The Effects of High Steady State Auxin Levels on Root Cell Elongation in Brachypodium

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Running title: High auxin steady state in Brachypodium roots

SYNOPSIS: Analysis of Brachypodium mutants with increased cellular steady state auxin content and root cell elongation sheds light on the importance of the Acid Growth Theory of Plant Cell Elongation in roots.
Abstract

The long-standing Acid Growth Theory of Plant Cell Elongation posits that auxin promotes cell elongation by stimulating cell wall acidification and thus expansin action. To date, the paucity of pertinent genetic materials has precluded thorough analysis of the importance of this concept in roots. The recent isolation of mutants of the model grass species *Brachypodium distachyon* with dramatically enhanced root cell elongation due to increased cellular auxin levels has allowed us to address this question. We found that the primary transcriptomic effect associated with elevated steady state auxin concentration in elongating root cells is up-regulation of cell wall remodeling factors, notably expansins, while plant hormone signaling pathways maintain remarkable homeostasis. These changes are specifically accompanied by reduced cell wall arabinogalactan complexity but not by increased proton excretion. On the contrary, we observed a tendency for decreased rather than increased proton extrusion from root elongation zones with higher cellular auxin levels. Moreover, similar to *Brachypodium distachyon*, root cell elongation is, in general, robustly buffered against external pH fluctuation in *Arabidopsis thaliana*. However, forced acidification through artificial proton pump activation inhibits root cell elongation. Thus, the interplay between auxin, proton pump activation and expansin action may be more flexible in roots than in shoots.

Keywords: Acid Growth Theory / auxin / Brachypodium / Arabidopsis / cell elongation / cell wall / AGP
### Introduction

Coordinated cell division and expansion is crucial for plant organogenesis because cell walls restrict the movement of cells relative to each other (Cosgrove, 1999; Wolf et al., 2012). The cell wall is a complex structure of intertwined and sometimes cross-linked polymers, comprising cellulose, xyloglucans, pectins and arabinogalactans, which resists the internal turgor pressure. Therefore, cell wall elasticity has to be regulated to permit cellular growth (Cosgrove et al., 1984; Cosgrove, 1993, 2005; Wolf et al., 2012). This is achieved through selective loosening of cell wall polymer interactions, which allows cellulose microfibrils and associated matrix polysaccharides to displace relative to each other. As cellulose microfibrils are typically arranged in a non-random, parallel orientation, most cells expand along one principal axis. This process is easily observed in organs with one principal growth vector, for instance in hypocotyls or root tips. In both organs, hormones strongly influence cell elongation. Among these, auxin is most prominent, because it not only orchestrates developmental programs, but also conveys environmental inputs to trigger adaptive responses such as tropisms (Sanchez-Rodriguez et al., 2010; Depuydt and Hardtke, 2011). It is generally assumed that auxin promotes cell elongation by inducing the expression of cell wall remodeling factors (Sanchez-Rodriguez et al., 2010; Wolf et al., 2012). These include expansins, which are considered facilitators of cell wall loosening by physically opening up the fiber network, thereby facilitating the access of other enzymes to their substrates (Cosgrove, 2005). Moreover, in parallel, auxin supposedly stimulates cell elongation via a non-genomic pathway, since it enhances cell elongation within minutes in classical assay systems, such as hypocotyls or coleoptiles. In these model organs, auxin treatment correlates with increased acidification of the apoplast, which presumably promotes cell elongation because central cell wall loosening factors and enzymes, e.g. expansins, polygalacturonases, endoglucanases and pectin methylesterases, work optimally under acidic conditions. Cell wall acidification, in turn, is thought to arise from auxin-induced activation of plasma membrane-localized proton pumps (PM-H\(^+\)-ATPases). This long-standing concept, named the Acid Growth Theory, was formulated in the 1970s (Rayle...
Whether the Acid Growth Theory is universally applicable to plant cell expansion remains controversial. For instance, even for the classic assay systems (coleoptiles or hypocotyls), some authors have concluded that cell wall acidification indeed stimulates growth, but that this is not an auxin-dependent effect (Kutschera and Schopfer, 1985b, a; Schopfer, 1989, 1993). Validation of the Acid Growth Theory is most difficult in roots, where analyses are complicated by the fact that as opposed to hypocotyls or coleoptiles, cell proliferation and cell expansion are deeply intertwined. Because both processes require variable threshold auxin activities, they are difficult to uncouple, which might account for the observation that auxin application generally inhibits, or at best only slightly promotes root growth (Moloney et al., 1981b; Evans et al., 1994). Likewise, *Arabidopsis thaliana* (Arabidopsis) roots are shorter upon both a genetically imposed strong decrease as well as a strong increase in auxin production (Chen et al., 2014). Moreover, reduced mature cell length is typically accompanied by reduced meristem size and *vice versa* (Moubayidin et al., 2010; Scacchi et al., 2010; Rodriguez-Villalon et al., 2015), which makes it difficult to distinguish whether observed phenotypes are primarily caused by altered cell elongation or cell proliferation and/or differentiation. Thus, a paucity of clear-cut conditions and pertinent genetic material has prevented conclusive analyses of the Acid Growth Theory in Arabidopsis roots.

In this study, we took advantage of recently isolated mutants of the model grass species *Brachypodium distachyon* (Brachypodium) in the *TAA1-RELATED2-LIKE* (*TAR2L*) and *ETHYLENE INSENSITIVE2-LIKE 1* (*EIN2L1*) genes. *TAR2L* encodes an enzyme of the TRYPTOPHAN AMINOTRANSFERASE OF ARABIDOPSIS1 (TAA1) and TAA1-RELATED (TAR) family of proteins, which catalyze conversion of tryptophan to indole-3-pyruvic acid (IPA) in the two-step auxin biosynthesis pathway (Stepanova et al., 2011; Won et al., 2011). IPA is subsequently converted to indole-3-acetic acid (IAA), the major active auxin, by the YUCCA cytochrome P450 enzymes. Two TAR homologs exist in Brachypodium, with *TAR2L* dominating in the seminal root elongation zone, where cell differentiation and elongation occur (Pacheco-Villalobos et al., 2013). Bd-*EIN2L1* is a homolog of Arabidopsis *EIN2*, an essential positive regulator of ethylene
signaling (Alonso et al., 1999; Qiao et al., 2012). Both hypomorphic Brachypodium

\textit{tar2}\textsuperscript{hypo} and \textit{ein2l1}\textsuperscript{hypo} mutants display different degrees of elevated IAA levels in the

seminal root elongation zone as well as dramatically enhanced cell elongation. This

initially counterintuitive phenotype could be explained by the observation that the

regulatory logic of the two-step auxin biosynthetic pathway is different in Brachypodium

as compared to Arabidopsis: whereas ethylene positively regulates both steps in

Arabidopsis, it negatively regulates the second, rate-limiting step in Brachypodium

(Pacheco-Villalobos et al., 2013). The pathway intermediate IPA is metabolically linked

to ethylene biosynthesis through the VAS1 enzyme, which catalyzes the formation of

tryptophan from IPA using hydrophobic amino acids, mostly L-methionine, as amino

group donor (Zheng et al., 2013a). Because the size of the L-methionine pool limits the

biosynthesis of the rate-limiting ethylene precursor 1-aminocyclopropane-1-carboxylate,

not only IPA, but also ethylene biosynthesis are increased in Arabidopsis \textit{vas1} mutants.

By analogy, the Brachypodium \textit{tar2}\textsuperscript{hypo} mutation apparently creates a situation where

reduced TAR activity leads to reduced IPA production and thereby also reduced

ethylene production, while ethylene signaling is directly dampened in \textit{ein2l1}\textsuperscript{hypo} mutants.

Thus, in both hypomorphic mutants, \textit{YUCCA} genes become de-repressed to different

degrees and eventually more IPA is converted to IAA than in wild type (Pacheco-

Villalobos et al., 2013). Because a unique feature of both Brachypodium \textit{tar2}\textsuperscript{hypo} and

\textit{ein2l1}\textsuperscript{hypo} mutants is that their root meristem size and activity are not affected, their

longer seminal roots are entirely explained by the increased mature cell length

(Pacheco-Villalobos et al., 2013). Thus, cell proliferation and cell expansion are

uncoupled in seminal roots of Brachypodium \textit{tar2}\textsuperscript{hypo} and \textit{ein2l1}\textsuperscript{hypo} mutants, which both

display elevated auxin levels in conjunction with greatly exaggerated cell elongation.

Therefore, they offer an unprecedented opportunity to monitor the consequences of

high steady state auxin levels in a monocotyledon root type.
Results

**Metabolic analysis confirms higher auxin levels in tar2\textsuperscript{hypo} root segments despite reduced tryptophan aminotransferase activity**

The strongly enhanced mature root cell length in Brachypodium tar2\textsuperscript{hypo} mutants (Figure 1A) (Pacheco-Villalobos et al., 2013) is apparently cell autonomous, because in regenerating excised tar2\textsuperscript{hypo} root tips (Supplemental Figure 1A), newly formed cells are still longer than in its wild type background, Bd21-0 (Figure 1B). The same is true for ein2\textsuperscript{hypo} compared to its wild type background Bd21-3 (Supplemental Figure 1B). In our subsequent analyses, we primarily concentrated on the tar2\textsuperscript{hypo} mutant because of the relatively strong phenotype of tar2\textsuperscript{hypo} seminal roots. For a more complete analysis of auxin metabolism in tar2\textsuperscript{hypo}, we measured the tryptophan aminotransferase background activity (Figure 1C) as well as total (Figure 1D) activity, in Bd21-0 and tar2\textsuperscript{hypo} seminal roots, which revealed a ca. 30% reduction in specific activity in the mutant (Figure 1D). Full-scale analysis of auxin biosynthesis intermediates in 1 cm segments from the root elongation zone of 4-day-old seedlings (Supplemental Figure 1C) with comparable fresh weight in Bd21-0 and tar2\textsuperscript{hypo} (Supplemental Figure 1D) produced a matching metabolic profile; that is, tryptophan levels were slightly increased while IPA levels were substantially decreased in the tar2\textsuperscript{hypo} mutant (Figure 1E). Concurrently, the abundance of a few (inactive) auxin conjugates was shifted, while as previously observed (Pacheco-Villalobos et al., 2013), the level of free auxin (IAA) was significantly increased. Also consistent with previous findings (Zheng et al., 2013b), down-regulation of a Brachypodium VAS1-LIKE gene (VAS1L) by RNA interference suppressed the tar2\textsuperscript{hypo} phenotype genetically (Supplemental Figure 1E and F). In summary, these observations confirm our previous finding that down-regulation of TAR2L results in increased rather than decreased cellular auxin levels and causes strongly enhanced root cell elongation (Pacheco-Villalobos et al., 2013).

**High auxin steady state is associated with remarkable transcriptional homeostasis of the auxin signaling network**

The tar2\textsuperscript{hypo} mutant offers a unique opportunity to survey a steady-state high auxin concentration transcriptome that is associated with enhanced cell elongation. To this
end, we performed mRNA sequencing (RNAseq) of 1 cm root elongation zone segments, grown and harvested in parallel with those used for the metabolic analysis. Complementary to this experiment, we also performed RNAseq on equivalent segments from wild-type plants that had been transferred onto medium containing L-kynurenine for two days. Mild concentrations of this tryptophan aminotransferase inhibitor induce
higher auxin levels and enhanced cell elongation in wild type roots, thus mimicking the tar2l hypop phenotype (Pacheco-Villalobos et al., 2013). The reads from the Bd21-0, tar2l hypop and L-kynurenine-treated Bd21-0 samples mapped onto more than 27,000 mRNA transcripts out of the 31,679 nuclear genes annotated in the Brachypodium reference genome sequence (version 2.1) (Supplemental Data Set 1), with a pairwise overlap between samples of more than 97%. Compared to wild type, 957 and 2,657 genes were differentially expressed in tar2l hypop and L-kynurenine-treated roots, respectively (q value < 0.01, fold change > 2X) (Figure 2A) (Supplemental Data Set 2).

Figure 2. Differential gene expression in root segments as determined by RNA sequencing. (A) Venn diagrams illustrating overlaps between the gene sets that were differentially expressed in root segments of Bd21-0 versus tar2l hypop, Bd21-3 versus ein2l hypop, and mock-treated versus L-kynurenine-treated Bd21-0. (B) Cluster analysis and (C) principal component (PC) analysis of the different RNA sequencing samples.
and 3). The higher number of differentially expressed genes in the L-kynurenine-treated samples is consistent with an organ-wide systemic effect of the treatment that includes transcriptome remodeling towards a new steady state. The overlap between the two sets was 344 genes, which represents ~4-fold enrichment over neutral expectation ($p < 0.0001$, $\chi^2$ test). A similar RNAseq experiment was carried out with root segments from Bd21-3 and ein2-1hypo seedlings. Again, over 27,000 transcripts were detected and 356 genes were differentially expressed (Supplemental Data Set 1 and 4). Overlap with the tar2-1hypo and L-kynurenine-treated sets was 140 and 112, respectively, which again represented high enrichment (~12-fold and ~3.5-fold, respectively) over neutral expectation ($p < 0.0001$, $\chi^2$ test). In summary, the RNAseq profiles indicated high overlap between the mutants and the L-kynurenine condition, with the extent of differentially expressed genes correlating with phenotype strength. Despite the similarities between their transcriptome profiles, the samples were clearly grouped apart. Both mutants were more similar to their wild type backgrounds than to each other, and the L-kynurenine-treated samples were most distant to all others (Figure 2B). A principal component analysis confirmed that parental background was the dominant factor in the grouping of samples (Figure 2C).

Analysis of the annotations of the differentially expressed genes revealed a rather low occurrence of genes involved in auxin or other hormone signaling pathways (Figure 3A). Significant differential expression was observed for six out of 26 annotated auxin response factors, seven out of 32 annotated AUX/IAA genes, and one out of five annotated auxin receptor genes (Figure 3B) (Supplemental Data Set 5). However, the expression changes were moderate throughout. Likewise, mostly small effects were observed for the few differentially expressed genes involved in polar auxin transport, which included two auxin influx facilitators and three auxin efflux carriers. Overall, the data indicate that the transcriptional steady state of the auxin-signaling network is well buffered with respect to variation in auxin levels. Interestingly however, some primary auxin target genes of the SMALL AUXIN UP-REGULATED (SAUR) category were differentially expressed (five out of 42 annotated genes) and mostly upregulated (4 out of the 5) (Figure 3C). SAUR genes are classic auxin signaling output genes, and it has been suggested that SAUR proteins antagonize post-translational inhibition of PM-H$^+$-
ATPases (Spartz et al., 2014). Finally, with the exception of two genes that were substantially down-regulated (Figure 3D), Brachypodium genes that encode PM-H\(^+\)-ATPases displayed no differential expression.
High auxin steady state is associated with transcriptional changes in cell wall remodeling factors

The majority of significantly enriched terms that stood out in a word cloud made from annotation of differentially expressed genes was related to the cell wall and associated processes (Figure 3A). Yet, only a small proportion of genes encoding cell wall remodeling proteins displayed differential expression, and these were restricted to a few groups. For example, while six out of fifteen annotated xyloglucan endotransglycosylases/hydrolases were up- or down-regulated at roughly equal measure (Figure 3E), no differential expression was observed among cellulose synthase genes. The most prominent differentially expressed cell wall modulators were expansins (23 out of 54 annotated expansin genes), which were significantly enriched ($p = 0.0078$ for the overlap between all samples, $\chi^2$ test) and are also considered classic auxin target genes. The majority (18 out of 23) displayed comparatively strong up-regulation (Figure 3F). Finally, three genes encoding arabino-galactan peptides or proteins stood out because of their consistent up-regulation (Figure 3G). Verification of differential expression by qPCR was carried out for a selected set of genes of interest in independent RNA samples from $\text{tar2lhypo}$ and L-kynurenine-treated root segments, confirming the RNAseq results (Figure 3H). In summary, the transcriptomic data indicate that in the presence of higher cellular auxin levels, the auxin signaling network maintains remarkable homeostasis at the transcriptional level, while the bulk of expression changes are observed in cell wall remodeling genes, notably expansins.

High auxin steady state is associated with a specific change in glycosidic cell wall linkages

The robust changes in the arabino-galactan protein/peptide genes were of particular interest in light of our results from chemical cell wall analyses. To monitor the structural effect of altered expression in cell wall remodeling genes, we performed cell wall polysaccharide analysis of root segments from parallel samples that were grown and harvested concomitantly with the segments analyzed by RNAseq. The analysis of the glycosidic linkages occurring in the cell wall polymers detected only few significant differences between Bd21-0 and $\text{tar2lhypo}$, or Bd21-3 and $\text{ein2lhypo}$. However, both
profiles were consistent, with a specific significant decrease in 1,3-galactosyl residues in the mutants relative to their wild type backgrounds (Figure 4A & B). 1,3-linked galactose is found specifically in the glycosidic moiety of the arabinogalactan proteins (AGPs)
1,3-linked galactosyl residues represented ~10% of all linked sugar residues detected in both wild type backgrounds and were reduced by about two- to three-fold in ein2l1hyp and tar2hyp, respectively. Moreover, the analysis of the cell wall neutral sugars indicated similar relative abundance of the different monosaccharides in the mutants and wild types, with one notable exception, fucose, a minor cell wall sugar (Figure 4C & D). Relative fucose abundance was more than halved in tar2hyp and reduced by about one third in ein2l1hyp. Interestingly, just like β-1,3-linked galactose, fucose is found in AGPs (van Hengel and Roberts, 2002). Thus, the analyses point to a very specific effect of elevated cellular auxin levels on arabinogalactan complexity or abundance in Brachypodium. To confirm this observation with an alternative technique, we performed in situ Yariv staining on Bd21-0 and tar2hyp roots. The Yariv reagent is known to specifically detect β-1,3-galactan (Yariv et al., 1967; Kitazawa et al., 2013). Staining was considerably reduced in the root elongation zone of tar2hyp, thereby corroborating the chemical cell wall analyses (Figure 4E). Moreover, we probed transverse sections in the root elongation zone with antibodies directed against de-methylesterified pectin (2F4 antibody; Figure 4F), methylesterified pectin (JIM7 antibody; Figure 4G) and arabinogalactan side chains (LM2 antibody; Figure 4H). None of these stainings showed a marked difference in epitope abundance or distribution, except that in general, the mean fluorescence signal of the LM2 antibody was reduced. This might hint to lower AGP abundance, however, it is unclear to what degree the antibody stainings are quantitative. Importantly, unlike the Yariv reagent, the LM2 antibody does not recognize the β-1,3-galactan linkages in the arabinogalactan backbone, but rather an epitope that comprises β-linked glucuronic acid, which is found at the side chain termini (Smallwood et al., 1996; Knoch et al., 2014). Therefore, collectively, the results point to reduced AGP complexity.

*High auxin steady state is not associated with markedly increased proton excretion*

Our transcriptomic and cell wall analyses indicate that elevated cellular auxin levels in Brachypodium roots are indeed associated with differential expression of cell wall remodeling genes and matching changes in cell wall composition. To determine
whether this also applies to the hallmark of the Acid Growth Theory, apoplastic acidification, we next investigated the capacity of Bd21-0 and tar2hypo roots to acidify the medium. To this end, we first visualized rhizosphere acidification by transferring...
seedlings onto medium supplemented with pH indicator. Acidification was readily detected within 4 h, but was not apparently stronger for tar2\textsuperscript{hypo} roots as compared to Bd21-0 roots (Figure 5A & B). Likewise, in a quantitative assay with liquid medium, acidification could be readily followed over time, however, no difference in proton pumping activity of root tips could be detected between Bd21-0 and tar2\textsuperscript{hypo} (Figure 5C). Finally, we measured apoplastic acidification more directly at the root surface by using fiber-optic pH microfiber sensors. To this end, five roughly equidistant measuring points from the root tip through the elongation zone were monitored along individual roots. As could be expected, these measurements revealed a gradient of increasingly acidic pH from the root tip to the differentiated cells. However, this gradient was less rather than more pronounced in tar2\textsuperscript{hypo} as compared to Bd21-0 (Figure 5D). Likewise, in general, reduced rather than increased acidification was observed in ein2\textsuperscript{l1 hypo} roots as compared to their Bd21-3 wild type background (Supplemental Figure 2A). Finally, we monitored the phosphorylation state of PM-H\textsuperscript{+}-ATPases in root segments. Phosphorylation of the penultimate amino acid within the autoinhibitory C-terminal domain of PM-H\textsuperscript{+}-ATPase and subsequent binding of 14-3-3 proteins is the major mechanism of enzyme activation (Palmgren et al., 1991; Portillo et al., 1991; Speth et al., 2010). We therefore monitored both the capability of 14-3-3 proteins to associate with PM-H\textsuperscript{+}-ATPase in microsomal membranes of root segments (14-3-3 overlay), reflecting its phosphorylation level, and the amount of PM-H\textsuperscript{+}-ATPase (Ottmann et al., 2007; Speth et al., 2010). Interestingly, phosphorylation-dependent binding of 14-3-3 proteins to the PM-H\textsuperscript{+}-ATPase was reduced, rather than increased, in microsomal preparations from tar2\textsuperscript{hypo} root segments as compared to Bd21-0 (Figure 5E). This is in striking contrast to the effect of auxin on PM-H\textsuperscript{+}-ATPase phosphorylation on hypocotyl elongation in Arabidopsis (Takahashi et al., 2012). Collectively, these experiments suggest that higher cellular auxin levels in Brachypodium roots are not associated with proton pump activation or markedly elevated proton excretion at the mesoscopic level.

*Forced apoplastic acidification inhibits root cell elongation*

Next, to conversely determine whether acidity affects Brachypodium root cell elongation, we monitored the response of the root to externally imposed pH changes. In Arabidopsis, strong acidity eventually impairs overall root growth by inhibiting
meristematic activity (Gujas et al., 2012), and the same applies to Brachypodium (see below). Reduced meristematic activity could alter mature cell length because it shifts the balance between proliferation and differentiation (Moubayidin et al., 2010; Scacchi et al., 2010). Therefore, we chose to examine mature cortex cell length after transfer of seedlings from standard medium (pH 5.7) to mildly more acidic conditions (pH 5.2), which nevertheless represent a ca. 3-fold increase in H⁺ concentration. Only cells formed after the transfer were scored. In these experiments, no significant length difference was observed between cells formed on either pH (Figure 6A) and overall root growth was not affected (Supplemental Figure 2B). At the same time, fiber-optic pH sensor measurements along the root surface performed in parallel revealed converging pH gradients under the two conditions (Figure 6B), to ca. pH 4.9 in the root elongation zone. Medium acidification by Brachypodium root tips to pH 4.8 to 4.9 was observed...
repeatedly and appears to represent a lower limit in tissue culture. Therefore, we challenged roots with pH 3.7, a respective ca. 10-fold increase in acidity. Surprisingly, while overall root growth was substantially reduced at this acidic pH (Figure 6C), which was entirely attributable to reduced meristematic activity. Mature cell length was again not affected (Figure 6D). Likewise, even shortly after transfer of root tips into acidic medium, at best a small and transient significant positive effect on cell elongation could be observed (Supplemental Figure 2C). The same applies to similar experiments where root tips were transferred into medium that contained fusicoccin, a proton pump stimulant (Supplemental Figure 2D). In summary, these results suggest that cell elongation in Brachypodium roots is robustly buffered against external pH fluctuations.

To further explore the relation between apoplastic acidification and cell elongation, we turned to a model system that allows more direct manipulations, Arabidopsis. Similar to Brachypodium, mature cortex cell length was scarcely sensitive to pH variations in the medium (Figure 6E), meaning that again reduced overall root growth on acidic medium (Supplemental Figure 2E) can be largely attributed to decreased meristematic activity, as previously observed (Gujas et al., 2012). Thus, apparently root cell elongation is also robustly buffered against external pH changes in Arabidopsis roots. To override this buffering effect, we sought to uncouple proton pump activity from homeostatic inputs and stimulate it at will. To this end, we again applied fusicoccin, which at low concentration again resulted at most in a small significant stimulation of cell elongation, while higher concentration clearly reduced cell elongation (Figure 6F). In a more direct, genetic approach, we investigated wild type seedlings that carried an inducible transgene for conditional expression of the Arabidopsis PM-H⁺-ATPase AHA2 devoid of its C-terminus (AHA2δ). This 95 amino acid deletion removes the auto-inhibitory domain of the protein, which is therefore turned into a hyperactive proton pump that is uncoupled from regulatory inputs (Regenberg et al., 1995; Axelsen et al., 1999). Similar to high fusicoccin concentrations, strong induction of the AHA2δ construct resulted in the cessation of meristem activity, massive root cell swelling and eventual rupture of the root tissues. By contrast, at lower induction levels, which maintained root growth, cell elongation was strongly reduced (Figure 6G). In these conditions, some cellular swelling was observed, consistent with an increased vacuole
size (Figure 6H), yet this vacuolar size increase by itself was apparently not sufficient to
drive significant cell elongation. Finally, in all conditions, both fusococcin exposure and
AHA2⁹⁵ induction resulted in reduced meristematic activity and thus reduced meristem
size (Supplemental Figure 2F and G). Yet even when strong vacuolar swelling was
induced, we did not observe longer cells.
Discussion

The importance of auxin in plant development cannot be overstated. Auxin impinges on a large variety of physiological and morphological processes, for which both absolute and relative auxin levels can be determinants. This is for instance illustrated in root development, where auxin biosynthesis, polar transport and signaling are required for proper morphogenesis, growth and integration of environmental signals (Hardtke and Berleth, 1998; Sabatini et al., 1999; Zhao, 2014; Adamowski and Friml, 2015). A wealth of genetic and physiological data underpins the role of auxin in root development, yet root responses to systemically applied external auxin have been difficult to interpret. While picomolar levels of auxin sometimes stimulate root growth, physiological, nanomolar concentrations in general suppress root growth (Sutcliffe and Sexton, 1969; Evans et al., 1994; Overvoorde et al., 2010). Likewise, genetically increased excess cellular auxin production through ectopic over-expression of YUCCA enzymes inhibits rather than enhances root growth (Chen et al., 2014). These results indicate that in the absence of correct tissue context, increased auxin levels fail to reveal the central role of auxin in root growth, possibly because crucial auxin gradients are overridden (Benjamins and Scheres, 2008). In summary, pertinent auxin biosynthesis, transport or signaling mutants and transgenic lines, mostly in Arabidopsis, do not display substantially enhanced root growth. Therefore, the Brachypodium tar2\textsuperscript{hypo} and ein2\textsuperscript{hypo} mutants represent a so far very unusual situation, because here locally increased auxin levels are associated with a specific and strong stimulatory effect on root cell elongation while meristematic activity and meristem size are not affected (Pacheco-Villalobos et al., 2013). This observation also contradicts the sometimes-voiced argument that root cell elongation is typically maximal and therefore cannot be stimulated further by hormone action.

A transcriptome associated with high auxin steady state

The remarkable phenotypic specificity of both Brachypodium mutants with respect to root cell elongation offered us the unique opportunity to survey a transcriptome that is associated with a high auxin steady state. Auxin-regulated genes have so far been mainly identified through their response to external auxin application. This approach has
been tremendously successful in identifying the principal auxin target genes and the 
auto-regulatory feedback in the auxin signaling networks. Most prominently, they 
include AUX/IAA genes, which encode repressors of auxin signaling and respond 
rapidly and strongly to auxin application. Compared to these classic auxin-responsive 
transcriptomes, components of the auxin signaling network are rare among the 
differentially expressed genes in our RNAseq analyses. Even when significant, their 
expression fold changes are very moderate throughout, typically smaller than 1.5. 
Overall, the differentially expressed auxin signaling genes are down-regulated in our 
samples, which could indicate a compensatory mechanism in response to higher 
cellular auxin levels. Thus, at the transcriptional level, the auxin signaling network 
maintains a remarkably buffered homeostasis. By contrast, a number of classic auxin 
target genes that are considered physiologically relevant immediate outputs of auxin 
action are differentially expressed at higher levels in the mutants and are up-regulated 
overall. Most notably, these include genes related to cell wall remodeling, among which 
substantial differential expression of genes encoding expansins is most robust. In 
summary, these observations suggest that our mutant transcriptomes could be 
indicative of physiologically relevant auxin targets.

Between the five types of transcriptomes, we not only observed increasing 
differential expression as a function of phenotypic strength, or, in the case of roots 
grown on L-kynurenine, systemic action, but also in relation to parental background. 
However, although all samples were harvested in parallel, the ein2l1\textsuperscript{hypo} and Bd21-3 
root segments were processed at a different time from the other samples, and their 
RNAseq was performed in a separate instrument run. Thus, it is possible that the 
ein2l1\textsuperscript{hypo} versus \textit{tar2}\textsuperscript{hypo} transcriptome comparison is to some degree not only 
constrained by parental background, but also by batch effect. Although this limits the 
validity of any derived analyses, it is noteworthy that across all possible cross-
comparisons, the by far most robust differential expression was observed for a gene 
that encodes an arabinogalactan peptide.

\textit{A specific effect of high steady state auxin levels on arabinogalactan complexity}

AGPs are a group of highly diverse cell surface glycoproteins (Seifert and Roberts, 
2007; Ellis et al., 2010). Their protein backbone is characterized by di-peptide motifs
that comprise hydroxyproline residues, which serve as attachment points for β-1,3-linked galactose oligosaccharides. These galactans can themselves serve as secondary branch points for additional side chains, which can contain a variety of other sugars, such as arabinose or fucose. The exact roles of AGPs in plant development remain somewhat unclear, in part because of their structural variety and the resultant fuzziness of analyses, but they have been implicated in various growth-related processes (Seifert and Roberts, 2007; Ellis et al., 2010). The most clear-cut evidence for a role in root development so far comes from genetic analyses of Arabidopsis plants with altered expression of enzymes that have an experimentally proven role in the biosynthesis or degradation of arabinogalactan side chains (Knoch et al., 2014). An interesting finding from this small set of studies is that while arabinogalactans appear to be generally required for cell elongation, in mutants or transgenic lines with mildly reduced arabinogalactan complexity, root elongation is substantially stimulated (van Hengel and Roberts, 2002; Eudes et al., 2008; Knoch et al., 2013). However, whether this is mainly due to enhanced cell elongation has not been reported.

The published genetic data on the role of arabinogalactans in Arabidopsis root development match our observations for Brachypodium. In our root segments, we did not observe any changes in the expression of Brachypodium homologs of proven arabinogalactan metabolism enzymes. Yet, we observed a clear reduction in β-1,3 galactan levels in biochemical and in situ analyses of our mutants. Overall, the data suggest reduced complexity, and possibly also abundance of arabinogalactans. It is conceivable that these observations could be linked to changes in AGP expression in response to higher steady state auxin levels. AGP protein backbones, which are typically ca. 100 amino acids long, are secreted and attached to the plasma membrane via a glycosylphosphatidylinositol membrane anchor that is added during their processing. The same applies to the much shorter arabinogalactan peptides, which are only 10 to 12 amino acids long (Schultz et al., 2004). Of the three differentially expressed arabinogalactan backbone genes in our dataset, the one encoding an 11 amino acid arabinogalactan peptide is most dynamically and robustly over-expressed (4-7X) in the high cellular auxin situation across all comparisons. Perhaps this shift in arabinogalactan protein backbone length distribution to a higher proportion of short
backbones could lead to a looser, less complex AGP network. Future approaches, for instance transgenic overexpression, could be used to address this notion directly.

Root cell elongation is robustly buffered against external pH fluctuation

The Acid Growth Theory of plant cell elongation has been formulated with respect to the elongation of shoot organs, with an experimental focus on coleoptiles and hypocotyls. To what degree it is pertinent for root cell elongation has been controversial from the beginning because of early observations that auxin application to intact roots generally inhibits growth or has no effect (Sutcliffe and Sexton, 1969). At best, growth stimulation could be observed with very low auxin concentrations or in treatments of auxin-depleted roots (Edwards and Scott, 1977; Pilet et al., 1979; Evans et al., 1994). However, in all cases, the size of the effect was small, and it was not reported whether the effect was due to altered root meristem activity or cell elongation. In accordance with these results, the impact of auxin on proton excretion also did not match observations in shoot organs and was generally variable. For example, proton efflux upon treatment of maize roots was reported for nanomolar concentrations of auxin, while proton uptake was observed with micromolar concentrations, with the caveat that these roots had been pre-treated with ethylene biosynthesis inhibitors (Mulkey et al., 1982). Others suggested that growing parts of barley (*Hordeum vulgare*) roots take up protons, while non-growing parts secrete them (Weisenseel et al., 1979). Finally, a recent study that monitored apoplastic pH using a fluorescent molecular marker *in planta* found that auxin treatment has little effect on pH in the meristem tip but leads to alkalization rather than acidification in the root cell elongation zone (Gjetting et al., 2012), corroborating similar earlier claims (Evans et al., 1980; Moloney et al., 1981a; Luthen and Bottger, 1993). Eventually, for maize roots, it was concluded that the pH growth curve exhibits a broad optimum ranging from pH 4.5 to 9, that any acid-induced growth is of very short duration, that the low sensitivity of root growth to external pH is independent of both pump activity and buffering capacity of the bathing solution, and that neither incubation in acidic buffer nor stimulation of the proton pump reverts auxin-induced root growth inhibition (Luthen and Bottger, 1993).

Our observations largely second these conclusions for *Brachypodium* as well as *Arabidopsis*. Although it is evident from our assays that the root tips acidify the
rhizosphere as could be expected, we did not detect enhanced proton excretion at the surface of mutant roots. Thus, although a few SAUR genes are up-regulated under the high auxin conditions in our transcriptomes and could possibly stimulate proton pumps indirectly, similar to Arabidopsis SAUR19 (Spartz et al., 2014), this apparently does not translate into a detectable increase in proton excretion at the mesoscopic level. Likewise, the two proton pump genes that are down-regulated under high auxin conditions should have little impact on overall proton pump abundance, because they are only weakly expressed. The by far preponderant proton pump gene that we detected (Bradi5g24690) was expressed very robustly across all conditions tested at levels over ten times higher than the other nine proton pump genes combined. Indeed, our protein gel blot analysis confirmed that PM-H+-ATPase abundance in tar2hypo was comparable to wild type or at best mildly reduced. Moreover, given that proton pump activity is mostly regulated through post-translational modifications, one would not expect that their mild transcriptional regulation would play a major role.

Based on our data, we cannot exclude the possibility that elevated proton pumping is induced by higher cellular auxin content, but it does not propagate beyond the immediate vicinity of the cell surface because of the concomitantly increasing membrane potential. Such proton pumping could therefore not be detected by our methods. Although our observation that PM-H+-ATPase phosphorylation is decreased rather than increased in tar2hypo argues against this scenario, it is important to note that the interpretation of our findings is constrained by the lack of single cell resolution of our observations and morphological features, such as the shorter root hairs in tar2hypo mutants (Pacheco-Villalobos et al., 2013). Perhaps very local and transient acidification is sufficient to trigger cell elongation, which would make the observation that cell elongation in both Brachypodium and Arabidopsis roots is robustly buffered against external pH fluctuation, including imposed acidity, even more remarkable. Consistently, throughout our experiments, excess acidity eventually suppressed root growth by impairing meristematic activity rather than cell expansion. This finding suggests that compensatory mechanisms act to keep apoplastic root cell pH optimal for cell elongation. Our finding that forced apoplastic acidification, for instance through
induction of a constitutively active proton pump, strongly impairs root cell elongation
underlines this notion.

In summary, our data suggest that elevated steady state auxin levels in Brachypodium seminal roots are associated with specific transcriptomic and cell wall changes. While some of those changes match expectations, e.g., enhanced expression of expansin genes, others come as a surprise, notably the specific effect on arabinogalactans. Whether all of the observed changes are direct effects of auxin or emerge indirectly, for example from hormonal crosstalk, notably with ethylene, remains to be determined. At this point, our data provide a phenotypic reference framework for future investigations that might also clarify to what degree our observations are specific for Brachypodium. Robust buffering of root cell elongation against pH fluctuation would surely make sense in the universal biological context of root growth. Unlike the shoot system, the root system is in close contact with a solid phase environment, the soil, which imposes its pH. Roots can modify the rhizosphere pH by proton pumping to increase the solubility of essential nutrients and promote their uptake, which has an optimum in the range of pH 5.0–6.5. Given the variability of soil pH values and their seasonal fluctuation, it appears advantageous for the plant if root growth capacity is not dictated by the soil environment pH. From this perspective, it would make sense that the interplay between auxin, proton pump activation and expansin action at the heart of the Auxin Growth Theory is possibly more flexible in the root than in the shoot.
Methods

Plant materials and growth conditions

The Brachypodium distachyon mutants \(\text{tar2}^{\text{hypo}}\) and \(\text{ein2l1}^{\text{hypo}}\) and their respective wild type backgrounds Bd21-0 and Bd21-3 have been described before (Pacheco-Villalobos et al., 2013). Unless indicated otherwise, analyses were performed on tissue culture-grown 4-day-old seedlings raised under previously described conditions (continuous light, 100–120 \(\mu\)E intensity, 22\(^\circ\)C, Philips F17T8/TL741 fluorescent light bulbs) (Pacheco-Villalobos et al., 2013). Solid media were prepared using Phytagel (Sigma) and Murashige and Skoog (MS) salts (Sigma). Care was taken to place the 10 cm square Petri dishes at a slight angle from the vertical to assure seminal root growth along the agar surfaces. Roots that had grown into the plate were excluded from analysis. For metabolic profiling, chemical cell wall analysis and RNAseq, parallel grown ~1 cm seminal root pieces harvested from 2-3 mm above the root tip were sampled (see Supplemental Figure 1D). To generate VAS1L RNAi knock down lines, a DNA fragment of 422 bp containing the VAS1L 3'-UTR was amplified using the oligonucleotides attB1-BdVAS1L-F 5'- GGG GAC AAG TTT GTA CAA AAA AGC AGG CTT GGT ACA G TA ACA GCC CAT C -3' and attB2-BdVAS1L-R 5'- GGG GAC CAC TTT GTA C AA GAA AGC TGG GTG GAA GTG GCA GTT CTG TCA G-3', and cloned into pDONR207 (Life Technologies). This 3'-UTR specific DNA fragment was then cloned into the pANDA RNAi vector (Miki and Shimamoto, 2004). Transformation of the pANDA-BdVAS1Li plasmid into \(\text{tar2}^{\text{hypo}}\) embryo-derived callus was performed as described (Pacheco-Villalobos et al., 2013). Arabidopsis experiments were performed with the standard Col-0 accession under the growth conditions described above. For dexamethasone-inducible expression of AHA2 (AT4G30190) devoid of its autoinhibitory C-terminal domain (AHA2\(^{955}\)) the corresponding cDNA was amplified by PCR and cloned into pTA7002 (Aoyama and Chua, 1997) via Xhol and SpeI restriction enzyme sites. Transgenic lines were obtained after transformation of Col-0 plants using standard procedures (Clough and Bent, 1998).

Tryptophan aminotransferase activity assays
Seminal roots were ground in liquid nitrogen with a TissueLyser II (Qiagen). Root tissue was homogenized in a pre-cooled mortar on ice with one volume of extraction buffer (100 mM 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid buffer pH 7.2, 250 mM sorbitol, 5 mM β-mercaptoethanol, 0.5% [v/v] Triton X-100, and 0.1% [w/v] phenylmethylsulfonyl fluoride). The protein extract was centrifuged at 16,000 g for 30 min at 4°C. The supernatant was used for determination of tryptophan aminotransferase activity with the Salkowski reagent as described (Szkop et al., 2012). Briefly, the reactions were carried out in 100 mM phosphate buffer pH 8.0, 10 mM L-tryptophan, 10 µM pyridoxal phosphate and 50 µg of soluble proteins. The mixture was pre-incubated for 3 min at 35°C. The transamination reactions were initiated by the addition of 3 mM 2-oxoglutarate and incubated for 15 min at 35°C. To estimate the basal tryptophan aminotransferase activity of the crude extracts, control samples without 2-oxoglutarate were also assayed. To terminate the reactions, 1 ml of Salkowski reagent (10 mM FeCl$_3$, 35% [v/v] H$_2$SO$_4$) was added and the samples were incubated in the dark for 10 min at room temperature. The absorbance at 530 nm of four replicates was measured.

**Detection of proton pump phosphorylation status**

To determine the abundance of activated PM-H$^+$-ATPase, microsomes were prepared from root segments and analyzed with the overlay assay as described (Ottmann et al., 2007), except that RGS-(His)$_6$-tagged 14-3-3 was applied. Bound 14-3-3 was visualized by means of an antibody raised against the RGS-His$_6$ epitope (20 µg/ml) (Qiagen, Cat No./ID 34610).

**Auxin metabolite profiling**

For full-scale profile of auxin metabolites, four independent replicate samples of pooled 1 cm root segments were analyzed. Upon harvest, samples were immediately frozen in liquid nitrogen and stored at -80°C until they were analyzed as described (Novak et al., 2012).

**RT-PCR**

To monitor the expression of $VAS1L$, full length transcript by RT-PCR, the full length transcript was amplified using oligonucleotides 5’-ATG AGC AGC TTT GCC AAG CT-3’
and 5'-GGA AGT GGC AGT TCT GTC AG-3'. **UBIQUITIN CONJUGATING ENZYME 18** (see above) was used as a control.

**RNA sequencing and data analysis**

For RNAseq, total RNA was extracted from 8-12 pooled root segments using RNA extraction kits (Qiagen). cDNA libraries for sequencing were then prepared with the TruSeq Stranded mRNA Library Prep Kit (Illumina) using 1 μg of RNA starting material. Sequencing was performed on HiSeq 2500 instruments (Illumina) to yield 100 bp reads. The Bd21-0, *tar2*<sup>hypo</sup>, mock-treated Bd21-0 and L-kynurenine-treated Bd21-0 samples were prepared and run in parallel, multiplexed in the same sequencing lane. The Bd21-3 and *ein2*<sup>l1hypo</sup> samples, although grown and harvested in parallel, were processed in a separate run in the same manner. The 100 bp single reads were then mapped onto the Brachypodium primary transcripts (version 2.1, http://phytozome.jgi.doe.gov/pz/portal.html) using kallisto software (version 0.42.1, http://pachterlab.github.io/kallisto/) (Bray et al., 2016) with default settings (100 bootstrap samples). Subsequent differential expression analysis was performed using sleuth software (version 0.27.3, http://pachterlab.github.io/sleuth/), again with default settings. The word cloud was produced using the wordle online tool (www.wordle.net).

**qPCR**

Quantitative PCR was performed on three biological replicates with a Stratagene MxPro 3005P Real-Time PCR System (Stratagene), using SYBR® Green to monitor DNA synthesis. Relative gene expression levels were calculated as described in Pacheco-Villalobos et al., 2013. The following oligonucleotides were used: reference gene **UBIQUITIN CONJUGATING ENZYME 18** (**Bradi4g00660**), 5'-GGA GGC ACC TCA GGT CAT TT-3' and 5'-ATA GCG GTC ATT GTC TTG CG-3', **EXPANSIN** (**Bradi1g74710**), 5'-GTC CTC TAC CAA CAG GTG AAG-3' and 5'-AGT TCC TGG ACA TCT GGA TC-3'; **EXPANSIN** (**Bradi3g50740**), 5'-CGC GTG CTA TCA GGT TAA ATG C-3' and 5'-TCT GTC CTC TAC CAA CAG GTG AAG-3' and 5'-AGT TCC TGG ACA TCT GGA TC-3'; **AG-peptide** (**Bradi2g31980**), 5'-AGT ACC CCC TTC GGT TTC GT-3' and 5'-TGG TCG ATG GAC GAT GCG TC-3'; **GH3 enzyme** (**Bradi2g20740**), 5'-ACC ACT TAC TCC GGG CTG TA-3' and 5'-CGT GTA CTC CAC TAA AGA CG-3'; **PM-H+-ATPase** (**Bradi1g12117**), 5'-AGA TGG GAG GTA CTC CAC TAA AGA CG-3'; **PM-H+-ATPase** (**Bradi1g12117**), 5'-AGA TGG GAG GTA CTC CAC TAA AGA CG-3'.
GAA AGA GAG TC-3' and 5'-AAT GGC TAG CTG ATC ACC TG-3'; PM-$H^+$-ATPase
(Bradi3g18790), 5'-CCA GAG GAT GAA GAA CTA CAC G-3' and 5'-GAT CGT CAT GAT TGT GCC ATC G-3'; ACC synthase (Bradi1g10030), 5'-CCA CTG GCA TCA ATC GAT C-3'; ETRL2 (Bradi3g55730), 5'-GCA GAA AGC TTG TGC AGA TG-3' and 5'-GCA TGA CGG CGA TGT ATA TTG C-3'.

**Yariv staining**

For arabinogalactan staining with Yariv reagent, roots isolated from Bd21-0 and tar2lhypo seedlings were embedded in 6% agarose and longitudinally sectioned with a Leica-VT 1000S vibratome. Root sections were then incubated in a Yariv reagent (Biosupplies) solution (freshly prepared according to the manufacturer's instructions) for 5 min and directly examined under a Leica™ DM5500B compound microscope.

**Rhizosphere acidification assays**

Media acidification assays were performed as described (Gujas et al., 2012). To visualize rhizosphere acidification, 3-day-old Brachypodium seedlings were transferred to half-strength MS-agar plates supplemented with 0.15 mM bromocresol purple (Sigma) (sensitivity range pH 5.2 - 6.8). The plates were incubated in the same culture chamber and scanned after 4 and 24 hours. For liquid medium pH assays, 2-day-old Brachypodium seedlings were transferred on a sterile mesh attached to a tube containing 10 ml of non-buffered half-strength liquid MS medium and 0.15 mM bromocresol green (Sigma) (sensitivity range pH 3.8–5.4). Measurement of pH was performed in a time series at 4, 24, 48 and 240 hours. Three replicates consisting of eight plants per tube were measured. Negative controls (mock) without plants were measured in parallel.

**Root tip regeneration assays**

For root tip regeneration experiments, ~1 cm root segments from above the root tip were excised with a razor blade. The isolated root tips were then incubated on the same plates under the same conditions. *De novo* root tissue formation from the tips was monitored by scanning the plates after 1, 2, 3, and 4 days. Length quantification of newly formed cortex cell was performed by microscopy at 4 days after excision.
Antibody staining of Brachypodium root sections

Bd21-0 and tar2hyp plants were grown on half strength MS plates containing 1% sucrose and 0.9% agarose under long day conditions (16 h light / 8 h dark) at 22°C. 4-day-old roots were then sectioned with a vibratome (Leica VT1000 S). For immunolabeling of de-methylesterified pectin, freshly cut 100 µm cross-sections were first rinsed with 2F4 buffer (Tris-HCl 20 mM pH 8.2, CaCl2 0.5 mM, NaCl 150 mM) for 10 min. The samples were then incubated with 2F4 monoclonal antibody (PLANTPROBES) diluted 1:250 in 2F4 buffer with 5% skim milk powder (w/v) under gentle stirring for 1 hour. After washing three times with 2F4 buffer, the samples were incubated with secondary antibody (Goat anti-Mouse IgG (H+L), Alexa Fluor® 488 conjugate, ThermoFischer A-11001) diluted 1:1000 in 2F4 buffer with 5% skim milk powder (w/v) for 3 hours in the dark and then washed three times in 2F4 buffer. For immunolabeling of methylesterified homogalacturonan and arabinogalactan, respectively, the sections were incubated in 1X PBS buffer with 1% BSA and 0.05% Tween for 1 hour at room temperature, washed with 1X PBS and then incubated with JIM7 or LM2 (PLANTPROBES) diluted 1:25 in 1X PBS buffer with 1% BSA and 0.05% Tween for 1 hour at room temperature. The samples were washed three times with 1X PBS and incubated with secondary antibody (Donkey anti-rat Cy3, Jackson ImmunoResearch 712165150) diluted 1:500 in 1x PBS for 1 to 3 hours in the dark. The samples were washed three times with 1X PBS buffer after incubation. Z-stacks were acquired using a Zeiss LSM 510 Meta confocal microscope.

Analysis of Cell Wall Polysaccharides

Biological replicates consisted of two pools of root segments harvested from 300-400 seedlings per genotype. The biological material was freeze-dried overnight and ground to a fine powder using a Mixer Mill MM 400 (Retsch). Cell wall preparation was performed by incubating the samples three consecutive times in 95% EtOH at 65°C for 30 min, followed by a treatment in chloroform:methanol (2:1, v/v) at room temperature for 1 hour under gentle agitation. The insoluble material was then successively washed in 70% (two times 1.5 h), 80% (1 h) and 95% (2 h) ethanol and dried under vacuum (SpeedVac Plus, Savant) after a final wash in acetone. The resulting alcohol-insoluble residue was resuspended in 500 µl of an α-amylase solution (5 U/ml; Sigma-Aldrich
A3176) in 0.01 M phosphate buffer pH 7 and incubated for 24 h at 37°C under continuous stirring (Melida et al., 2009). The resulting de-starched cell wall residue was washed three times with 70% ethanol, followed by three times with acetone, and stored at room temperature for further analysis. Neutral sugar composition was determined after sulfuric acid hydrolysis (Saeman et al., 1954). For this purpose, the cell wall samples were resuspended in 72% sulfuric acid and kept in this solution for 3 h at room temperature before being heated at 100°C for 3 h. Myo-inositol was used as an internal standard. The samples were then passed through 0.2-µm nylon filters and diluted 5X with deionized water. The hydrolysates were then subjected to high-performance anion-exchange chromatography (HPAEC) using a Dionex Carbopac PA1 column and a Dionex HPLC fitted with a pulsed amperometric detector (PAD, Dionex ICS 3000 system). The samples were eluted over 20 min with deionised water and the neutral monosaccharides were detected following post-column addition of 300 mM sodium hydroxide at a flow rate of 0.5 ml/min. Glycosidic linkage analyses were performed using 0.8 mg of cell wall samples. The latter were swollen in 400 µl dry dimethyl sulfoxide for 3 h with stirring at room temperature prior to sugar derivatization to permethylated alditol acetates and GC/MS analysis, as described earlier (Melida et al., 2013). Neutral sugar and linkage analyses were performed in triplicate.

**Fusicoccin treatment and AHA2\textsuperscript{695} induction**

Col-0 or AHA2\textsuperscript{695} seeds were stratified for 4 days in water at 4°C, then germinated and grown for 4 days in half strength MS medium adjusted to pH 5.7. For fusicoccin treatment, Col-0 seedlings were transferred to solid medium supplemented with DMSO (mock), or either 3 µM or 10 µM fusicoccin (Sigma) and grown for an additional 24 hours. For analysis of forced apoplastic acidification, AHA2\textsuperscript{695} seedlings were transferred to solid medium supplemented with DMSO (mock) or 1 µM dexamethasone (Sigma) and grown for an additional 24 hours. Propidium iodide-stained roots were then analyzed under a Zeiss LSM780 confocal microscope.

**pH microelectrode measurements**

For root surface pH measurements, we used a pHOptica micro fiber optic pH system (World Precision Instruments) with pH1-micro-AOT-06-059 fiber-optic microsensors.
(PreSens) (pH range 4.0–9.0). Five equidistant points along the root tip surface were measured in 4-day-old seedlings grown in tissue culture. Media pH was verified in a distant location from the seedlings.

Accession numbers

Sequence data from this article can be found in the GenBank/EMBL libraries under accession numbers \textit{Bradi2g04290} (\textit{TAR2L}), \textit{Bradi4g08380} (\textit{EIN2L1}), \textit{Bradi2g04860} (\textit{VAS1L}) and \textit{AT4G30190} (\textit{AHA2}). The RNAseq raw data are available at the National Center for Biotechnology Information Sequence Read Archive under accession number SRP072551.

Supplemental Data

Supplemental Figure 1. Supplemental illustrations of Brachypodium phenotypes.

Supplemental Figure 2. Supplemental illustrations of pH experiments.

Supplemental Data Set 1. RNAseq quantification, read counts for all samples.

Supplemental Data Set 2. Gene expression level comparison, \textit{tar2}^{hypo} versus Bd21-0.

Supplemental Data Set 3. Gene expression level comparison, L-kynurenine-treated versus mock-treated Bd21-0.


Supplemental Data Set 5. Gene expression level data for selected genes of interest plotted in Figure 3B-G.

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Author Contributions

DPV, SW, CO, KL, VB and CSH designed the research. DPV, SMDM, AvdS, TT, YHK, BG, ON, NJ and ZL performed the research. All authors analyzed data. DPV and CSH wrote the paper with input from the other authors.

Figure legends

Figure 1. Reduced tryptophan aminotransferase activity in seminal roots of hypomorphic tar2l mutants results in higher cellular auxin levels and strongly enhanced cell elongation. (A) 4-day-old tissue culture-grown Brachypodium wild type (Bd21-0) and hypomorphic tar2l mutant (tar2lhypo) seedlings (left) and light microscopy images of their mature cortex (c) and epidermal (e) cell layers (right). Double-sided arrows point out the longitudinal dimensions of individual cells. (B) Mature cortex cell length in roots 4 days after regeneration from isolated Bd21-0 and tar2lhypo root tips (40 cells per root, 10 roots). (C) Tryptophan aminotransferase background activity in Bd21-0 and tar2lhypo roots. (D) Total tryptophan aminotransferase activity in Bd21-0 and tar2lhypo roots, background portion indicated in grey. (E) Quantification of auxin and auxin metabolites in 1 cm segments from the root elongation zone of 4-day-old Bd21-0 or tar2lhypo seedlings. Error bars indicate standard error of the mean (3-4 biological replicates). Differences were not statistically significant (Student’s T-test) unless indicated as follows: * = p < 0.05; ** = p < 0.01; *** = p < 0.001;

Figure 2. Differential gene expression in root segments as determined by RNA sequencing. (A) Venn diagrams illustrating overlaps between the gene sets that were differentially expressed in root segments of Bd21-0 versus tar2lhypo, Bd21-3 versus ein2lhypo, and mock-treated versus L-kynurenine-treated Bd21-0. (B) Cluster analysis and (C) principal component (PC) analysis of the different RNA sequencing samples.
Figure 3. Differential expression of auxin- and cell wall-related genes in root segments. (A) Word cloud from annotations of genes differentially expressed between Bd21-0 versus tar2<sup>hypo</sup>, or mock-treated versus L-kynurenine-treated Bd21-0. (B-G) Expression changes (fold changes) for individual members of the indicated gene classes in Bd21-0 versus tar2<sup>hypo</sup> and mock-treated versus L-kynurenine-treated Bd21-0 (Bd21-0 / mock set to 1 on the left, tar2<sup>hypo</sup> / L-kynurenine-treated Bd21-0 values on the right). Only genes that showed differential expression and a q value < 0.01 in at least one comparison are plotted. See Supplemental Data Set 5 for gene identifiers and expression values. (H) qPCR verification of differential gene expression in independent RNA samples prepared from independent root segments (3 biological replicates). Error bars indicate standard error of the mean. Differences were not statistically significant (Student’s T-test) unless indicated as follows: * = p < 0.05; ** = p < 0.01; *** = p < 0.001.

Figure 4. Cell wall analyses of tar2<sup>hypo</sup> and ein2<sup>l1</sup>hypo root segments compared to their wild type backgrounds. (A) Glycosidic linkage analysis of Bd21-0 and tar2<sup>hypo</sup> root segments. (B) Glycosidic linkage analysis of Bd21-3 and ein2<sup>l1</sup>hypo root segments. (C) Neutral sugar analysis of Bd21-0 and tar2<sup>hypo</sup> root segments. (D) Neutral sugar analysis of Bd21-3 and ein2<sup>l1</sup>hypo root segments. (E) Yariv staining (brownish) against β-1,3-galactan linkages in AGPs on longitudinal sections of Bd21-0 and tar2<sup>hypo</sup> root tips. (F) 2F4 antibody staining against de-methylesterified pectin (green) on transverse sections from the elongation zone of Bd21-0 and tar2<sup>hypo</sup> root tips. (G) JIM7 antibody staining against methylesterified pectin (red) on transverse sections from the elongation zone of Bd21-0 and tar2<sup>hypo</sup> root tips. (H) LM2 antibody staining against AGP side chains (red) on transverse sections from the elongation zone of Bd21-0 and tar2<sup>hypo</sup> root tips. Error bars indicate standard error of the mean (2 technical replicates per each of 3 biological replicates). Differences were not statistically significant (Student’s T-test) unless indicated as follows: * = p < 0.05; ** = p < 0.01; *** = p < 0.001.

Figure 5. Medium acidification by Bd21-0 and tar2<sup>hypo</sup> root tips. (A) Medium acidification through proton excretion from seminal roots of 3-day-old seedlings, 4 hours after transfer onto fresh medium with pH indicator. (B) Same as (A), 24 hours after transfer. (C) Progressive acidification of liquid medium through proton excretion from
seminal root tips starting with 2-day-old seedlings (6 biological replicates). (D) pH traces
along the surface of root tips, measured with a fiber-optic pH microsensor at five
equidistant points as indicated (10-12 biological replicates). (E) Protein gel blot antibody
detection of H⁺-ATPases in protein samples isolated from microsomes of root segments
and overlay with 14-3-3 protein binding. Error bars indicate standard error of the mean.
Differences were not statistically significant (Student's T-test) unless indicated as
follows: * = p < 0.05; ** = p < 0.01; *** = p < 0.001.

Figure 6. Root cell elongation in response to external pH variation or forced
apoplastic acidification. (A) Mature cortex cell length in Bd21-0 wild type roots 2 days
after transfer of 2-day-old seedlings from standard pH (5.7) to standard pH or more
acidic pH (5.2) (103-122 cells from 6-8 roots). Only cells formed after transfer were
scored. (B) Root tip surface pH traces obtained with a fiber-optic pH microsensor at five
equidistant points as indicated (8 biological replicates). (C) Root length of Bd21-0 wild
type seedlings 2 days after transfer of 2-day-old seedlings from standard pH (5.7) to
standard pH or very acidic pH (3.7) (8-10 roots). Note that on pH 3.7, root growth is
severely inhibited and meristematic activity gradually ceases. (D) Mature cortex cell
length of roots in (C) (75-97 cells). Only cells formed after transfer were scored. (E)
Mature cortex cell length in roots of 5-day-old Arabidopsis Col-0 wild type seedlings
grown on media with different pH (145-157 cells from 19-20 roots). (F) Mature cortex
cell length in roots of 4-day-old Arabidopsis Col-0 wild type seedlings, 24 hours after
transfer from standard medium onto mock- or fusicoccin-supplemented medium. Only
cells formed after transfer were scored (28-53 cells from 6 roots). (G) Mature cortex cell
length in roots of 4-day-old transgenic Arabidopsis AHA2δ seedlings, 24 hours after
transfer on medium supplemented with 1 µM dexamethasone to induce expression of
the hyperactive proton pump (25-74 cells from 5 roots). Cells were scored in mock-
treated roots, as well as before (pre-treatment) and after induction in the same roots.
(H) Confocal microscopy of epidermal root cells in mock-treated or dexamethasone-
induced AHA2δ seedlings, 4 hours after induction, FM4-64 staining. Vacuoles can be
easily distinguished in the inverted grey scale images. Error bars indicate standard error
of the mean. Differences were not statistically significant (Student's T-test) unless
indicated as follows: * = p < 0.05; ** = p < 0.01; *** = p < 0.001.


The Effects of High Steady State Auxin Levels on Root Cell Elongation in Brachypodium
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