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
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Hormone-regulated expansins: Expression, localization, and cell wall biomechanics in Arabidopsis root growth

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The authors responsible for distribution of materials integral to the findings presented in this article in accordance with the policy described in the Instructions for Authors (<https://academic.oup.com/plphys/pages/General-Instructions>) are Marketa Samalova and Jan Hejatko.

Abstract

Expansins facilitate cell expansion by mediating pH-dependent cell wall (CW) loosening. However, the role of expansins in controlling CW biomechanical properties in specific tissues and organs remains elusive. We monitored hormonal responsiveness and spatial specificity of expression and localization of expansins predicted to be the direct targets of cytokinin signaling in *Arabidopsis* (*Arabidopsis thaliana*). We found EXPANSIN1 (EXPA1) homogeneously distributed throughout the CW of columella/lateral root cap, while EXPA10 and EXPA14 localized predominantly at 3-cell boundaries in the epidermis/cortex in various root zones. EXPA15 revealed cell-type-specific combination of homogenous vs. 3-cell boundaries localization. By comparing Brillouin frequency shift and AFM-measured Young's modulus, we demonstrated Brillouin light scattering (BLS) as a tool suitable for non-invasive in vivo quantitative assessment of CW viscoelasticity. Using both BLS and AFM, we showed that EXPA1 overexpression upregulated CW stiffness in the root transition zone (TZ). The dexamethasone-controlled EXPA1 overexpression induced fast changes in the transcription of numerous CW-associated genes, including several EXPAs and XYLOGLUCAN:XYLOGLUCOSYL TRANSFERASEs (XTHs), and associated with rapid pectin methylesterification determined by in situ Fourier-transform infrared spectroscopy in the root TZ. The EXPA1-induced CW remodeling is associated with the shortening of the root apical meristem, leading to root growth arrest. Based on our results, we propose that expansins control root growth by a delicate orchestration of CW biomechanical properties, possibly regulating both CW loosening and CW remodeling.

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Introduction

The cell wall (CW) is a fundamental component of plant cells that shapes the plant body and plays key roles in the development and growth of organs, the movement of solutes and nutrients, and in protecting plants from the environment. The CW is also important in controlling cell differentiation and intercellular communication during development (Wolf et al. 2012). CWs provide rigidity; however, they also have the ability to expand. Recent studies of growth regulatory networks suggest that turgor-driven cell expansion is the result of a fine-tuned balance between wall relaxation and stiffening linked via a mechanosensing feedback loop (Sassi and Trass 2015; Braybrook and Jönsson 2016; Willis et al. 2016; Trinh et al. 2021). These regulatory networks comprise transcription factors and plant hormones and allow tight control over the equilibrium between cell division and differentiation, a process fundamental to the growth and development of individual organs in any multicellular organism.

The primary CW consists predominantly of polysaccharides—cellulose, hemicellulose, and pectins. While cellulose microfibrils are responsible for the main load-bearing characteristics of the CW, the presence of hemicellulose and pectins can alter viscoelastic properties of the wall matrix (Wolf et al. 2012; Cosgrove 2018a; Zhang et al. 2021). Once the final cell size is reached, a secondary CW can be deposited in specific cell types, e.g. xylem tracheary elements (for review, see Didi et al. 2015), to provide additional mechanical resistance. CW mechanical properties are modulated by controlling the biochemical composition, as well as the degree and nature of linkages between the CW polysaccharides. Interestingly, wall extensibility may be controlled in discrete regions—so-called biomechanical hotspots—where cellulose–cellulose contacts are made, potentially mediated by trace amounts of xyloglucan (Cosgrove 2014, 2018a, 2018b). These relatively few contact points between cellulose microfibrils may be key sites of CW loosening—a complex process allowing targeted wall expansion.

Expansins, originally described as CW loosening agents during “acid growth” (McQueen-Mason et al. 1992), become activated during CW acidification triggered by a number of stimuli acting on the plasma membrane H⁺-ATPase proton pump (Cosgrove 2005). Mechanistically, expansin-mediated CW loosening neither requires polysaccharide hydrolytic activity nor changes in the composition of the CW; instead, expansins are proposed to disrupt non-covalent bonds between cellulose and components of the surrounding CW matrix, most probably xyloglucans, thus relaxing wall stresses and allowing turgor-driven expansion (McQueen-Mason et al. 1995; Cosgrove 2005; Park and Cosgrove 2012). *Arabidopsis* (*Arabidopsis thaliana*) has 36 members of the expansin superfamily, 26 of them belonging to the family of α -expansins (EXPA) that were experimentally proven to promote CW loosening (Li et al. 2002; Sampedro and Cosgrove 2005). EXPA1 (At1g69530) has been known for decades from experiments with beads loaded with purified expansin that induced

leaf primordia formation on the tomato shoot apical meristem (Fleming et al. 1997). Apart from leaf organogenesis (Reinhardt et al. 1998) and vascular tissue differentiation (Cho and Kende 1998), expansins are involved in root development and growth (Lee and Kim 2013; Pacifici et al. 2018; Ramakrishna et al. 2019), root hair initiation (Cho and Cosgrove 2002), and seed germination (Ribas et al. 2019; Sanchez-Montesino et al. 2019); for more examples, see the recent review by Samalova et al. (2022). Interestingly, NbEXPA1 was shown to be plasmodesmata-specific and functions as a host factor for potyviral infection (Park et al. 2017).

Biomechanical interactions of cells with the extracellular matrix have been shown to be an important regulator of cell fate specification in animal models (Engler et al. 2006; Yang et al. 2016). In plants, changes in CW mechanics are the driving force of growth and development, as predicted by a number of biomechanical models (Geitmann and Ortega 2009; Hamant and Traas 2010; Sassi and Traas 2015; Braybrook and Jonsson 2016; Haas et al. 2020). For instance, in vivo chemical modification (demethylesterification) of homogalacturonan by pectin methylesterases was shown to be sufficient for the initiation of novel flower and floral organ primordia in *Arabidopsis*. In contrast, inhibition of homogalacturonan demethylesterification resulted in the inhibition of normal organ formation (Peaucelle et al. 2008). Importantly, demethylesterification of homogalacturonan was shown to be associated with an increase in CW plasticity as measured via atomic force microscopy (AFM), suggesting that higher elasticity of CWs might be instructive for newly formed organ primordia (Peaucelle et al. 2011). Nonetheless, the opposite response, i.e. CW stiffening, has also been observed upon pectin demethylesterification (reviewed in Levesque-Tremblay et al. 2015). In plants, the importance of biomechanical CW properties has been described mostly in shoot tissues (Reinhardt et al. 1998; Pien et al. 2001; Hamant et al. 2008; Sampathkumar et al. 2014; Landrein et al. 2015; Gruel et al. 2016; Hervieux et al. 2017; Majda et al. 2017; Takatani et al. 2020). However, the biomechanical interactions associated with controlling CW properties are emerging as an important mechanism also guiding root growth and development (Vermeer et al. 2014; Barbez et al. 2017; Pacifici et al. 2018; Ramakrishna et al. 2019; Hurny et al. 2020).

Phytohormones, including auxins and cytokinins, are key players in growth regulation responses and are thus determinants of plant architecture and CW development. The role of auxins in the “acid growth theory” is well known (Cleland 1971; Hager et al. 1971), in which auxin-induced extrusion of protons into the apoplast activates expansins, leading to CW loosening and growth. Recently, auxin signaling has been shown to play a role in both transcriptional and non-transcriptional regulations leading to apoplast acidification (reviewed in Li et al. 2021). Cytokinins are factors controlling the equilibrium between cell division and differentiation (Dello Ioio et al. 2007, 2008) in the root apical meristem (RAM) by delimiting the auxin minimum that triggers the

developmental switch (Di Mambro et al. 2017). Pacifici et al. (2018) proposed that EXPA1 is a direct target of multistep phosphorelay signaling in cytokinin-regulated cell differentiation in the RAM. Recently, cytokinins were also implicated in root growth control, attenuating cell elongation by inducing CW stiffening (Liu et al. 2022).

Brillouin light scattering (BLS) microscopy is an all-optical label-free spectroscopic technique that allows one to spatially map the inelastic Brillouin frequency shift (BFS, ν_B) with a lateral resolution near the diffraction limit (Elsayad et al. 2019; Prevedel et al. 2019), and has been used on live cells (Scarcelli et al. 2015) and tissues (Elsayad et al. 2016; Bacete et al. 2022). The distance of the Brillouin peaks (in GHz) from the central laser frequency is a measure of the local mechanical properties within the confocal volume. Despite probing a distinct elastic modulus in a different frequency regime, the measured BLS has been empirically found to (semi-logarithmically) scale with the lower frequency stiffness defined by Young's modulus. BFS can be expected to be higher for "stiffer" samples and smaller for "softer" cells and tissue (Scarcelli et al. 2015; Andriotis et al. 2019; Gouveia et al. 2019). Young's modulus is typically measured by AFM. This method is based on micrometer or nanometer tissue compressions (indentations) and was developed to precisely measure the mechanical properties of CWs in developing organs and across entire tissue regions (Peaucelle 2014). The measured stiffness (resistance to deformation/indentation) is defined by the measurement of an indentation modulus that best describes the elasticity of the CW scaffolding in the given tissue. AFM can be also used to image CW surface topology at high resolution to detect individual cellulose microfibrils (app. 3 nm in diameter, Zhang et al. 2014) and can be carried out under water, allowing CW imaging of under near-native conditions.

In this work, we set out to localize EXPA1 and its homologs EXPA10, EXPA14, and EXPA15 *in planta* and describe the relationship between the expression of expansins and CW mechanical properties during root cell differentiation. Using both BLS and the more established AFM, we demonstrate CW stiffening upon EXPA1 overexpression accompanied by changes in CW chemical composition leading to remarkable root shortening in Arabidopsis seedlings.

Results

Cytokinins and auxins control EXPA transcription in the Arabidopsis root

For our study, we selected the following expansins, AtEXPA1 (EXPA1, At1g69530), AtEXPA10 (EXPA10, At1g26770), AtEXPA14 (EXPA14, At1g69530), and AtEXPA15 (EXPA15, At2g03090), that have previously been suggested to be under hormonal control (Lee et al. 2007; Bhargava et al. 2013; Pacifici et al. 2018; Ramakrishna et al. 2019).

Type-B Response Regulator binding assays and cytokinin responsiveness (Taniguchi et al. 2007; Zubo et al. 2017; Pacifici et al. 2018; Xie et al. 2018) indicate that EXPA1 is the direct

target of cytokinin-responsive ARABIDOPSIS RESPONSE REGULATOR 1 (ARR1) and its homologs ARR10 and ARR12 (Fig. 1A). EXPA1 responsiveness to auxin could be potentially regulated by AUXIN RESPONSE FACTOR 5 (ARF5) since the corresponding DNA affinity purification (DAP)-sequencing peaks (O'Malley et al. 2016) are located in its promoter (Fig. 1B). To confirm hormonal control over EXPA1, we quantified EXPA1 transcripts using reverse transcription quantitative PCR (RT-qPCR) in wild-type (WT) Arabidopsis seedlings treated with 5 μ M 6-benzylaminopurine (BAP) or 5 μ M 1-naphthaleneacetic acid (NAA, Fig. 1C, Supplemental Materials and Methods, Supplemental Table S1). With the cytokinin treatment, transcript levels were transiently and moderately (3 to 4 times compared with mock-treated control) upregulated over the 4-h time span tested. By comparison, EXPA1 responded slightly more distinctly to the auxin treatment, and its transcript level increased continuously up to 5- to 10-fold at 4 h. However, no or only very weak increase in EXPA1 promoter activity was detected after cytokinin or auxin treatment, respectively (Supplemental Fig. S1A), in either the RAM or the transition zone (TZ) in a *pEXPA1::nls:3xGFP* transcriptional fusion line (Ramakrishna et al. 2019).

EXPA10, EXPA14, and EXPA15 were upregulated by exogenous cytokinins in the WT *Col-0*, and this induction was reduced in the line overexpressing ARR7, type-A ARR, and the negative regulator of cytokinin signaling (Lee et al. 2007). Based on our *in silico* analysis, EXPA10, EXPA14, and EXPA15 might also be direct targets of cytokinin-activated type-B ARRs (Supplemental Fig. S1B). In line with this finding, both EXPA14 and EXPA15 were upregulated by cytokinins; nonetheless, in contrast to a previous report (Pacifici et al. 2018), no positive response was detected with EXPA10 (Fig. 1D).

Altogether, our data suggest moderate and transient EXPA1 and EXPA14 upregulation by cytokinins (in the case of EXPA1 taking place outside the RAM/TZ), with a stronger response being seen for auxin- and cytokinin-mediated upregulation of EXPA1 and EXPA15, respectively. However, no clear effect of exogenous hormone application was detectable for EXPA10.

Expansins localize to the root CW in a specific pattern

Expansins were previously shown to be localized in the CW by immunogold labeling of CWs and Golgi-derived vesicles using an antibody against α -expansin (Cosgrove et al. 2002). However, attempts to visualize expansins in the CW of living plants using a translational fusion with a green fluorescent protein (GFP) have so far failed (Pacifici et al. 2018), perhaps due to the high sensitivity of GFP to the low pH environment. Therefore, we created translational fusions of EXPA1, EXPA10, EXPA14, and EXPA15 with the red fluorescent protein mCherry (Shaner et al. 2004; Supplemental Table S2) under the control of native promoters and confirmed their CW localization in a highly tissue-specific manner in roots (Figs. 1 and 2).

In Arabidopsis root, EXPA1 revealed the strongest expression in the columella and lateral root cap (LRC) of both

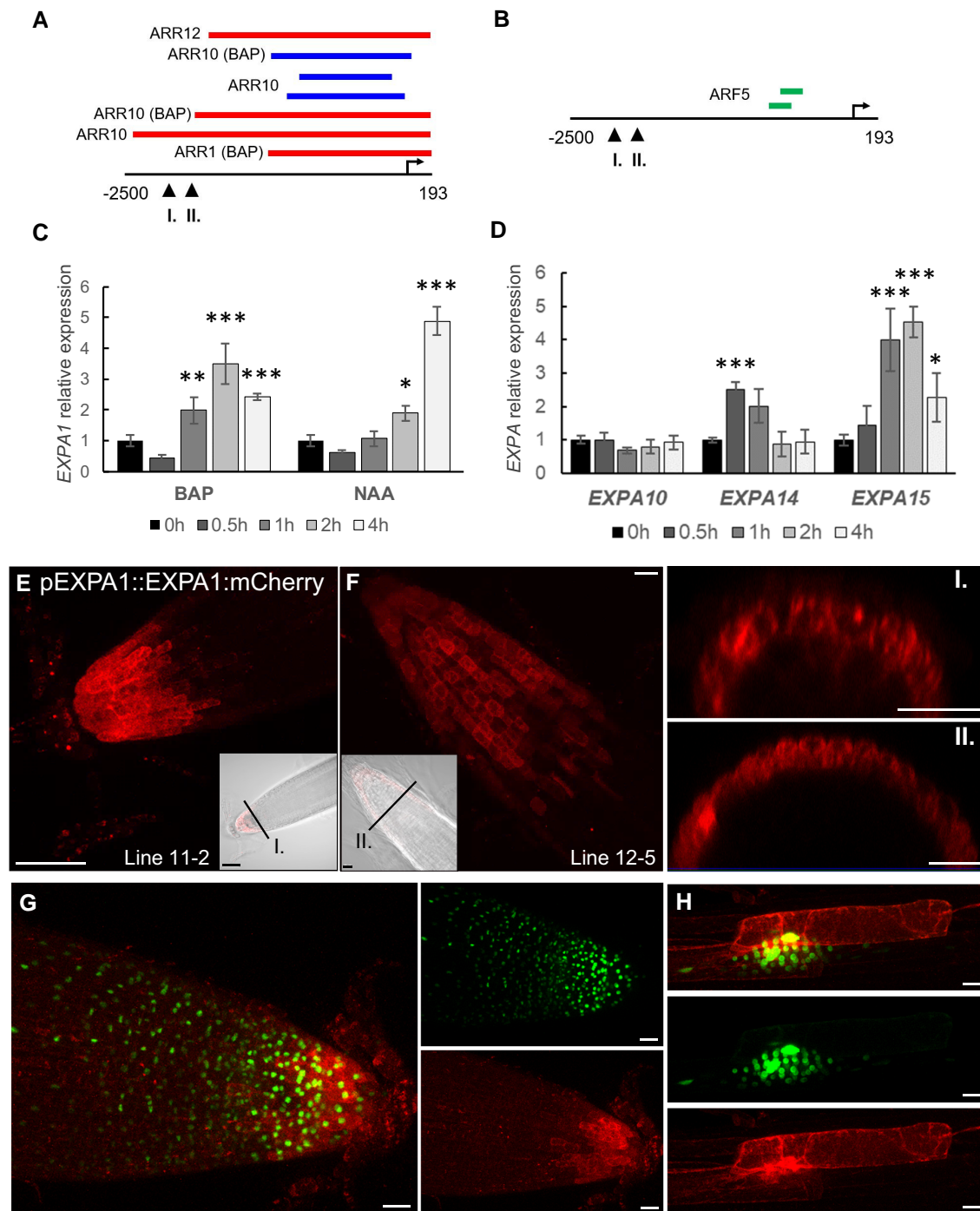


Figure 1. *EXPA1* promoter activity and its localization in roots, and transcript profiling of expansins in response to hormones. **A**) *EXPA1* promoter analysis identifies ChIP-seq- and DAP-seq-derived binding events for transcription factors involved in cytokinin and **B**) auxin signaling pathways. In both **A**) and **B**) red, blue, and green colors depict peaks from Xie et al. (2018), Zubo et al. (2017), and O'Malley et al. (2016), respectively. The coordinates represent positions relative to the transcription start site marked by the arrow. Arrowheads indicate the 5'-end of the *pEXPA1* promoter in this publication (I.) and Pacifici et al. (2018) (II.). **C**) Quantitative real-time PCR and *EXPA1* transcript abundance in 7-d-old Arabidopsis WT roots treated with 5 μ M BAP or 5 μ M NAA for 0.5, 1, 2, and 4 h. **D**) Quantitative real-time PCR-assayed *EXPA10*, *EXPA14*, and *EXPA15* transcript abundance in 7-d-old Arabidopsis WT roots treated with 5 μ M BAP for 0.5, 1, 2, and 4 h. In both **C**) and **D**), *EXPA*s transcript abundance is double normalized to *UBQ10* and mock-treated controls. The experiment was repeated twice with 3 replicas of each sample ($n = 6$), and error bars represent SD. Both experiments were evaluated using simple ANOVA and post hoc Tukey test. Statistically significant differences at alpha 0.05 (*), 0.01 (**), and 0.001 (***) are shown. **E, F**) Z-stack projections of *pEXPA1::EXPA1:mCherry* fusion localized in the LRC and columella of 2 independent single-copy transgenic lines 11-2 **D**) and 12-5 **E**) and their transversal xz optical sections (I. and II.) as indicated by the black lines in the transmitted-light micrograph inserts shown as a single optical section. **G, H**) Z-stack projections of F1 line *pEXPA1::EXPA1:mCherry* (11-2) crossed with *pEXPA1::nls:3xGFP* illustrating a similar pattern of *EXPA1* expression by GFP (green) and mCherry (red) signals in the RAM **F**) and the lateral root primordium **G**). Scale bars correspond to 20 μ m except in **D**) and inset I., where it corresponds to 50 μ m.

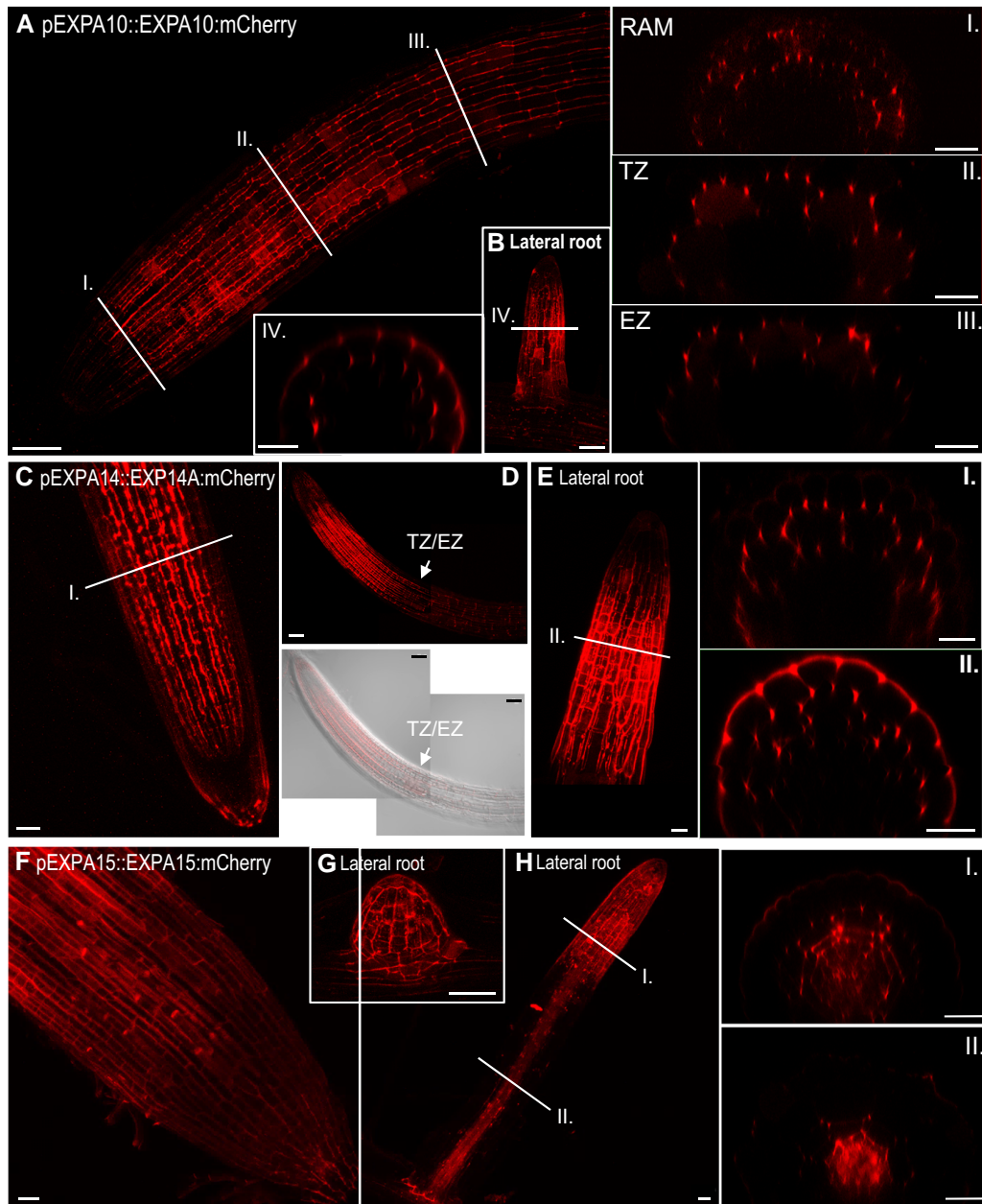


Figure 2. EXP10, EXP14, and EXP15 reveal specific localization in the root tip. **A, B**) *pEXPA10::EXPA10:mCherry* in the primary (**A**) and lateral (**B**) root and transversal xz optical sections (I–IV.) through the RAM, transition (TZ) and elongation (EZ) zones, and the lateral root as indicated by white lines. **C–E**) *pEXPA14::EXPA14:mCherry* in the primary (**C, D**) and lateral (**E**) root and transversal xz optical sections (I. and II.) as indicated by white lines; white arrows point to the TZ/EZ boundary. The specimen in the bottom panel of (**D**) is shown by composite images comprised of 2 partially overlapping fields of view. **F–H**) *pEXPA15::EXPA15:mCherry* in RAM (**F**), emerging lateral root (**G**), lateral root (**H**), and its transversal xz optical sections (I. and II.) as indicated by white lines. Z-stack projections from optical sections taken from the top of the root to the center are shown in all longitudinal plane views. Transmitted-light micrograph inserts show a single optical section. Scale bars correspond to 20 μm except in (**A, B, D, G**), where they correspond to 50 μm .

the primary and lateral roots (Fig. 1, E to G). Interestingly, cells immediately surrounding the developing lateral roots and primordia also strongly expressed *EXPA1* (Fig. 1H). These results were confirmed using an independent transcriptional *pEXPA1::nls:3xGFP* fusion line (Ramakrishna et al. 2019) crossed into the mCherry line background (Fig. 1, G and H). Occasionally, very weak *EXPA1* promoter activity

was detectable in the boundary between the root TZ and the elongation zone (EZ) (Supplemental Fig. S2A); however, no *EXPA1:mCherry* was detectable there (Supplemental Fig. S2B).

EXPA10 was visibly expressed in the cortex layer of the primary root from the meristematic zone up (proximally) to the first lateral roots (Fig. 2A). Unlike *EXPA1*, *EXPA10* was not

expressed in the LRC. Interestingly, in contrast to the rather homogenous distribution of EXPA1 throughout the CW surrounding the LRC/columella cells, we observed a distinct “spotty” localization of EXPA10, predominantly in the cortex/endodermis and cortex/epidermis 3-cell boundaries that were particularly visible on cross-sections of the roots (Fig. 2A, insets I. to III.). In the lateral roots, the EXPA10:mCherry fusion localized predominantly to the epidermis and cortex layers of the transition/elongation zones (Fig. 2B and inset IV.).

EXPA14:mCherry fusion localized to the cortex layer of the RAM up to the TZ/EZ boundary, with the signal not being detectable more proximally (Figs. 2, C and D). The distinct pattern of strong accumulation of the protein in the apoplastic space at the boundary of 3 cells (insets I. and II. for Fig. 2, C and E, Supplemental Fig. S3A) resembles the one observed for the EXPA10:mCherry fusion. In lateral roots, EXPA14 is expressed dominantly in the TZ/EZ boundary (Fig. 2E). However, in contrast to the primary root, EXPA14 is located not only in the lateral root cortex, but also in the epidermal cell layers (Fig. 2E, inset II.).

EXPA15:mCherry was localized to the RAM epidermis (Fig. 2F) and emerging lateral roots (Fig. 2G, Supplemental Fig. S3B) in a relatively uniform pattern; however, the “spotty” pattern at the 3-cell boundary was apparent in

the internal cortex/endodermis. Proximally from the meristem zone, EXPA15 re-localized into deeper (vasculature) layers, again showing a rather homogenous distribution (Fig. 2H, insets I. to II.). It is worth mentioning that in contrast to homogeneously distributed EXPA1 and (partially) EXPA15, EXPA10 and EXPA14 seem to be localized mainly in the CWs oriented in parallel with the longitudinal root axis walls in the RAM/TZ of the main root (Supplemental Fig. S3C).

To confirm the extracellular localization, we activated the (naturally very weak) expression of EXPA1:Cherry in all plant tissues (Fig. 3) using the dexamethasone (Dex) inducible pOp6/LhGR system (Craft et al. 2005; Samalova et al. 2005, et al 2019). After both long (7 d) and short (24 h) Dex induction, the fusion protein accumulated in the cell periphery/apoplastic space in roots but was also visible in the transit through the secretory pathway from the endoplasmic reticulum to the CW. However, because the resolution of a confocal microscope did not allow us to distinguish between CW and plasma membrane localization, we treated the roots with 550 mM sorbitol to allow for plasmolysis. Figure 3G shows that unlike the plasma membrane marker (Supplemental Fig. S4), EXP A1:mCherry remained located at the outer edges of the cells, suggesting that EXPA1 is indeed localized in the CW. Importantly, the CW localization

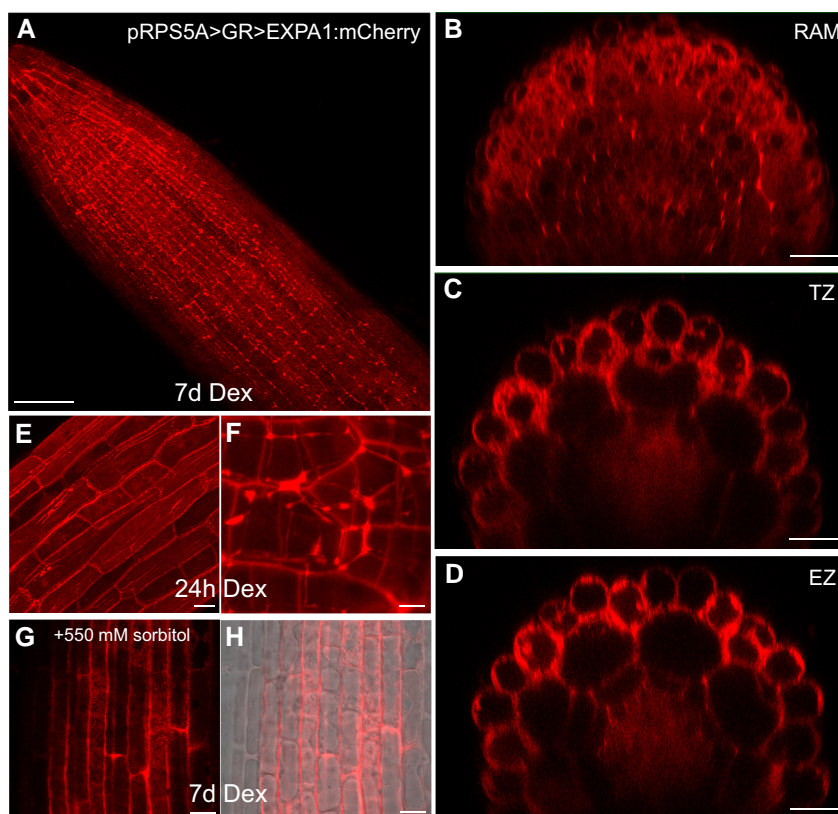


Figure 3. Overproduced EXPA1 localizes to the CW. Z-stack projections of EXPA1:mCherry fluorescence in *pRPS5A>GR>EXPA1:mCherry* seedlings induced by Dex on solid MS medium for 7d in the primary root (A) and its xz optical cross-sections through RAM (B), TZ (C), and EZ (D). E, F) *pRPS5A>GR>EXPA1:mCherry* seedlings induced by Dex in liquid MS medium for 24 h; EZ of the primary root (E) and lateral root (F). G, H) *pRPS5A>GR>EXPA1:mCherry* seedlings induced as in (A) and after 10 min-treatment of the primary root with 550 mM sorbitol. Fluorescence channel (G) and its overlay with transmitted light (H). Scale bars correspond to 20 μm except in (A) (50 μm), (F) (5 μm), and (G & H) (10 μm).

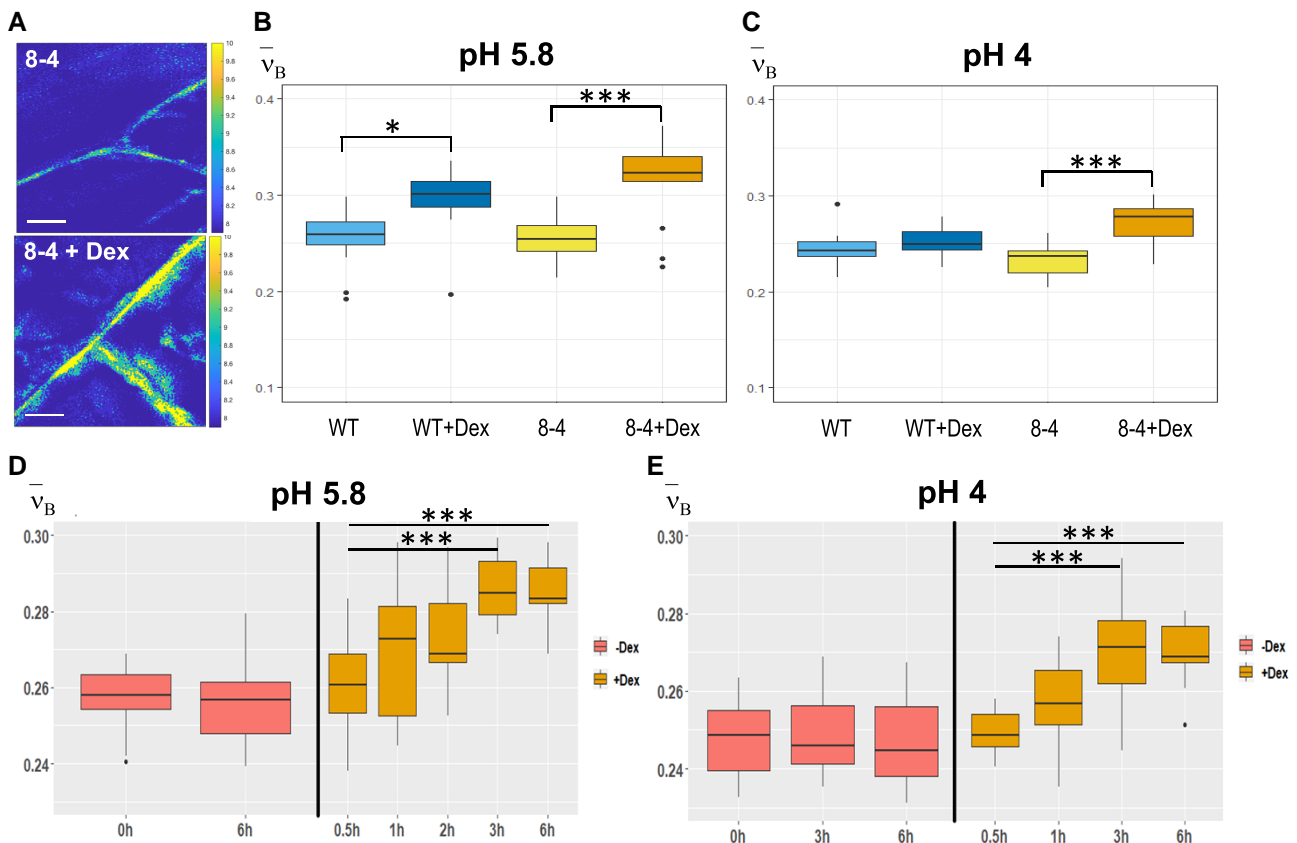


Figure 4. *EXPA1* overexpression stiffens CWs measured via BFS. **A**) Representative images of 2D (*xy*) BFS maps in root cells of 7-d-old Arabidopsis seedlings of *pRPS5A>GR>EXPA1* overexpressing *EXPA1* (line 8-4) grown on MS media pH 5.8 +/- Dex. BFS expressed as Brillouin Elastic Contrast (v'_B) was determined in roots of WT and the 8-4 line grown on solid MS media **(B)** +/- Dex pH 5.8, **(C)** +/- Dex pH 4, **(D)** grown on solid MS (without Dex) for 7 d and induced in Dex-supplemented liquid MS media pH 5.8 for 0.5 to 6 h, or **(E)** liquid MS media pH 4 for 0.5 to 6 h; DMSO was used in -Dex treatments. Medians shown are from at least 4 seedlings and 10 measurements in each category ($n \geq 10$). All experiments were evaluated using simple ANOVA and post hoc Tukey test. Statistically significant differences at alpha 0.05 (*) and 0.001 (***) are shown. Scale bars in **(A)** correspond to 10 μm .

pattern we observed in the case of the Dex-induced *pRPS5A > GR > EXPA1:mCherry* line still resembled the homogenous distribution of *EXPA1:mCherry* driven by its natural promoter in the LRC (compare Figs. 1 and 3). This suggests that the CW localization pattern of *EXPA1* is independent of cell type and level of expression.

To conclude, all assayed expansins show distinct expression and localization patterns. The differences in the localization pattern between *EXPA1* revealing homogenous distribution all around the cell and the “spotty” localization of *EXPA10*, *EXPA14*, and (partially) *EXPA15* suggest differential roles of individual *EXPA*s in controlling root CW properties.

EXPA1 overexpression stiffens the root CW

To characterize its potential importance in controlling CW biomechanical properties, we overexpressed *EXPA1* (without any tag) and generated *pRPS5A>GR>EXPA1* Dex-inducible lines (8-4 and 5-4) using the pOp6/LhGR system as above. Representative 2D BFS maps (Fig. 4A) display the BLS shift in the CW of plants overexpressing *EXPA1* before and after

induction. We quantified the v'_B in roots (TZ/EZ boundary) of 7-d-old Arabidopsis WT and *EXPA1* overexpressing seedlings (line 8-4) grown on MS media pH 5.8 or pH 4 (Fig. 4, B and C, respectively) with or without Dex induction. For technical reasons (to obtain sufficient overlap of point spread function with the CW and, hence, a good CW signal), and following the expression profile of assayed *EXPA* genes (epidermis and/or cortex), we focused on the anticlinal portion of longitudinal CWs in the root epidermis. The plants overexpressing *EXPA1* showed a higher Brillouin Elastic Contrast [v'_B , a BFS normalized to the BFS of distilled water, a measure of longitudinal elastic modulus (Antonacci et al. 2020); see Materials and Methods], for the root CWs on both pH media, suggesting that their CWs are stiffer. Dex-induced stiffening was also observed in the WT at pH 5.8. However, this effect was smaller compared with the v'_B in line 8-4 and was not observed under pH 4 (Fig. 4, B and C). The *pRPS5A>GR>EXPA1:mCherry* lines induced on Dex also displayed higher v'_B , but it was not significantly different from non-induced plants (Supplemental Fig. S5A), perhaps due to lower expression levels of the *EXPA1:mCherry* (Supplemental Fig. S5B). It may also be due to the

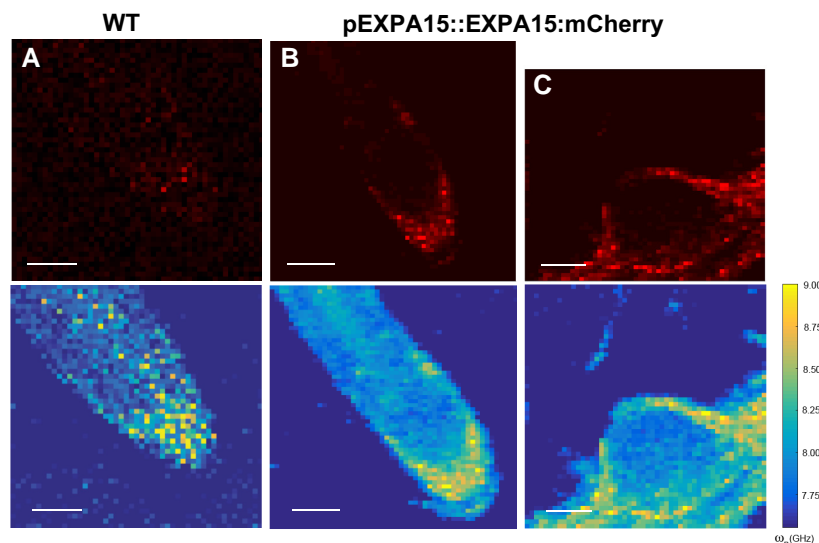


Figure 5. EXPA15 localization overlaps with higher CW stiffness in the root tip as determined using fluorescence emission-Brillouin scattering. Fluorescence images (top row, in red) and BFS (bottom row, false color-coded) of 7-d-old Arabidopsis root from (A) WT and (B) *pEXPA15::EXPA15:mCherry* primary root and (C) emerging lateral root. The fluorescent signal corresponds to EXPA15 expression and overlaps with higher BFS. Scale bars correspond to 100 μm .

possibility that the recombinant protein is less functional in plants. Also, these data clearly demonstrate no effect of Dex alone on the CW stiffening. Importantly, we detected an increase in the ν'_B (CW stiffness) in response to even short-term Dex induction (up to 6 h) that was sufficient to upregulate EXPA1 expression (Supplemental Fig. S6A) and EXPA1:mCherry fluorescence (Supplemental Fig. S6B). With the strong expression line *pRPS5A>GR>EXPA1* (8-4), a statistically significant change in ν'_B was observed as soon as 3 h after EXPA1 induction (Fig. 4, D and E), similarly to long-term induction on media at both pH values (4 and 5.8). To exclude unspecific or side effects of gene overexpression, we determined the spatial map of cell stiffness using fluorescence emission-Brillouin imaging (Elsayad et al. 2016) in the Arabidopsis root over a larger area in a *pEXPA15::EXPA15:mCherry* line, revealing stronger fluorescent signal when compared with *pEXPA1::EXPA1:mCherry*. The regions of higher BFS associated well with the EXPA15 expression domain (Fig. 5), implying a possible role for EXPA15 in controlling CW stiffness. On the other hand, neither BLS nor AFM detected any change in CW stiffness in the root TZ of the *expa1-2* knock-out line (Ramakrishna et al. 2019) when compared with WT (Supplemental Fig. S7).

Since the refractive index (RI) in cells directly associates with the mass content, we applied quantitative cell tomography (using a holotomographic microscope) to measure the RI directly in the Arabidopsis roots in water. Representative maximum intensity projections of RI tomograms are shown in Fig. 6A. Quantitative data analysis confirmed that there are no statistically significant differences across all genotypes and treatment performed in both longitudinal (upper graphs) and transverse (lower graphs) CWs of the early elongating cells in Arabidopsis roots grown at both pH 4 and 5.8 (Fig. 6, B and C). Based on this finding, the effect of RI on the modulus

calculation from the Brillouin shift can be expected to be minimal, supporting the use of ν'_B as a direct indicator of CW stiffness.

To directly measure the “stiffness” of root CWs, we used AFM. AFM-based microindentations apply precisely known forces on a cell through a cantilever and give a deformation value, from which we can extract Young’s modulus of the cell (Peaucelle 2014, 2015). In complex plant tissues and given small deformations, the force to deform material is proportional to the area of indentation allowing the determination of a coefficient of proportionality that is named “apparent Young’s modulus” (Peaucelle 2014). This coefficient depends on the speed of deformation and the mechanical characteristics of the sample. Representative heat maps of the apparent Young’s modulus (E_A) show clear differences when measured in the CW of root TZ/EZ boundary in 7-d-old Arabidopsis WT and EXPA1 overexpressing seedlings (Fig. 7A). The data confirm that the Dex-induced EXPA1 is associated with significantly stiffer root CW ($P < 0.001$) irrespective of whether stiffness is measured on anticlinal or periclinal CWs (Fig. 7, B and C, respectively) on growth media at either pH 5.8 or pH 4 (Fig. 7D). The stiffening effect of EXPA1 overexpression thus seems to be observable at indentation speed of the order of seconds (measured by AFM) and at GHz frequencies through the Brillouin technique.

To validate the ability of BLS to detect not only CW stiffening but also CW softening, we chose the well-described PECTIN METHYLESTERASE 5 overexpressor (PME5oe) lines that upon induction show substantially reduced stiffness of Arabidopsis epidermal hypocotyl cells due to pectin demethylesterification (Peaucelle et al. 2015). Both BLS and AFM revealed softer CW of PME5oe lines compared with the corresponding controls (Supplemental Fig. S8).

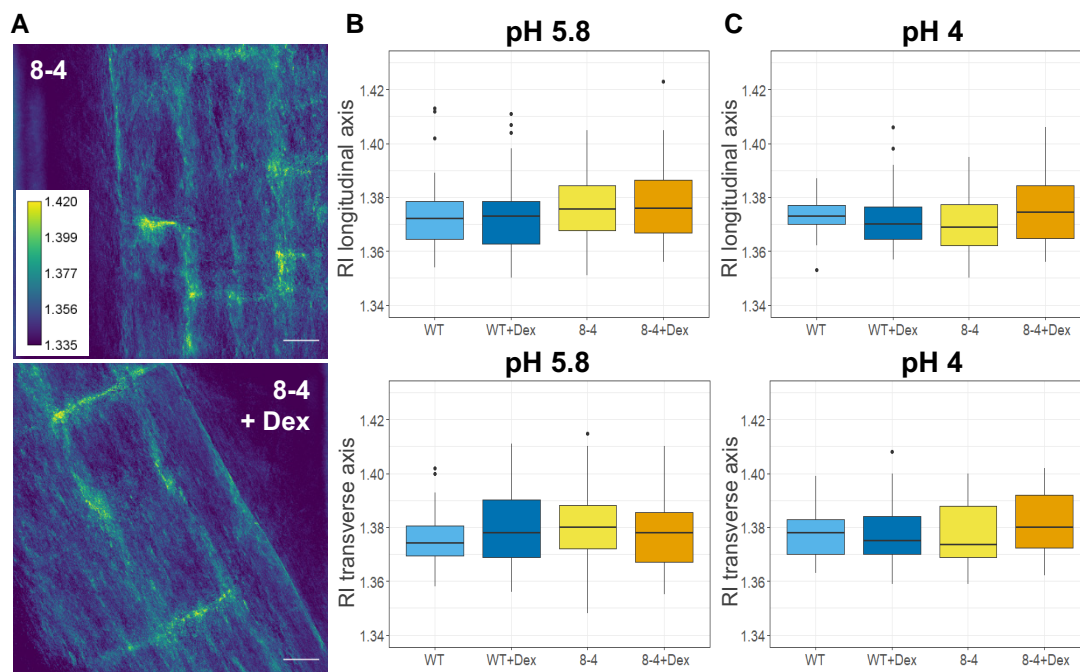


Figure 6. RI measurements of Arabidopsis root CWs reveal comparable CW density in WT and *EXPA1* overexpressing roots. **A)** RI tomograms (maximal projections) of root cells of 7-d-old Arabidopsis seedlings *pRPSSA>GR>EXPA1* overexpressing *EXPA1* (line 8-4) grown on MS media pH 5.8 +/- Dex. Scale bars correspond to 20 μm . The graphs show RI measurements in water (RI 1.330) of roots of WT and the 8-4 line grown on MS media **(B)** +/- Dex pH 5.8 and **(C)** +/- Dex pH 4. Medians from a minimum of 6 seedlings and 30 measurements in each category ($n \geq 30$) of longitudinal (upper) and transverse (lower) CW axes are shown. All experiments were evaluated using simple ANOVA and post hoc Tukey test; no statistically significant differences were observable ($P > 0.1$).

In sum, overexpression of *EXPA1* results in CW stiffening measured at the TZ/EZ boundary using both BLS and AFM. Interestingly, even endogenous *EXPA15* expression seems to be associated with cells displaying higher stiffness within the Arabidopsis root tip.

EXPA1 overexpression leads to CW remodeling

To get mechanistic insight into *EXPA1* overexpression-induced CW stiffening, we chose to use a highly reproducible in situ Fourier-transform infrared (FT-IR) spectroscopy that provides a powerful and rapid assay for wall components and putative cross-links by identifying polymers and functional groups non-destructively (McCann et al. 1992; Mouille et al. 2003; Andres-Robin et al. 2018). FT-IR measurements (spectra 829 to 1,800 cm^{-1}) were performed on the root TZ/early EZ of Arabidopsis WT and the *EXPA1* overexpressing line. Dex induction was performed for a short 3 h period (Fig. 8A) or the seedlings were grown on Dex for a week (Fig. 8B). The most remarkable difference of absorbance observed at both time intervals (appearing after 3 h and being more pronounced after 7 d of *EXPA1* upregulation; arrows in Figs. 8, A and B) could be assigned to ester linkages (around 1,730 cm^{-1}) according to McCann et al. (1992) and Mouille et al. (2003). This suggests a relatively higher esterification degree of pectin in the WT compared with the Dex-treated expansin overexpressor. Collectively, assaying CW composition via FT-IR shows that

induction of *EXPA1* expression leads to rapid (less than 3 h after induction) demethylesterification of pectin.

To elucidate the mechanism underlying the observed *EXPA1*-induced CW remodeling, we assayed the genome-wide transcriptome changes both early (3 h) and later (7 d) after induction of *EXPA1* overexpression. After 3 h of Dex induction, 336 differentially expressed genes (DEGs) were significantly up-regulated and 287 downregulated when compared with non-induced controls (Dex-treated WT *Col-0*, Supplemental Data Set 1). Among the DEGs, significantly enriched gene ontology (GO) terms included plant-type CW loosening (GO0009828), modification (GO0009827), and organization (GO0009664), and apart from the intentionally overexpressed *EXPA1* gene, pointed out a number of other expansin genes being misregulated (Supplemental Table S3). Of those, the most striking was a change in the transcriptional activity of *EXPA2*, reaching almost the fold change levels observed for Dex-induced *EXPA1*; however, the relatively smaller but still highly significant changes were observed also in the activity of several other expansin and expansin-like genes including *EXPA4*, *EXPA8*, *EXPA15*, *EXPB3*, and *EXLB1* (Supplemental Table S4). GO term “CW organization or biogenesis” (GO0071554) comprising *XYLOGLUCAN:XYLOGLUCOSYL TRANSFERASEs* (*XTHs*), particularly *XTH5*, *XTH7*, *XTH11*, and *XTH32*, was also found to be significantly enriched among the DEGs (Supplemental Table S3). After 7 d of Dex induction of the *EXPA1* overexpressing line, only 127 genes were upregulated and 37 downregulated (compared with

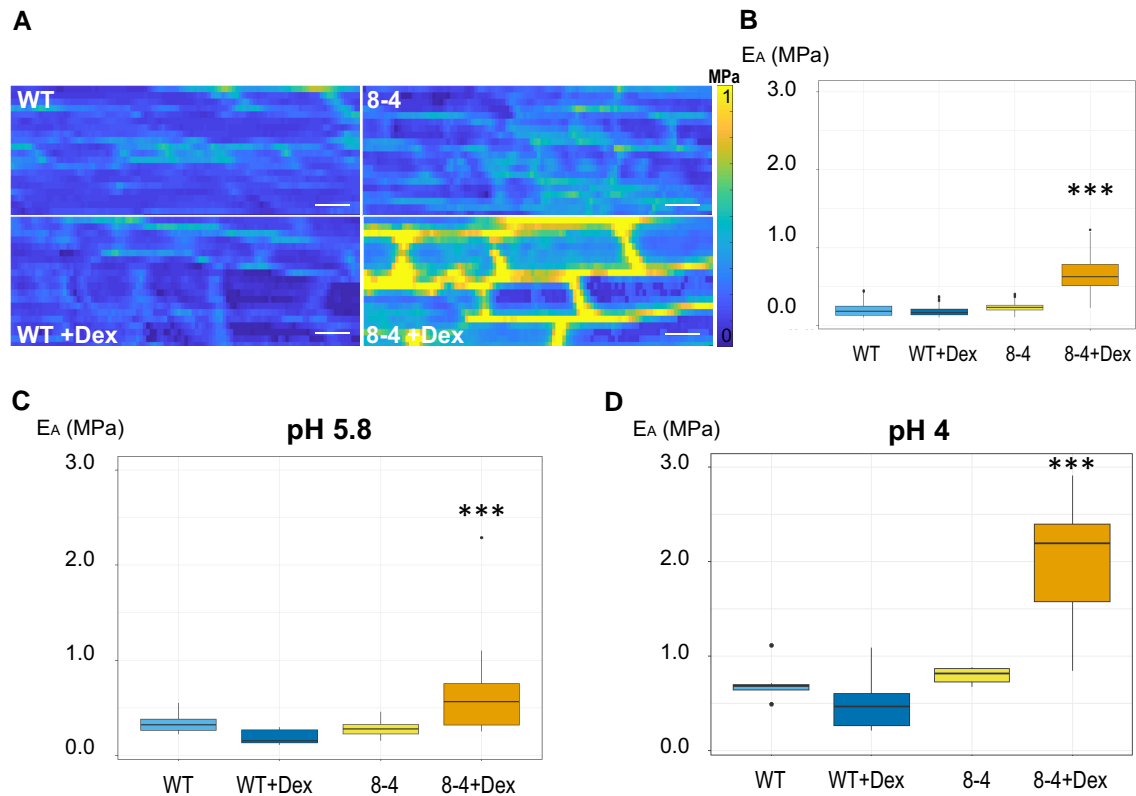


Figure 7. *EXPA1* overexpression stiffens CWs when determined using AFM. **A**) Representative maps of the apparent Young's modulus (E_A) of root cells of 7-d-old Arabidopsis WT and *EXPA1* overexpressing seedlings (line 8-4) grown on MS media +/- Dex, showing differences in E_A (representative of >50). The E_A maps are presented as heat maps, with the respective scale, and show data from 2 successive maps of 60×80 and 60×80 force scans. Each pixel in the E_A map represents the E_A calculated from a single force-indentation curve, and each map consists of 4,800 data points. Images are $100 \mu\text{m}$ in length, scale bars correspond to $10 \mu\text{m}$. **B**) Graphs present the E_A of the anticlinal CW of roots as in **(A)** grown on MS media +/- Dex pH 5.8. Medians shown are from a minimum of 345 measurements in each category ($n \geq 345$). **C, D**) Graphs present the E_A of the periclinal CW of roots as in **(A)** grown on MS media +/- Dex at pH 5.8 **(C)** and pH 4 **(D)**. The E_A plotted on the graphs was determined by sampling data points within the area of interest. Medians shown are from a minimum of 6 measurements in each category ($n \geq 6$). Statistically significant differences at alpha 0.001 (***) are shown. All experiments were evaluated using Mixed Model ANOVA and post hoc Tukey test. Statistically significant differences at alpha 0.001 (***) are shown.

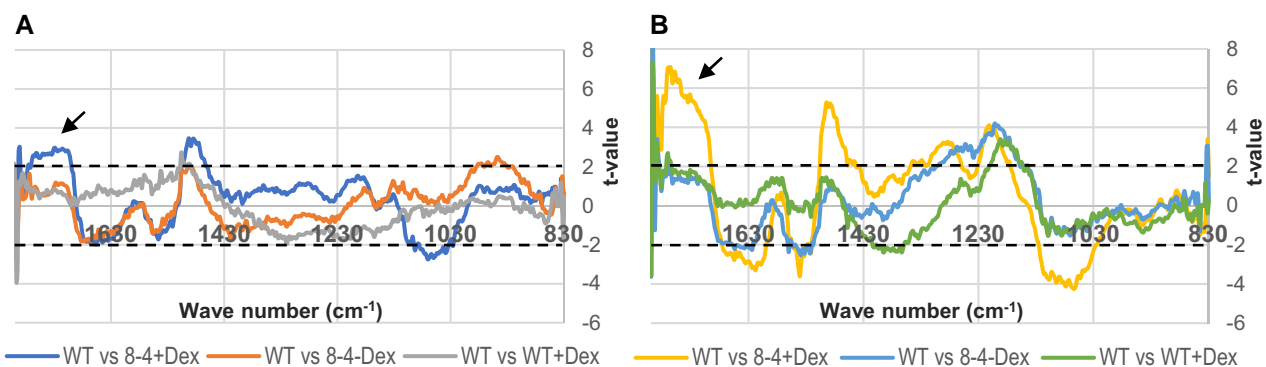


Figure 8. *EXPA1* overexpression changes physical-chemical properties of the CW as determined by FT-IR analysis of root CWs. FT-IR measurements of 7-d-old Arabidopsis WT and *EXPA1* overexpressing seedlings (line 8-4) grown on MS media pH 5.8 without Dex (-Dex) and **(A)** induced with Dex (+Dex) for 3 h or **(B)** grown on Dex for 7 d. Root TZ/early EZ of 4 to 5 individual seedlings from at least 3 experiments ($n > 12$) were analyzed. Student's *t*-test of FT-IR spectra (829 to $1,800 \text{ cm}^{-1}$) was performed to compare WT and the various treatments and genotypes. Graphs show the *t*-value (Y-axis) plotted against the wave numbers (x-axis). Horizontal dashed lines indicate the $P = 0.95$ significance threshold. The black arrows point towards 1730 cm^{-1} .

Dex-treated WT *Col-0*, Supplemental Data Set 1). Compared with the changes observed in the short-term response, apart from the significantly changed expression of *EXPA1*, *EXPA2*, and *EXPA15*, barely any changes in genes related to the CW processes were found (Supplemental Table S5). Dex caused no significant changes in CW modifications and only 15 and 20 genes were up- and downregulated, respectively.

To sum up, our data suggest that *EXPA1* overexpression stimulates prompt changes in the expression of several CW modifying genes and associates with CW remodeling.

EXPA1 overexpression restrains root growth by reducing RAM size

We examined the phenotype of WT and *EXPA1* overexpressing seedlings (*pRPS5A>GR>EXPA1* lines 5-4 and 8-4) grown on Dex continuously for a week. The Dex-induced plants had significantly reduced root length [by 25% to 30% (Figs. 9, A and B)]. The reduction was further enhanced to 40% to 73% when the pH of the growth media dropped from 5.8 to 4. To determine the possible mechanism of root shortening in more detail, we assayed the impact of *EXPA1* overexpression on longitudinal root zonation as defined by Takatsuka et al. (2018). Our data show that RAM (meristematic zone/proximal meristem + TZ, Supplemental Fig. S9) size was significantly reduced by 18% and 29% for line 8-4 grown on media with normal (5.8) and acidic (4) pH, respectively (Fig. 9C). Similarly, cell number (counting in the cortex layer from the quiescent center to the first cell of the EZ) was significantly reduced (Fig. 9D). However, the RAM size/cell number ratio remained the same for each line with or without Dex induction (Fig. 9E), suggesting that the RAM cell number, but not cell length is reduced in the smaller roots. That was confirmed by a more detailed examination revealing that while the number of cells in the TZ remained similar to those of WT, the number of cells in the RAM proliferation zone (called the proximal meristem in Takatsuka et al. 2018) was significantly reduced upon *EXPA1* overexpression (Fig. 9, F and G). Noteworthy, while the aforementioned effects on root length were significant in both lines (5-4 and 8-4) at both pH tested (5.8 and 4), the inhibitory effects on the RAM size in case of weaker overexpressor (line 5-4) were significant only at pH 4, suggesting acidic pH being beneficial for the *EXPA1*-mediated RAM shortening.

In contrast to previous reports, in our hands, *expa1-1* (Pacifci et al. 2018), *expa1-2* (Ramakrishna et al. 2019), as well as our Dex-inducible amiRNA (*amiEX1* lines, Supplemental Table S6), designed to downregulate *EXPA1* and the closely related *EXPA14* and *EXPA15*, did not display any significant phenotype in terms of root or RAM size (Supplemental Fig. S10). However, it should be noted here that following Dex induction, the *amiEX1* lines only reduced *EXPA1* expression by ~50%, *EXPA14* by 60%, and *EXPA15* by 90% (RT-qPCR, Supplemental Fig. S11).

Taken together, while we do not see any effect of *EXPA1* absence/downregulation on root growth and/or RAM size, the overexpression of *EXPA1* results in fewer proliferating

cells in the root meristem thus reduces Arabidopsis root growth.

Discussion

Is there a role for hormonal regulation of *EXPA1* in root growth?

Recently, a role for Arabidopsis *EXPA1* was proposed in the control of cell differentiation (expressed as a function of cell elongation) in cells leaving the meristematic zone of the RAM (Pacifci et al. 2018). Similar to a more recent study (Ramakrishna et al. 2019), our results do not show any statistically significant change in root length and/or RAM size in *expa1-1*, the CRISPR/Cas9 line *expa1-2*, or in our amiRNA lines. Nonetheless, it should be stressed here that *exp1-2* is a hypomorphic allele (Ramakrishna et al. 2019) and our amiRNA lines are knock-down (not knock-out) lines. We did not detect *EXPA1*:mCherry outside the columella/LRC in the root tip and promoter activity (*pEXPA1::nls:3xGFP*) was only occasionally seen in the TZ/EZ boundary and in elongated cells proximal to that. In line with this, we did not detect any changes in CW stiffness in the root TZ of *expa1-2* and no statistically significant upregulation of *EXPA1* was detectable (using absolute fluorescence measurement) in the RAM using the *pEXPA1::nls:3xGFP* lines after 6 and 12 h treatment with both BAP and NAA. These findings suggest that the cytokinin-mediated transcriptional regulation of *EXPA1* may take place in other parts of the root (e.g. cells surrounding LR primordia). Based on our data, we do not exclude a role for *EXPA1* in controlling RAM size, but it probably does so in concert with other EXPAs (see below) and in the columella/LRC, where *EXPA1* is active. The role of (cytokinin-regulated) auxin accumulation in the LRC in the control of RAM size has been proposed recently (Di Mambro et al. 2019). However, even with a more distinct transcriptional regulation of *EXPA1* by NAA, we do not see any substantial and consistent *EXPA1* upregulation in the LRC, both at the level of promoter activity and *EXPA1* protein, thus leaving the functional importance of cytokinin- and auxin-mediated regulation over *EXPA1* in the root tip rather unclear.

In planta expansin localization: specificity rather than redundancy

In previous studies, expansins were located in the CWs using immunolocalization techniques of fixed plant materials (Zhang and Hasenstein 2000; Cosgrove et al. 2002; Balestrini et al. 2005). Transgenic lines carrying *EXPA* genes in translational fusions with mCherry allowed observing the localization of *EXPA* proteins in living plants. Interestingly, our data suggest that the assayed EXPAs differ not only in the spatiotemporal specificity of expression, but their protein products also reveal a distinct localization pattern in specific domains of the root apoplast. Firstly, we see *EXPA10* and *EXPA14* localized predominantly in the longitudinal CWs of elongated root cells. This is similar to the

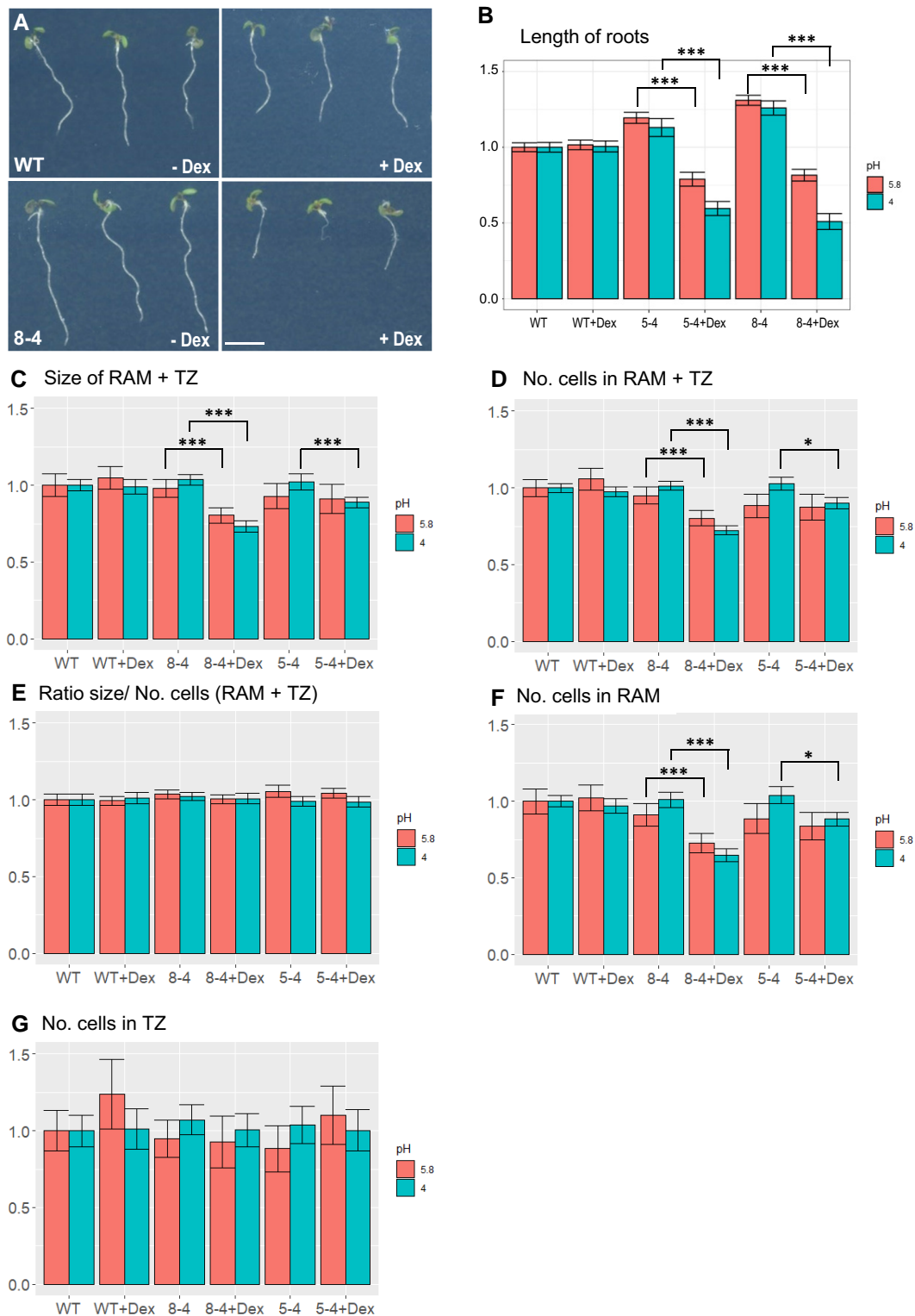


Figure 9. *EXPA1* overexpression reduces root growth by shortening the RAM. **A**) 7-d-old Arabidopsis seedlings of WT (top row) and *pRPSSA>GR>EXPA1* line 8-4 (bottom row) grown on MS media pH4 supplemented either with DMSO (-Dex) or dexamethasone (+Dex). The scale bar is 5 mm. **B**) Length of roots, **(C)** the size of RAM + TZ, **(D)** the total number of cells (No. cells) in RAM + TZ, **(E)** the RAM size/cell number ratio, **(F)** No. of cells in RAM, and **(G)** No. of cells in TZ of 2 independent *EXPA1* overexpressing lines (8-4 and 5-4) grown on MS media +/- Dex pH 5.8 and pH 4 relative to WT. Each experiment was repeated at least 3 times with a minimum of 10 seedlings in each category ($n \geq 30$), and the error bars represent a 95% confidence interval. For all experiments, mixed model ANOVA using random effects for the different experiments was used with the Tukey test as a post hoc test. Statistically significant differences within genotypes and treatments at alpha 0.05 (*) and 0.001 (***) are shown.

situation observed in maize xylem, where the signal obtained after immunolocalization using anti-cucumber expansin antibody was homogeneously distributed in isodiametric (non-elongated) xylem cells, while located mainly in the longitudinal CWs of elongated xylem (Zhang and Hasenstein 2000). Secondly, EXPA10, EXPA14, and EXPA15 (partially) showed a punctate pattern, spatially colocalizing with 3-cell boundaries, possibly in the surrounding intercellular space. These results imply the existence of a factor determining specific localization of individual EXPAs in different CW compartments. The homogenous distribution of EXPA1 throughout the CWs even outside its natural expression domain as seen in the *pRPS5A>GR>EXPA1:mCherry* line suggests that the specific localization pattern is not cell-type-specific, but rather encoded in the amino acid sequence of EXPA1. The existence and the molecular and/or biophysical nature of this factor responsible for homogenous expansin distribution in the CW and/or targeting a subset of EXPAs to specific apoplast domains remain elusive.

The Arabidopsis genome contains 26 genes for α -expansins (Li et al. 2002; Sampedro and Cosgrove 2005), suggesting functional diversification within the subfamily. Specific expression and localization of EXPA1, EXPA10, EXPA14, and EXPA15 together with differential hormonal sensitivity imply possible functional crosstalk among individual expansins. Overlap in their targeted action in the apoplastic continuum covering individual cells might result in the final vectorial change of CW expansion and the highly coordinated cellular behavior underlying root growth including its longitudinal zonation. A similar functional and spatiotemporal specificity including differential hormonal response and shoot cell growth-based zonation was described for *LeEXP2*, *LeEXP9*, and *LeEXP18* in tomato (Caderas et al. 2000; Vogler et al. 2003). The functional importance and possible crosstalk and/or redundancy within the EXPA family in Arabidopsis, however, remains to be demonstrated.

Mechanism of EXPA-regulated CW biomechanics and root growth

Expansins were identified as factors facilitating pH-dependent CW loosening, allowing the cell to extend up to 100 times when compared with its length as a meristematic initial (see Cosgrove 2016 and references therein). To be able to do that, expansins are proposed to modify non-covalent bonds in the cellulose microfibril network, laterally interconnected via xyloglucans bound to the hydrophobic face of cellulose microfibrils (see Cosgrove 2018b and references therein). Consequently, fibril–fibril sliding seems to allow CW extension and in-plane stress release of the multi-lamellate CW structure (Zhang et al. 2019; Zhang et al. 2021). Importantly, expansin-controlled CW loosening allows in-plane stress relaxation and enlargement of CWs, but does not associate with changes in CW viscoelastic properties, as measured by tensile tests (Yuan et al. 2001; Cosgrove 2018a). Thus, the CW “stiffening” or “softening”

(i.e. making the CW less or more deformable to out-of-plane mechanical force, respectively, measurable by, e.g. indentation techniques) is a mechanistically different process and should be distinguished from the EXPA-mediated CW loosening (Zhang et al. 2019; Samalova et al. 2022).

Using 2 independent approaches—non-invasive BLS imaging and AFM—we show here that overexpression of EXPA1, homogeneously distributed throughout the CW, results in increased CW stiffness in root cells. Together with FT-IR analysis, these data suggest that besides its role in CW loosening, EXPA1 could be the modulator of CW viscoelastic properties by regulating CW remodeling. In general, EXPAs might modulate CW chemical composition by controlling the access of CW remodeling enzymes to their substrate either by relaxing CW structure (thus allowing different hydrolases to access their substrates) or by changing substrate availability due to EXPA binding to non-cellulose components of the CW matrix including hemicelluloses and pectin (McQueen-Mason and Cosgrove 1995; Georgelis et al. 2011; Wang et al. 2013). This was experimentally demonstrated for the putative carbohydrate-binding module (CBM) from strawberry expansin 2 (CBM-FaExp2). The binding of CBM-FaExp2 to CW carbohydrates was associated with a reduction in the activity of CW degrading enzymes including pectinase in the in vitro assays. Interestingly, CBM-FaExp2 shows a high level of similarity with AtEXPA1 and AtEXPA2 (Nardi et al. 2013). Thus, the high abundance of EXPA1 might interfere with the accessibility of pectin to pectin-modifying enzymes like methylesterases, pectin acetyl esterases, polygalacturonases, or pectate-like lyases. This would potentially lead to pectin modifications and subsequent changes in CW rheological properties (Senechal et al. 2014). Our findings imply an interference between EXPA1 overexpression and methylesterification of pectin, previously demonstrated to be responsible for controlling CW viscoelasticity (reviewed in Levesque-Tremblay et al. 2015). Specifically, the FT-IR data suggest that EXPA1 is able to trigger rapid pectin demethylesterification that has been demonstrated to allow both CW stiffening and softening, possibly dependent on the demethylesterification pattern (Levesque-Tremblay et al. 2015). Potential functional artifacts due to high levels of EXPA1 overexpression used in our study cannot be completely excluded. Nonetheless, the ability of EXPA1 to induce CW stiffening as soon as 3 h after Dex addition (leading to an order of magnitude lower EXPA1 mRNA amount compared with the long-term induction) and an association between CW stiffness and EXPA15 expression (partially overlapping with EXPA1 expression pattern) in living Arabidopsis roots implicates expansins in the spatial-specific control of CW viscoelasticity, as proposed recently (Samalova et al. 2022). On one hand, it should be highlighted here that the co-localization of EXPA1 and EXPA15 with the CWs revealing higher relative stiffness in columella/LRC may be coincidental and does not provide evidence of any causal link between EXPA1/EXPA15 and CW stiffening. On the other hand, however, changes in the Raman spectra of pericycle CWs in *expa1-1* provide strong

evidence for the association of EXPA1 activity and CW composition in the EXPA1-dependent asymmetric pericycle (lateral root founder cell) expansion (Ramakrishna et al. 2019).

Besides the interference with pectin-modifying enzymes, another mechanism of EXPA1-induced CW remodeling could be EXPA1-mediated transcriptional regulation of CW modifying/loosening genes. By far the most upregulated one was EXPA2 ($\log_2FC = 5.718$). The gibberellic acid-inducible EXPA2 was shown to control endosperm expansion, being necessary for seed germination in Arabidopsis (Yan et al. 2014; Sanchez-Montesino et al. 2019). Interestingly, EXPA1 was found to be strongly downregulated in the mutant deficient in NAC25, the TF mediating the gibberellic acid-induced expression of EXPA2. Furthermore, the regulators mediating the GA-controlled endosperm expansion comprised also a number of genes we found to be misregulated by EXPA1 overexpression including several EXPAs (EXPA8, EXPA15) and XTHs (XTH5) (Sanchez-Montesino et al. 2019). This suggests that EXPA1 and EXPA2 together with several genes encoding for enzymes involved in the CW remodeling might be a part of the regulatory network involved in the (GA-regulated) cell elongation/expansion.

CW stiffness associated with upregulated EXPA1 seems to downregulate root growth by reducing RAM size. This suggests a mechanism connecting biomechanical CW properties with control over cell division in the RAM, as also implied by the previously demonstrated cell proliferation regulatory role for CW integrity (CWI) signaling (Gigli-Bisceglia et al. 2018). To maintain the functional CW, plants use specialized systems for monitoring and initiating compensatory responses often amplified through feedback processes. The sensors recognizing the CWI status and triggering the downstream transcriptional regulations are kinases from the *Catharanthus roseus* receptor-like kinase1-like (CrRLK1L) and Wall-associated kinase subfamilies. These kinases are able to interact with pectin, suggesting their possible role in sensing the disturbance in the CW matrix (see Baez et al. 2022 and references therein). CWI reacts to CW perturbation by changes in gene expression leading to CW deposition and remodeling as possible compensatory mechanisms (Voxeur and Hofte 2016; Baez et al. 2022). This implies time-dependent changes in the CWI-induced responses. In line with that, we see much stronger transcriptome changes early (3 h) after inducing EXPA1 OE (623 DEGs), while a much lower number of DEGs (164) was identifiable after the long-term (7 d) EXPA1 upregulation, thus hinting to the involvement of a mechanism mediating adaptation to EXPA1 overexpression. We may speculate on the involvement of CWI signaling-activated CW remodeling as a possible compensatory mechanism responding to CW disturbance and allowing to reach the new equilibrium (see Vaahtera et al. 2019; Samalova et al. 2022 and references therein).

BLS vs. AFM

In terms of the methodological approach used, there are important differences between the longitudinal elastic modulus

(M) measured by BLS and Young's modulus (E) measured via AFM (Prevedel et al. 2019). The BLS measured M is well known to be very sensitive to the level of hydration (Palombo et al. 2014; Wu et al. 2018; Andriotis et al. 2019) and temperature (Berne and Pecora 2000), and any comparisons thus need to be made under the same thermodynamic conditions and hydration levels. However, as these can be assumed to be similar between the different samples measured, variations between samples can be interpreted as being due to changes in the mechanical properties in the probed regime. The comparable trend of the quasi-static Young modulus (measured by AFM) and BLS measured v'_B observed here is consistent with observations in a diverse set of biological samples (e.g. Scarcelli et al. 2015; Andriotis et al. 2019; Gouveia et al. 2019) suggesting that here too the latter may serve as a proxy for stiffness. We note, however, that at very high hydration levels (much higher than in the system studied here), the relation between the 2 can be expected to break down (Wu et al. 2018).

Conclusions

Based on our and others' results, we propose an important regulatory role in controlling root growth and development in Arabidopsis for the tightly controlled spatiotemporal specificity of expansin expression and the localization of their protein products into distinct domains of the extracellular matrix, possibly in conjunction with hormone-regulated pH distribution in the root apoplast (Barbez et al. 2017; Pacifici et al. 2018; Großholz et al. 2021; Li et al. 2021; Serre et al. 2022). We also propose that apart from mediating CW loosening, expansins may control viscoelastic properties of CWs by regulating CW remodeling. That implicates expansins as important modulators of plant development controlling CW biomechanical properties through a complex regulatory mechanism.

Materials and methods

Plant cultivation conditions and dexamethasone (Dex) induction

Standard MS medium supplemented with 1.5% (w/v) sucrose, 0.8% (w/v) plant agar (Duchefa), and pH 5.8 (adjusted with KOH) or pH 4 (adjusted with H_2SO_4) was used. Arabidopsis (*A. thaliana*) plants were cultivated in growth chambers under long-day conditions (16 h light/8 h dark) at 21 °C in Petri dishes or in soil, with a light intensity of $150 \mu\text{M m}^{-2} \text{ s}^{-1}$ and 40% relative humidity. Dex induction was performed by adding 20 μM Dex into MS media on plates; for short time exposure (1 to 6 h or 24 h), seedlings were placed into liquid MS media containing 20 μM Dex (as described in Samalova et al. 2019). DMSO at the same concentration was used instead of Dex as a control.

Confocal laser scanning microscopy and image analysis

We used a Zeiss LSM 880 laser scanning microscope to localize EXPA:mCherry fusions. mCherry fluorescence was detected at 580 to 650 nm with 561-nm HeNe laser excitation and eGFP at 490 to 550 nm with a 488-nm Argon laser line with a detector gain set to 800. Z-stack projections are shown at maximum intensity. Fluorescence quantification was done using CellProfiler (mCherry) and Imaris (nls:3xGFP) software. To measure RAM size, 7-d-old Arabidopsis seedlings were stained with $30 \mu\text{g ml}^{-1}$ propidium iodide for 5 min, scanned at 590 to 650 nm with 488-nm excitation, and measured using ZEN 3.0 software. The roots were imaged using the 40 \times /1.2 water-corrected C-Apochromat or the 25 \times /0.8 immersion-corrected Plan-Apochromat objectives.

BLS microscopy

BLS is the inelastic scattering of light from inherent or stimulated high-frequency acoustic vibrations in a sample, the speed of which is directly related to the elastic modulus of the material (Berne and Pecora 2000). The BFS (ν_B) is proportional to the acoustic phonon velocity, which is in turn proportional to the square root of the high-frequency longitudinal elastic modulus (M). As such, the Brillouin frequency can serve as a proxy for the mechanical properties of the sample. In particular, M is closely related to the compressibility of the sample and has empirically been observed to scale semi-logarithmically with Young's modulus (E) as measured by AFM in diverse samples including live cells (Scarcelli et al. 2015). An accurate calculation of M requires knowledge of the ratio n^2/ρ , where n and ρ are the RI and the mass density, respectively, in the probed region of the sample. While it can often be assumed by virtue of the Lorentz–Lorenz (LL) relation that n^2 will scale with ρ (Zhao et al. 2011), such that explicit knowledge of the ratio n^2/ρ at each probed region is not required, the validity of the LL-relation in complex multicomponent structures including the CW cannot be rigorously justified. As such, we present results in terms of a dimensionless frequency shift previously introduced as the Brillouin Elastic Contrast, $\nu'_B = \nu_B/\nu_B^{(w)} - 1$, where $\nu_B^{(w)}$ is the measured BLS frequency of distilled water (Antonacci et al. 2020). As is the case for the BFS, ν'_B scales with the square root of M . Moreover, it is independent of the probing wavelength; it can correct for slight variations in temperature between measurements and allows for more direct comparisons of measurements between instruments employing different probing wavelengths.

Brillouin microscopy was performed using the in-house Brillouin confocal microscope described in Elsayad et al. (2016). Excitation was via a single-mode 532 nm laser (Torus, Laser Quantum, DE). A dual cross-dispersion Virtual Imaged Phase Array (VIPA) spectrometer (Scarcelli et al. 2015) with a Lyott Stop (Edrei et al. 2017) was used to measure BLS spectra. The spectral projection was

measured on a cooled EM CCD camera (ImageEMX II, Hamamatsu, JP). The spectrometer was coupled to an inverted microscope frame (IX73, Olympus, JP) via a physical pinhole with an effective size of 1 Airy Unit to ensure optimum confocal detection. After the pinhole, a dichroic mirror was used to outcouple light with wavelengths longer than 536 nm to a fluorescence spectrometer (Ocean Optics QE Pro, USA) to detect the fluorescence signal assuring pixel-to-pixel association with the measured Brillouin spectra. To acquire Brillouin maps, samples were scanned in x , y , and/or z using either a 3-axis long-range Piezo-stage (Physik Instrumente, DE) or a motor stage (ASI, USA), both mounted on top of the inverted microscope frame. Light could also be coupled out through a second port on the microscope frame using a long-pass filter (AHF, DE) and a tube lens to a compact sCMOS camera (Thorlabs, DE) allowing us to locate samples and regions of interest (in wide-field transmitted-light conditions illuminating the sample from the top with a halogen lamp) as well as to monitor the position being probed during scanning.

All hardware was controlled using LabView (National Instruments, USA)-based software developed by THATec (DE), especially for our microscope. The 16-bit depth spectral projection image for each position in the spatial scan was exported from the native THATec format into MATLAB (Mathworks, DE), where a custom-written code was used for analysis. This code (see also Elsayad et al. 2016) used 2 calibration spectra (of triple distilled water and spectroscopic grade ethyl alcohol) measured before and after each set of scans. These were used for registration of the spectral projection onto a frequency scale, based on the calculated dispersion for a dual-VIPA setup in the paraxial approximation regime. The alignment of the spectrometer was such that maximal energy was transferred into a single diffraction order. Due to the spatial masking of the elastic scattering peaks at the 2 intermediate imaging planes in the spectrometer, the spectral projection consisted of only 2 inelastic scattering peaks corresponding to the so-called Brillouin Stokes and anti-Stokes scattering peaks.

All data analysis was performed in MATLAB (Mathworks, DE) using custom-written scripts (Elsayad et al. 2016). Spectral phasor analysis (Elsayad 2019) was used to obtain initial parameter estimates for peak positions and widths which were subsequently inserted into a non-linear least squares fitting algorithm that fitted 2 broadened Lorentzian functions (Voigt functions) to obtain the 2 peak positions, from which the BFS could be obtained. The BLS spectra were also deconvolved in phasor space using a response function obtained from measuring the attenuated Rayleigh scattering inside the respective samples (by opening the spatial masks).

For all scans, the laser power at the sample was between 1 and 5 mW, and the dwell time per point, which was also the acquisition time of each spectrum, was 100 ms. Cells were inspected (by transmitted-light wide-field illumination) and appeared healthy and unperturbed after the experiments, suggesting that BLS measurements had no negative or phototoxic effects.

A 1.4 NA objective was used for excitation and detection (back-scattering geometry). As such a broad range of scattering wavevectors is probed, and one effectively probes directionally averaged elastic moduli. As a direct consequence of probing a broad spectrum of wavevectors, the Brillouin spectra are broadened as predicted from the momentum-energy conservation equations describing the scattering processes. The so-called Brillouin scattering peak position is, however, not noticeably modified to within experimental uncertainties, as was verified by reducing the numerical aperture of excitation and detection on the studied samples using an iris in the beam path.

Roots of 7-d-old Arabidopsis seedlings were scanned at the early EZ, the size of the scan was $25\ \mu\text{m} \times 25\ \mu\text{m}$ (50×50 pixels), with step size typically 500 nm (or 250 nm for larger scans) using the piezo-stage.

RI tomography

RI tomograms were acquired on a holotomographic microscope with rotational scanning 3D Cell Explorer (Nanolive SA, Lausanne, Switzerland) with Nikon BE Plan 60 \times NA 0.8. The size of the acquired tomogram was $93.1 \times 93.1 \times 35.7\ \mu\text{m}$ (*xyz*). Samples were measured in water (reference RI 1.330). Steve 1.6.3496 software (Nanolive SA) was used for image acquisition. Subsequent image analysis was performed in ImageJ 1.52q (NIH, USA) on a max projection of tomography data. The median RI at the CW of the longitudinal and transverse axes of the cell was then extracted.

Atomic force microscopy

Roots of 7-d-old Arabidopsis seedlings were immobilized on glass slides and surrounded by stiff agarose. Approximate TZ/early EZ was defined based on the visual landmark observed through a bright field microscope. In order to extract the mechanical properties of only the outer CW, the maximum indentation force was set to 60 nN to archive a maximum indentation of no more than 80 nm. The cantilever used was “Nano World” (Nanosensors Headquarters, Neuchâtel, Switzerland) SD-R150-T3L450B tips with a spring constant of 0.15 to 1.83 N/m (the one used was estimated to be 0.781 N/m) with silicon point probe tips with a 150-nm radius.

In order to extract the mechanical properties of the periclinal CW, the maximum indentation force was set to 70 nN to achieve a maximum indentation of no more than 80 nm. The cantilever used was “Nano World” (Nanosensors Headquarters, Neuchâtel, Switzerland) SD-Sphere-FM-S-10 tips with a spring constant of 0.15 to 1.83 N/m (the one used was estimated to be 1.1 N/m) with silicon point probe tips with an 800-nm radius.

All force spectroscopy experiments were performed as previously described (Peaucelle 2014; Peaucelle et al. 2015; Feng et al. 2018). Briefly, the stiffness of samples was determined as follows: an AFM cantilever loaded with a spherical tip was used to indent the sample over a $60 \times 100\ \mu\text{m}$ square area, within the area 60×100 measurements were made resulting in 6000 force-indentation experiments; each force-indentation experiment was treated with a Hertzian indentation model to extract the apparent Young modulus (E_A); each pixel in the stiffness

map represents the apparent Young modulus from 1 force-indentation point. The E_A was calculated using JPK Data Processing software (ver. Spm-4.0.23, JPK Instruments AG, Germany), which allows for a more standardized analysis than the estimation of the E_A using a standard Hertzian contact model (Peaucelle 2014, 2015). As is typically the case in nano-indentation experiments, only the retraction curve was used in our analyses. A Poisson ratio of 0.5 was assumed for the material. Range distribution of E_A from 0.2 to 3 MPa in 1-MPa binned groups was calculated using MATLAB. The topography map generated from the contact point calculation permitted the selection of the periclinal or anticlinal CW perpendicular to the indentation for the analysis.

FT-IR analysis

Seven-day-old Arabidopsis seedlings of WT and *EXPA1* overexpressing line (8-4) grown on MS media pH 5.8 \pm Dex were collected and stored in ethanol. They were then rehydrated in water overnight prior to being laid out on gold-coated slides and left to dry for a few hours at room temperature. Spectra were collected using a Nicolet iN10 Infrared Imaging Microscope (ThermoFisher) in a $30 \times 30\ \mu\text{m}$ window. Root TZ of 4 to 5 individual seedlings from at least 3 experiments ($n > 12$) were analyzed for each treatment/genotype. Eight interferograms were collected in transmission mode with $8\ \text{cm}^{-1}$ resolution and coadded to improve signal-to-noise ratios. The collected spectra were baseline corrected and normalized using previously described procedures (Mouille et al. 2003).

Statistical analysis

For statistical analyses, simple ANOVA and post hoc Tukey test were used. For pairwise comparisons in repeated experiments, mixed model ANOVA using random effects for the different experiments was used with the Tukey test as a post hoc test. In the case of non-normal count data (e.g. No. of cells), a Poisson mixed model was used to identify differences between genotypes. The lme4 package in R was used for the implementation of the mixed models (Bates et al. 2015).

Accession numbers

Sequence data from this article can be found in the GenBank/EMBL data libraries under the following accession numbers: *EXPA1* (At1g69530), *EXPA10* (At1g26770), *EXPA14* (At5g56320), and *EXPA15* (At2g03090).

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Author contributions

M.S. and J.H. designed the research, M.S., A.M., K.E., A.P., E.G., J.G., E.V.Z., E.V.U., D.B., V.B., and G.M. performed the research, M.S., K.E., A.P., J.G., I.S., E.V.Z., M.S., N.B., P.A., V.B., G.M., and J.H. analyzed the data, and M.S., K.E., A.P., J.G., G.M., and J.H. wrote the manuscript.

Supplemental data

The following materials are available in the online version of this article.

Supplemental Figure S1. *EXPA1* is not inducible by cytokinins in the root tip, and cytokinin-responsive transcription factors bind to *EXPA* promoters.

Supplemental Figure S2. *EXPA1* activity is only occasionally detectable in the root TZ, while *EXPA1* signal is delimited to the columella/lateral root cap.

Supplemental Figure S3. Cell-type-specific localization of *EXPA14*, *EXPA15*, and *EXPA10* that localizes predominantly to the longitudinal cell walls.

Supplemental Figure S4. Confocal imaging of a plasma membrane marker line U_{BQ10}::YFP-PIP_{1,4} before and after plasmolysis.

Supplemental Figure S5. Low-level *EXP1A::mCherry* overexpression does not reveal cell wall stiffening, as determined using Brillouin light scattering microscopy and RT-qPCR of Dex-induced levels of expression of *EXP1A::mCherry* and *EXPA1* in selected lines.

Supplemental Figure S6. Short-time Dex induction is sufficient to increase *EXPA1* expression, as detected by RT-qPCR and *EXPA1::mCherry* fluorescence.

Supplemental Figure S7. *expa1-2* knock-out does not show changes in cell wall stiffening in the TZ, as determined using Brillouin light scattering microscopy and atomic force microscopy.

Supplemental Figure S8. Comparison of BLS and AFM measurements on *PME5oe* overexpressing seedlings.

Supplemental Figure S9. An example of CLSM images used to determine the size of RAM.

Supplemental Figure S10. Root growth is not affected in *expa1* knock-out and *EXPA* knock-down lines.

Supplemental Figure S11. RT-qPCR of Dex-induced reduction of expansin expression in *amiEX1* lines.

Supplemental Materials and Methods.

Supplemental Table S1. List of primers used for RT-qPCR.

Supplemental Table S2. List of primers used for expansin cloning.

Supplemental Table S3. GO terms significantly enriched among DEGs identified 3 h after Dex-induced *EXPA1* overexpression.

Supplemental Table S4. CW-related DEGs identified 3 h after Dex-induced *EXPA1* overexpression.

Supplemental Table S5. CW-related DEGs identified 7 d after Dex-induced *EXPA1* overexpression.

Supplemental Table S6. List of primers used for cloning of *amiEX1*.

Supplemental References.

Supplemental Data Set 1. DESeq2 data analysis.

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