene families peaking in the REZ. These included a large family of peroxidases (which produce the reactive oxygen species (ROS) needed for cell expansion), and three xyloglucan endo-transglycosylase/hydrolase genes (XTH17, XTH18, and XTH19). The significance of the latter may be related to a role in breaking and re-joining xyloglucan cross-bridges between cellulose microfibrils, a process which is required for wall expansion. Knockdowns of these XTHs resulted in shorter root lengths, confirming a role of the corresponding proteins in root extension growth.
homogalacturonan (HGA), rhamnogalacturonan-I (RG-I), and rhamnogalacturonan-II (RG-II). The tethering of the adjacent cellulose microfibrils occurs primarily through xyloglucan in dicotyledonous and non-commelindian monocotyledonous walls (Hayashi, 1989), or by glucuronoxarabinoxylan (Nishitani and Nevin, 1991; Carpita and Gibeaut, 1993) and mixed-linkage (1-3),(1-4)-β-D-glucans in the walls of Poales and Equisetales (Kato et al., 1982; Scheller and Ulvskov, 2010; Mohler et al., 2013).

As well as pectin and hemicelluloses, several different classes of glycoproteins and enzymes are also present (Albenne et al., 2009). This complex composition results in the mechanical properties of the cell wall and greatly influences its growth potential. The volume increase of a plant cell was described in the Lockhart equation (1965), \( dV/dT = \phi \cdot (P-Y) \), where \((P-Y)\) is the turgor above a yield threshold \(Y\) that must be exceeded before plastic wall extension can occur, and \(\phi\) is the extensibility coefficient that represents the time-dependent yielding properties of the cell wall in the direction of growth (Schopfer, 2006). Several proteins that can influence the cell wall's yielding parameters have been described, including expansins (McQueen-Mason et al., 1992), xyloglucan endotransglycosylase/hydrolases (XET/XTHs; Nishitani and Vissenberg, 2007; Miedes et al., 2013), peroxidases (Passardi et al., 2006), \(\beta\)-(1-4)-glucanases (Labrador and Nevin, 1989), yield-ins (Okamoto-Nakazato et al., 2001), and lipid transfer proteins (LTPs; Niewland et al., 2005).

With the emergence of different molecular biological approaches and tools, many genes that encode enzymes with a role in the synthesis of the various cell wall polysaccharides and proteins found in cell walls have been identified and their mutants described (e.g., Harcholt et al., 2010; Carpita, 2011; Mewalal et al., 2014). The synthesis of complete cell-wall components, their trafficking and final assembly in cell walls are, however, very complex (McCann and Rose, 2010) and there is often not a simple link between genotype and a growth phenotype. Furthermore, the content, architecture and biophysical characteristics of the walls of a cell change at any point along the growth axis as a consequence of both the cell's history (i.e., the multiple processes since the cell originated) and the needs of its current location.

As a result of this complexity, point measurements are extremely difficult to interpret and the use of a single “omics” technique to uncover cell-wall processes underpinning plant growth might not be sufficient (Somerville et al., 2004; Farrokhi et al., 2006). Therefore, model organisms, such as Arabidopsis thaliana, and appropriate research tools are needed. The A. thaliana root has a relatively simple anatomy and develops in a highly predictable manner (Dolan et al., 1993), lending itself to investigation of growth mechanisms, and their regulation as evidenced by numerous reports (e.g., Ubeda-Thomás et al., 2009; Band et al., 2011; Bruex et al., 2012; De Rybel et al., 2012). In addition, its genome sequence is published (Arabidopsis Genome Initiative, 2000) and many research tools already exist (e.g., Fukao et al., 2013; Jacques et al., 2013; Moussaieff et al., 2013).

We used a combination of point measurements and three techniques to characterize the different developmental zones along the A. thaliana root, looking at cell wall composition by means of quantitative assessment of cell-wall epitopes (epitomics), epitope localization (localisomics), and gene expression (transcriptomics), and combined the -omics data in this study to provide an integrated perspective. This revealed that individual -omics-techniques are inadequate and can even result in misleading conclusions. In contrast, the multi-omics approach has identified three gene families that appear to play a role in regulating root growth, and mutant analysis for one of these families (XTHs) supports these findings.

**MATERIALS AND METHODS**

**PLANT MATERIAL AND HANDLING**

Seeds of Arabidopsis thaliana (L.) Heynh. (ecotype Columbia-0) were surface-sterilized by incubation in 5% (v/v) sodium hypochlorite for 5 min, washed three times in sterile water and sown on vertical 125 × 125 mm square Petri plates. Each plate contained 60 ml 1/2 strength Murashige and Skoog media (Sigma) solidified with 1% (w/v) agar. For material used for transcriptomic and glycan microarray profiling (epitomics), sterile 9 × 9 cm square sections of 100 μm nylon mesh (ClaroCor) were placed onto the media surface before sowing to facilitate root dissection and harvesting of cut sections. After 2 days at 4°C, plates were transferred to controlled-environment chambers at 23°C under continuous light at a photon flux density of 150 μmol m\(^{-2}\) s\(^{-1}\) for 7 days.

Roots were dissected into five sections as shown in Figure 1: (1) meristem (from the root tip to the top of the lateral root cap, approximately 350 μm from the tip); (2) rapid elongation zone (from the top of the lateral root cap to the first visible root hair bulge, approximately 850 μm from the shootward boundary of zone 1); (3) late elongation (deceleration) zone (from the first root hair bulge to the first fully elongated root hair); (4) mature root (500 μm shootward of the first fully elongated root hair); and the lateral root zone (2.5 cm in length, from the shootward boundary of zone 4 in a shootward direction). Dissected samples were immediately frozen in liquid nitrogen.

**EPITOMICS**

Techniques involving glycan microarrays were used for this (Shin et al., 2005; Moller et al., 2007, 2008). Cell wall material was isolated from dissected material as alcohol-insoluble residue (AIR). Frozen material was ground in liquid nitrogen using a micro-pestle, 1 ml of 70% (v/v) ethanol was added to each tube and the mixture shaken at 4°C for 1 h. The mixture was centrifuged (5 min, 10,000 ×g) and the supernatant discarded. This process was repeated 4 times. Pellets were resuspended in 1 ml of acetone for 5 min and then air-dried overnight. A total of 600 roots were dissected, yielding 10–20 mg AIR for each of the
five sections, which were subjected to three sequential extractions as previously described (Sørensen and Willats, 2011). Briefly, the extractions were performed using 15 μl each of 50 mM diamino-cyclo-hexane-tetra-acetic acid (CDTA), 4 M sodium hydroxide (NaOH) with 0.1% v/v sodium borohydride (NaBH4) and cadoxen [31% (v/v) 1,2-diaminoethane with 0.78 M cadmium oxide (CdO)]. These respectively enrich for pectin, hemicelluloses, and cellulose-associated molecules. The extracted fractions were printed as microarrays with six replicates and three dilutions using a Microgrid II microarray robot (Genomic Solutions, Ann Arbor, MI, USA) and the arrays were probed with a range of primary antibodies or carbohydrate-binding modules and appropriate alkaline phosphatase (AP) conjugated secondary antibodies before developing as previously described (Sørensen and Willats, 2011). All JIM- and LM-monoclonal antibodies and 2F4 were obtained from PlantProbes (http://www.plantprobes.net), CCRC antibodies from CarboSource (http://www.ccrc.uga.edu/~carbosource/CSS_home.html) and secondary antibodies from Sigma-Aldrich (http://www.sigmaaldrich.com). The arrays were scanned and analyzed using the microarray software ImaGene 6.0 (http://www.biodiscovery.com) to obtain raw signal values. These were then treated in the same way as fluorescent microarray data. Specifically, the value from the negative control where the secondary antibody was omitted was subtracted, and then the median and median-absolute deviation values were calculated. A heatmap was produced using Microsoft Excel.

LOCALISOMICS
Four-day-old seedlings were fixed and prepared for whole-mount immunolocalization analyses requiring some cell-wall permeabilization steps as described previously (Peret et al., 2012). Cell wall antibodies were used at 1:100 dilutions, whereas Alexafluor488 or Alexafluor543 coupled anti-rat or anti-mouse secondary antibodies were used at 1:200 dilutions to give green or red fluorescence, respectively. Counter staining was performed using either propidium iodide (for the AlexaFluor488 coupled secondary antibody) or Sytox Green (for AlexaFluor543 coupled secondary antibody). Seedlings were mounted in 50% glycerol and images were taken using a Leica SP2 confocal laser scanning microscope (Leica Microsystems UK Ltd). The specific antibodies, their epitopes and localization (primarily as determined in this study) are listed in Table 1.

TRANSCRIPTOMICS
Three biological replicates from separate pools of seeds were used. For each biological replicate, plants were grown and approximately 50 roots dissected as described in Section Plant Material.

Table 1: Cell wall epitopes assessed in the glycan microarray and their localization from in situ fluorescence studies.

| Antibody | Epitope | Localization | References/image |
|----------|---------|--------------|------------------|
| CAL 1→3-β-glucan | n/a | All walls | Figure 3 and Freshour et al., 2003 |
| CCRC-M1 Xyloglucan | n/a | Secondary cell walls | McCartney et al., 2006 |
| CBM22 Xylan | n/a | Mainly lateral root cap and young epidermis; in meristem some cell-wall plates in epidermis and a stele cell file | Supplementary Figure S1, pp. 12–17 |
| 2F4 Calcium-stabilized homogalacturonan chains | n/a | Mainly lateral root cap | Supplementary Figure S1, p. 18 and McCartney et al., 2003 |
| LM1 Extensin (HRGP) | n/a | Mainly lateral root cap | Supplementary Figure S1, p. 21 and Willats et al., 2008 |
| LM2 A6 (β-linked glucuronic acid) | n/a | Mainly lateral root cap | Supplementary Figure S1, p. 21 and Willats et al., 2008 |
| LM5 1→4-β-D-galactan | n/a | Quiescent center and mature epidermis, especially at interface between cell files and root hairs | Supplementary Figure S1, p. 21 and Willats et al., 2008 |
| LM6 1→5-α-L-arabinan/AGPs | n/a | Stele files especially zones 2–5, faint signal in epidermis | Supplementary Figure S1, p. 21 and Willats et al., 2008 |
| LM8 Xyloglacturonan | n/a | | Supplementary Figure S1, p. 21 and Willats et al., 2008 |
| LM10 1→4-β-D-xylan | n/a | | Supplementary Figure S1, p. 21 and Willats et al., 2008 |
| LM15 XXXG motif of xyloglucans | n/a | | Supplementary Figure S1, p. 21 and Willats et al., 2008 |
| LM5 Partially methylstified-homogalacturonan | n/a | | Supplementary Figure S1, p. 21 and Willats et al., 2008 |
| JIM7 Partially methylstified-homogalacturonan | n/a | | Supplementary Figure S1, p. 21 and Willats et al., 2008 |
| JIM8 AGP glycan | n/a | | Supplementary Figure S1, p. 21 and Willats et al., 2008 |
| JIM13 AGP glycan | n/a | | Supplementary Figure S1, p. 21 and Willats et al., 2008 |
| JIM19 Extensin | n/a | | Supplementary Figure S1, p. 21 and Willats et al., 2008 |
| JIM20 Extensin | n/a | | Supplementary Figure S1, p. 21 and Willats et al., 2008 |
| MAC207 AGP glycan | n/a | | Supplementary Figure S1, p. 21 and Willats et al., 2008 |
and Handling. RNA was extracted using the Qiagen MicroRNA Kit following the manufacturer’s instructions (Qiagen, Crawley, UK) and quantified using a Nanodrop ND100 spectrophotometer (Nanodrop, Wilmington, USA). All RNA samples were approximately 50 ng μL$^{-1}$ in a total volume of 10 μL. Labeling of RNA samples was conducted using the Affymetrix IVT-Express Eukaryotic Target Labeling Assay kits following standard Affymetrix protocols (Affymetrix UK Ltd., High Wycombe, UK). RNA labeling and hybridization to Affymetrix ATH1 arrays were performed by the Nottingham Arabidopsis Stock Centre (NASC).

Data were normalized from .cel files with RMA statistics performed using the Limma package in R/Bioconductor (Smyth, 2005) and a custom CDF file (ATH1121501_AT_TAIRT v17; Dai et al., 2005). A gene was considered to be expressed if a t-test between two zones was significant at a q-value of 0.05 after Benjamini and Hochberg false discovery rate correction (Smyth, 2005). Further analyses were performed using Excel 2010 (Microsoft Corporation, Redmond, USA). Genes were further annotated into cell-wall functional subclasses using Cell Wall Navigator (Girke et al., 2004) and PlnTFDB (Pérez-Rodríguez et al., 2010). Transcriptomics data used in these experiments have been made available through ArrayExpress (www.ebi.ac.uk) with accession number E-MEXP-2912.

**xth MUTANT ANALYSIS**

Atxth17-1(SALK_015077), Atxth19-1 (SALK_034274), Atxth20-1(SAIL_575_H09), and XTH18-RNAi were kindly provided by Prof. K. Nishitani (Tohoku University, Japan). Atxth17-2 (SALK_008429), Atxth19-2 (SAIL_62_A10), and Atxth20-2 (SALK_066689) were obtained from NASC. All lines are in the Columbia-0 background. To assess basal root growth, the root length of seedlings grown vertically for 7 days was measured from the hypocotyl to the root tip. Root lengths were measured using the NeuronJ plugin of ImageJ 1.41j (http://rsb.info.nih.gov/ij/). Two-tail Student t-tests were performed using Excel 2010 to determine significance (p-value < 0.05).

Confocal microscopy for imaging of *A. thaliana* roots was performed using a Leica SP5 confocal laser-scanning microscope (Leica, Milton Keynes, UK). For cell quantification and cell length measurements, seedlings were treated with propidium iodide (10 μg ml$^{-1}$; Sigma) to visualize cell walls. Cell lengths were measured using the Cell-o-Tape image-analysis tool (French et al., 2012). Data are presented as the mean ± the standard error and two-tail Student t-tests, used to determine significance (p-value < 0.05), were performed using Microsoft Excel software.

**RESULTS**

**EPITOMICS**

Figure 2 depicts the epitope intensities in the different fractions and root zones, and highlights the zonal difference between antibodies to related epitopes. The signals corresponding to the pectin I and xyloglucan (XyG) binding antibodies, suggests that the CDTA and NaOH treatments were effective in terms of extracting the associated polysaccharides (pectin and hemicelluloses, respectively), since it was expected that some cellulose-associated XyG would be extracted using cadoxen due to the fact that XyG tethers adjacent cellulose microfibrils.

The more soluble (1→3)-β-glucans, recognized by CAL, peaked in zones 2 and 4 in the hemicellulosic fraction while in the pectin fraction they actually peaked in zone 5, consistent with the highest pectin-associated signal of crystalline cellulose. The xylans (recognized by LM10 and CBM22) were present in all three fractions. The CBM22 signal peaked in zones 1–3 in the pectin fraction, in zones 2–4 in the hemicellulosic fraction and in zone 5 in the cellulose fraction. LM10, however, showed an increasing signal in the hemicellulosic fraction as the root matured, a trend that was also present in the cellulose fraction where it was accompanied by a reduction in zones 2 and 3.

Xyloglucans (probed by CCRC-M1 and LM15) peaked in the REZ and were specific to the hemicellulosic fraction, with only minor binding in the cellulose fraction, a pattern becoming more evident as the root matures. The homogalacturonan/pectin I epitope, probed by 2F4, JIM5, and JIM7, which differ in their sensitivity to the degree of esterification, shows a similar pattern and is specific to the pectin fraction. JIM7 peaked in zones 1 and 2; JIM5 peaked in zone 2, and 2F4 peaked in zones 2 and 3.

The pectin II epitopes show a broad range of patterns, consistent with the complexity of its polymer constituents. LM6, specific for arabinan, peaked in the hemicellulosic fraction in zone 1 and rapidly decreased as the root matures. LM8, specific for xylogalacturonan, peaked in zones 1 and 2 in the same fraction and to a lesser extent in the cellulose fraction before trailing off rapidly after zone 3, whilst in the pectin fraction it was consistently present at a low level throughout. LM5, recognizing galactan, shows one of the most complex signal patterns, peaking in zones 2–3 in the pectin fraction, zone 2 in the hemicellulosic fraction while showing the inverse pattern in the cellulose fraction.

Three antibodies detect variants from the extensin family. The LM1 and JIM20 epitopes are present at high levels in all zones, with the JIM20 signal being higher in the hemicellulosic fraction, peaking in zone 2, and the LM1 signal being nominally highest in zone 5. The JIM19 epitope, however, was absent from the meristematic region, was mostly pectin-associated in the elongation zones, and showed its highest signal in zones 4 and 5, but was specifically non-pectin associated. Several antibodies recognize different epitopes associated with AGPs. Their signal levels tended to be higher from zone 2 onwards, but they were also found in all extractions and developmental zones.

**LOCALISOMICS**

Figure 3 contains localization results for four classes of cell wall-related epitopes, namely XyGs, pectins, AGPs, and extensins. Crucially, the images show that antibodies were able to bind to epitopes throughout the root, rather than only at the surface, giving us confidence that cell-wall epitope localization can be obtained when using permeabilization procedures that will lead to the loss of some cell wall structures. However, five antibodies were not available for localization studies and a further five gave no signal, possibly due to the epitope being modified beyond recognition by the permeabilization process. The full image dataset can be found in the Supplementary Material.
The results for each antibody are scaled relative to the maximum signal for that antibody. The heatmap is supplemented by Supplementary Figure S2, which shows the actual signals in graph form for each antibody. Pectin I corresponds to homogalacturonan epitopes and Pectin II to rhamnogalacturonans.

**FIGURE 3 | Immunolocalization with key antibodies.** CCRC-M1 and LM15 detect xyloglucans, JIM5 detects pectins, JIM20 detects extensins, and JIM13 and LM6 bind to AGPs. An AF488-linked secondary antibody was used for all antibodies except CCRC-M1 for which an AF543-linked antibody was used. The scale bars correspond to 100 μm for all images except LM15 for which it is 25 μm.

**Table 2 | Transcriptomic data for the five zones of the A. thaliana root.**

| Zone  | Genes expressed (% of 21,331) | Significantly up-regulated | Significantly down-regulated |
|-------|-----------------------------|---------------------------|-----------------------------|
| Zone 1 | 7741 (36.3%)                | 1632                      | 1939                        |
| Zone 1–2 | 1632                      | 1939                      | 1343                        |
| Zone 2 | 7539 (35.3%)                | 1343                      | 1750                        |
| Zone 3 | 7801 (36.6%)                | 155                       | 369                         |
| Zone 4 | 7854 (36.8%)                | 155                       | 184                         |
| Zone 5 | 8044 (37.7%)                | 1322                      | 2627                        |

A gene was considered to be expressed if its expression was greater than 100 and differentially expressed if a t-test between two zones was significant at a q-value of 0.05.

**TRANSCRIPTOMICS**

A large percentage of the genome is expressed in the root with about 7500–8000 genes detectably expressed in any given root zone (see Table 2 and Supplementary Table 1). Zones 1–3, despite their physical proximity, show large inter-zonal differences consistent with their very different developmental roles. Zones 3–5 have more similar gene expression levels, highlighting the general developmental quiescence of the mature root. However, ~10% of significantly differentially expressed genes in these zones are annotated as cell-wall genes, which is twice the number of genes changing between the earlier zones. It is the changes between zones 1–2 and 2–3 that are likely to inform the expression changes that allow rapid elongation and deceleration, respectively. At their peak, 52% of reactive oxygen species (ROS) and 63% of aquaporin genes are expressed in the root. Only 25% of annotated transcription factors are detectably expressed in the root, which suggests that they may have specific roles in the cells where they are expressed but that their general role in root growth may be difficult to interpret. Finally, binding of the LM6 antibody, recognizing xylogalacturonan, was restricted to the lateral root cap in a manner reminiscent of LM6 (see Supplementary Material).

(Supplementary Figure S1) and epitope localization arising from this dataset is described in **Table 1**.

The XyGs recognized by LM15 were present at low levels throughout, but with a particularly high signal in the quiescent center, although no such specificity was seen with the CCRC-M1 antibody, recognizing fucosylated XyG. JIM5 showed the partially methylesterified pectins to be localized to the lateral root cap and radial walls in the inner tissues, while the extensins detected by JIM20 were most prominent in the epidermis and cortex of the REZ and to a lesser degree in the lateral root cap. JIM13 and LM6 binding patterns suggested that individual AGPs might be specific to different locations, suggesting that they play specific roles in the cells where they are expressed but that their general role in root growth may be difficult to interpret. Finally, binding of the LM8 antibody, recognizing xylogalacturonan, was restricted to the lateral root cap in a manner reminiscent of LM6 (see Supplementary Material).
is lower than other annotated gene families—possibly reflecting tight developmental differentiation.

**DISCUSSION**

The *A. thaliana* root develops in a highly predictable manner. Cells pass through consecutive developmental phases during which the post-mitotic elongation of cells contributes the majority of the increase in the root length. Cell wall metabolism is very important in allowing and controlling cellular expansion. The synthesis, trafficking, deposition, integration, and remodeling pathways of cell-wall components are, however, very complex and not completely resolved, so multiple omics-approaches are needed to establish which genes are contributing to the observed changes in composition and consequent mechanical properties.

**FIGURE 4** | Combined xyloglucan, XyG biosynthesis genes, and XTH expression profiles. (A) epitomic (columns) and XyG biosynthesis mRNA expression (lines). (B) zonal transcriptomic expression profiles for selected members of the XTH family. Error bars are ±1 SD.
These issues are complicated further by the observation that the different tissues differentially contribute to the mechanics of root elongation (Dyson et al., 2014).

**EPITOMICS ANALYSIS**

Epitomics (also known as glycan microarray analysis and previously described as Comprehensive Microarray Polymer Profiling) is a high-throughput microarray-derived technique that allows the handling of multiple samples. In this study, it was used to indicate which polysaccharides and glycoproteins were found in specific root developmental-zones. Many of the results support current knowledge of root cell-wall composition, for example that pectin is synthesized and deposited into the existing cell wall in a highly esterified form (Liners and Van Cutsem, 1992) and that pectinesterases modify the pectins while the cells age (Micheli, 2001). Carbohydrate polymer synthesis is complicated and does not necessarily correlate well with wall composition. Little is known of extant synthesis pathways, let alone the reality that pectin is synthesized and deposited into the existing cell wall. This suggests a role in maturation rather than cell size.

**LOCALISOMICS ANALYSIS**

In contrast to epitomics, whole-mount immunolocalization provides high spatial resolution and revealed cell and tissue-specific locations for some cell wall epitopes that would not have been clear from the epitomic data alone. The AGPs appear to be the most remarkable in this respect, suggesting that individual proteins play a specific and subtly different role. The localization of extensins to zone 2 epidermis and cortex is intriguing, given the predictions of Dyson et al. (2014) that the outer layers of the root have most influence on growth. Unfortunately, some of the antibodies were not available for this technique and others did not yield any signal in the cell walls, probably because of masking of the epitope by other cell wall components or more likely due to the use of enzymes in the permeabilization procedures required for whole-mount preparations. This -omics has refined the roles of certain epitopes, but cannot be used to link to specific genes.

**TRANSCRIPTOMICS ANALYSIS**

Analysis of the transcriptomic data in isolation can also be misleading. Three gene families account for nearly as much of the expression as all the other cell-wall-related families together. The largest cell wall family is the AGPs with 61 members representing more than 18% of cell wall-related gene expression, but the role of individual genes in expansion and maturation is unclear as the localization shows that different epitopes are found in different places.

Aquaporins play a role in vacuolar filling and contribute to turgor pressure, the driving force of expansion. Their expression is present in the elongation zone (zones 2 and 3), naively suggesting that expansion is effected by an increase in turgor pressure. However, recent work (Dyson et al., 2014) shows that pressures remain constant, implying that the role of the aquaporins is to ensure that turgor pressure is not lost by the rapid expansion in cell size.

The third highly-expressed gene family encodes peroxidases, which contains members that are involved in the generation of ROS. These have long been implicated in root growth and development (Gapper and Dolan, 2006; Manzano et al., 2014). However, our data (Supplementary Table 1) show that their expression rises in zone 2, peak in zone 3 and remain high in the later zones. This suggests a role in maturation rather than

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**Table 3** | Mean root lengths in wild type and xth mutants, measured at 7 days after germination, asterisks denote significance at a $p > 0.05$.  

|                | Length (mm ± SEM) | % vs. Col-0 | n  |
|----------------|-------------------|-------------|----|
| **SINGLE MUTANTS** |                   |             |    |
| Col-0          | 37.87 ± 0.39      | –           | 71 |
| xth17-1        | 34.01 ± 0.40*     | 89.8%       | 44 |
| xth17-2        | 34.09 ± 0.46*     | 90.0%       | 39 |
| xth18-RNAi     | 32.75 ± 0.65*     | 86.5%       | 26 |
| xth18-2        | 38.74 ± 0.45      | 102.3%      | 40 |
| xth19-1        | 41.31 ± 0.29*     | 109.1%      | 47 |
| xth19-2        | 32.36 ± 0.48*     | 85.5%       | 19 |
| xth20-1        | 35.17 ± 0.48*     | 92.9%       | 41 |
| xth20-2        | 33.73 ± 0.57*     | 89.1%       | 36 |
| **DOUBLE MUTANTS** |                |             |    |
| Col-0          | 38.02 ± 0.39      | –           | 69 |
| 17-1 × 18-1    | 32.09 (± 0.60)*   | 84.4%       | 25 |
| 17-2 × 18-1    | 27.77 (± 0.61)*   | 73.0%       | 31 |
| 17-2 × 19-1    | 33.63 (± 0.54)*   | 88.4%       | 25 |
| 19-1 × 20-2    | 34.02 (± 1.21)*   | 89.5%       | 20 |
| 17-1 × 20-1    | 35.29 (± 0.40)*   | 92.8%       | 40 |
| 18-1 × 20-1    | 36.13 (± 0.73)*   | 95.0%       | 24 |

**Table 4** | Effect of reduced XTH17 and XTH18 expression on root growth.  

|                | Growth rate (mm/h) | Average mature cell length (μm) |
|----------------|--------------------|---------------------------------|
|                | 3 day  | 4 day  | 5 day  | 6 day  | 7 day  | 3 day  | 4 day  | 5 day  | 6 day  | 7 day  | 172-172 | 172-172 |
| Col-0          | 0.086  | 0.219  | 0.314  | 0.370  | 0.407  | 199.18 | 66       |
| xth17-2        | 0.075  | 0.206  | 0.255  | 0.354  | 0.367  | 184.42 | 66       |
| XTH18-RNAi     | 0.070  | 0.172  | 0.286  | 0.359  | 0.357  | 189.66 | 66       |
| xth17-2 × XTH18-RNAi | 0.054  | 0.193  | 0.240  | 0.319  | 0.319  | 182.06 | 66       |

Growth rates were measured at the indicated days after germination (dag) using NeuronJ. Cortical cell lengths were measured from confocal microscope images, asterisks denote significance at a $p > 0.05$.  

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elaboration, and also possibly in lignification and Casparian strip formation. Although the other -omics data indicated a role for extensins, the genes for these proteins are difficult to define because sequences/functions overlap with other gene families. Hence they were not considered further in this context. Initial carbohydrate biosynthesis peaks in zone 2, where large amounts are needed for both accelerating and subsequent decelerating expansion, but in order to get enough RNA these transcriptomic data face the same issue of resolution as the epitomics work.

**MULTI-OMICS ANALYSIS**

As mentioned above, all these techniques in isolation have their benefits and drawbacks. We therefore combined data from all analysis in a multi-omics approach to identify genes that play an important role in the elongation of *A. thaliana* root cells.

To investigate xyloglucans, we looked for expression patterns in the transcriptome that correlated with the epitope pattern shown by LM15 (peaking in the zone 2). The transcriptomic dataset showed a large number of genes with a similar pattern (Supplementary Table 1). However, filtering the data to include only genes known to affect XyG biosynthesis, we found genes involved in fucose biosynthesis (MUR3, AT2G20370 and GER1, AT1G73250) and a xyloglucan xylosyltransferase, (XXT3, AT5G07720) with $R^2$-values of 0.96–0.98, suggesting that these members from large gene families may be responsible for the observed LM15-XyG signal (Figure 4).

XyG is highest in zone 2, the rapid elongation zone, correlating with studies showing that in vivo, XET activity by XTHs is also highest in this zone (Vissenberg et al., 2000). The general XTH expression signal peaks in zone 2 and 3 (Figure 4), suggesting an important role for XTHs in root elongation. XTHs remodel the cell wall and some are believed to promote growth acceleration (Van Sandt et al., 2007), while others could aid in the deceleration. Different family members have distinct activity dependencies and pH optima, suggesting that some can act as loosening factors, while others could do the opposite (Maris et al., 2009, 2011). If they were only associated with accelerated cell wall actions, then the plant might override the loosening in the deceleration zone, probably by pectin modifications (Micheli, 2001) or peroxidase-mediated cross-linking of other cell-wall components such as structural proteins (Ma et al., 2004; Passardi et al., 2004; De Cnodder et al., 2005). The key XTHs controlling expansion might be expected to be expressed as early as possible to tightly regulate wall extensibility and disruption of these XTHs may therefore show growth defects.

Looking for XTHs with a peak very early in root development and associated with gibberellic acid (GA), a known regulator of cell expansion (Middleton et al., 2012), showed that XTH17 and 18 increase more than 30 fold between zones 1 and 2. One of these is GA-induced (XTH17) and both belong to a subclade of group 1 XTHs (Rose et al., 2002) consisting of 4 genes (XTH17, 18, 19, and 20). XTH17, 18, and 19 are expressed in the elongation zones of the root, and XTH20 is expressed in the vascular tissue of the mature root (Vissenberg et al., 2005). We therefore investigated phenotypes in mutant lines for this subclade. Putative knock out (KO) lines were identified for three of the genes and an RNAi line was created for the fourth. All these lines showed significant growth defects with shorter mature root lengths, shorter mature

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**FIGURE 5 | Combined profiles of pectin epitopes, GAUT1, PME, and PMEI expressions.** Epitomic (columns) and biosynthesis (GAUT1) and modification (PME and PMEI) mRNA expression (lines). Error bars are ±1 SD.
root cells and reduced growth rates (Tables 3, 4). The one exception to this was xth19-1 which, in contrast to the xth19-2 line, showed an increase in root length. The insertion in the xth19-1 line is 3′ to the gene, which may stabilize the mRNA and hence act as an over-expression line. The reduction in growth phenotype was increased in the xth17-2xXTH18-RNAi double mutant (Table 4).

Regarding pectin (or more specifically homogalacturonan), JIM5, and JIM7 recognize esterified pectin, which is present at highest levels in zones 1 and 2 and drops off in the mature root. Pectin is deposited into the cell wall in a highly esterified form, and is typically a component of a loosened cell wall. 2F4 recognizes cross-linked pectin with no more than 40% esterification (Liners et al., 1992) and this is low in zone 1 but rises to high levels.
in zone 2 and beyond. This suggests zone 2 has a mixture of esterified and non-esterified forms and the distribution is increasingly biased toward non-esterified along the shootward axis. This follows the expression of several pectin methyltransferases [PMEs; e.g., PME2 (AT1G53830)], some of which are highly correlated with the epimorphic pattern (Figure 5).

Several PME Inhibitors (e.g., AT5G04970, AT3G10720) also correlate with this pattern (Figure 5), suggesting that the root uses a tight balance of these two groups of enzymes to control the rate of de-esterification (i.e., stiffening) rather than using one to shut the other off. With regard to pectin biosynthesis, the galacturonic acid transferase 1 enzyme (GAUT1, At3g61130) is expressed in zones 1 and 2 and then decreases, which correlates (R² > 0.9) with the pattern of the three antibody epitopes recognizing homogalacturonan (Figure 5 and Supplementary Figure S2).

The transcriptomics data revealed multiple patterns for AGP expression, dominated by maximum levels in zones 2 and 3 (Figure 6). In contrast, the epimorphic signals peak in zone 3, which could be accounted for by several hypotheses. One possibility is that there are delays in synthesis and transport either because of long synthesis and transport times of the proteins, or accumulation for use in zone 3 walls. Alternatively the epimorphic profiles could simply be reflecting AGP accumulation over time.

Different AGPs may play different roles in the cell wall as revealed by the localisome, which was also mentioned before (Ellis et al., 2010). For example, LM6 (AGP or pectic arabinan) localizes to the lateral root cap and epidermis and its epitope may play a role in expansion, while the JIM13 epitope is specific to stel lar cell files and could be part of the process of vascular patterning.

OVERARCHING CONCLUSIONS

It is evident that the individual -omics approaches provide an incomplete picture, and a combination of multiple analyses aids in establishing a clearer picture of the processes involved. In general terms, the transcriptomic dataset suggests the location of cell wall synthesis, whereas the glycan microarray analyses show the accumulation and dilution as these polymers are modified or additional material added. This is best shown with regard to pectins and XyGs. Deceleration in the root elongation rate appears to be linked to a change in the ratio of esterified to non-esterified pectins. The mutation studies have confirmed a role for XTH17, XTH18, and possibly XTH19 in root growth, probably affecting yield threshold, as this was the case in dark-grown hypocotyl cells (Miedes et al., 2013).

POTENTIAL APPLICATION TO OTHER SYSTEMS

Root growth is a system in which the genes for synthesis of cell wall material are expressed in one location, while the molecules themselves might not appear in the wall until later. Different molecules contribute to different aspects of the cell-wall mechanics, which are further affected by subsequent modification and interaction. The multi-omics approach used in this study could be used for other plant structures and translated to other systems where the chronology of gene expression, macromolecular synthesis, and modification contribute to growth or mechanical properties of an organ as a whole. Potential applications include musculoskeletal growth, strength, and brittleness in health, aging and disease states, as well as plant lodging (i.e., the bending or even falling over of stalks leading to reduced crop yields).

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: http://www.frontiersin.org/journal/10.3389/fcell.2015.00010/abstract
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