Auxin-dependent control of cytoskeleton and cell shape regulates division orientation in the Arabidopsis embryo

Highlights

- Auxin responses regulate directional cell expansion in Arabidopsis embryos
- Cell shape and division orientation are tightly coupled
- Transcriptome analysis identifies MT-associated IQD proteins in division control
- Cytoskeletal dynamics control division orientation

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In brief

By combining 3D imaging of cell shapes, nuclear and polarity markers, transcriptome profiling, genetic analysis, and targeted perturbation of cytoskeletal structures, Vaddepalli, de Zeeuw et al. reveal that auxin-dependent control of cell division orientation in Arabidopsis embryos involves regulation of cytoskeleton and cell shape.
SUMMARY

Premitotic control of cell division orientation is critical for plant development, as cell walls prevent extensive cell remodeling or migration. While many divisions are proliferative and add cells to existing tissues, some divisions are formative and generate new tissue layers or growth axes. Such formative divisions are often asymmetric in nature, producing daughters with different fates. We have previously shown that, in the Arabidopsis thaliana embryo, developmental asymmetry is correlated with geometric asymmetry, creating daughter cells of unequal volume. Such divisions are generated by division planes that deviate from a default “minimal surface area” rule. Inhibition of auxin response leads to reversal to this default, yet the mechanisms underlying division plane choice in the embryo have been unclear. Here, we show that auxin-dependent division plane control involves alterations in cell geometry, but not in cell polarity axis or nuclear position. Through transcriptome profiling, we find that auxin regulates genes controlling cell wall and cytoskeleton properties. We confirm the involvement of microtubule (MT)-binding proteins in embryo division control. Organization of both MT and actin cytoskeleton depends on auxin response, and genetically controlled MT or actin depolymerization in embryos leads to disruption of asymmetric divisions, including reversion to the default. Our work shows how auxin-dependent control of MT and actin cytoskeleton properties interacts with cell geometry to generate asymmetric divisions during the earliest steps in plant development.

INTRODUCTION

Mechanisms controlling plant division plane orientation have been an area of focus for over a century.\(^1\) Starting from the establishment of the early embryo to the development of post-embryonic tissues and organs, plants need to constantly calibrate the coordination between cellular and genetic inputs for proper cell and tissue patterning. Failure in the coordination leads to aberrant phenotypes with severe developmental defects.\(^4\,\,5\) Proliferative mitotic cell divisions select symmetric division plane, resulting in cells with approximately equal size. In formative divisions, however, division planes strongly deviate from the symmetric position, leading to daughter cells of different sizes. Such asymmetric divisions often lead to the formation of new cell identities and tissue layers, and these divisions can thus lead to differential developmental fates.

Plant cells by default divide along the minimal surface area (in 3D) following the “shortest wall” (in 2D) rule.\(^1\) Thus, cell geometry influences its plane of division and therefore is a fundamental input in determining the size and shape of daughter cells. Genetic regulation can interfere with the default symmetric division to facilitate division plane orientation.\(^6\) Recent evidence suggests that cytoskeleton dynamics may bridge the coordination of geometric and genetic input to influence the re-orientation of the division plane.\(^9\,\,10\) During the first asymmetric division of the zygote and in lateral root founder cells, dynamics of cytoskeletal pattern determine the correct orientation of division plane. In both these systems, however, cells are elongated, and the various orientations of division are dramatically different in terms of surface area and volume partitioning. A key question is whether similar mechanisms operate in smaller, polyhedral cells, where such differences are less extreme. The signaling cue for
biasing division plane orientation likely involves cell polarity mechanisms, but how the intracellular position of polarity proteins directs division plane orientations remains elusive. In several cell types, nuclear position co-aligns with the preprophase band (PPB), and migration of the nucleus is correlated with positioning of the division plane wall in the zygote, lateral root founder cells, and leaf epidermis. Again, all those cell types are either large, relative to nuclear size, or have extreme aspect ratios, and it remains a question whether the same principles apply to division control in other types of cells.

Developing from a fertilized egg cell, the early plant embryo is a hotspot for formative events: new cell types are established with many divisions, which in Arabidopsis are highly predictable. Using advanced imaging and cellular segmentation approaches, a 3D description of early Arabidopsis embryogenesis has been generated. From this work, it surfaced that divisions leading to the 2-, 4-, and 8-cell embryo stages follow the minimal surface area rule, corroborating classical (2D) models from the 19th century. However, the next asymmetric divisions that generate the protoderm and inner cells at the 16-cell stage deviated from this rule. Using mutant embryos in which response to the plant hormone auxin was blocked by ubiquitous expression of a transcriptional repressor (RPSSA>>bdl), it was demonstrated that transcriptional response to auxin is required to suppress the geometric default division, implicating that the regulation of oriented cell division by geometric and genetic cues can be uncoupled. Thus, the 8-cell Arabidopsis embryo represents a unique case where the activity of a transcriptional regulator (bdl) allows to switch between default, symmetric, and regulated asymmetric division. Based on a more recent computational model, it has been proposed that all division planes observed in wild-type and mutant cells conform to a default rule, provided that new walls can be curved when inserted. In the same study, it was suggested that, also in these cells, nuclear position may provide input into division plane position. Analysis of live embryos found little to no curvature in newly formed walls, and it is therefore an open question through what cellular processes, genes, and mechanisms the division orientation in early embryos is controlled. Conceptually, the axis of cell polarity, cell shape, and nuclear position can each influence and collectively interact to guide division plane position and orientation. The auxin-dependent switch in ultimate division plane allows to dissect at what point these processes can be regulated. Here, we explore mechanisms underlying division plane selection in the embryo.

Figure 1. Auxin response is required for asymmetric embryonic cell division

(A) 3D comparison of wild-type, RPSSA>>bdl, and tir1/afb mutant embryos. Cell colors correspond to cellular volume as indicated in the color scale. The percentage of symmetric orientation defects at 16-cell stage and number of cell pairs analyzed (% (n)) are indicated at the bottom.

(B) Violin plots representing distribution of division plane areas and volume ratios. The relative area is the actual division plane normalized by the smallest (0) and largest (1) possible division wall areas. Division planes were simulated through the center point of the actual division plane between two sister cells. The cell volume ratio plot of the daughter cells resulting from these divisions is presented in the right panel. Whiskers represent median ± 95% confidence intervals (CIs). At least two individual embryos were used per condition.

RESULTS

TIR1/AFB-dependent auxin response controls cell division orientation in the early embryo

Auxin promotes degradation of AUX/IAA proteins through TIR1/AFB receptors, thus promoting ARF-dependent gene expression. A mutation in the degron of AUX/IAA proteins prevents interaction with TIR1/AFB proteins, causing accumulation of mutant protein and permanent ARF inhibition. Expression of mutant iaa12/bdl protein in early embryos (RPSSA>>bdl) prevents asymmetric divisions at the 8-cell stage. However, because the mutant protein can accumulate to unnaturally high levels, this may lead to inhibition effects beyond the normal activity of auxin. Hence, it is not yet clear whether an endogenous auxin response process controls division orientation.

Therefore, we scrutinized 3D division orientation in mutant embryos lacking all 6 TIR1/AFB receptors, the tir1/afb sextuple (tir1afb12345) mutant. This mutant arrests during embryogenesis with defects that superficially resemble those observed in RPSSA>>bdl embryos. To compare division defects more directly, we made use of a sextuple homozygote that carries a heterozygous complementation transgene carrying TIR1-M妆:2::AFB5-mCherry::AFB2-mCitrine. The hemizygous tir1afb12345 mutant thus has 25% sextuple mutant progeny. Cell segmentation analysis using MorphoGraphX on mutant embryos confirmed the resemblance of division plane defects with the RPSSA>>bdl phenotype (Figure 1A). Division plane orientation did, however, show variability. Next, we measured volumes of pairs of sister cells at 8- and 16-cell stages to determine the volume distribution ratios as a proxy for division (a-) symmetry. We also analyzed the division plane surface area relative to the minimal and maximal planes cutting through the center of the actual division. In tir1afb12345 mutant embryos, divisions leading to 8-cell embryos are symmetric and use the
minimal surface area, similar to wild-type and RPS5A>>bdl embryos (Figure 1B). In tir1abf12345 mutant embryos, divisions leading to 16-cell embryos show a high variation in relative division plane surface area and volume ratios ranging between the values seen for RPS5A>>bdl and those in wild-type (Figure 1B). The average division is therefore more symmetric than wild-type. The observed variation in division represents a spectrum of division plane defects that clearly include those observed in RPS5A>>bdl embryos but also display weaker aberrations. It is unclear whether this is due to residual TIR1/AFB activity in the hextuple mutant. Nonetheless, this analysis shows that endogenous auxin response is required for promoting asymmetric cell division in the embryo through regulating division plane orientation.

Cell shape, not polarity axis or nuclear position, correlates with division orientation
To analyze the role of cell polarity, nuclear position, and cell shape in division orientation, we focused on 4-cell and 8-cell stage embryos that give rise to the 8-cell and 16-cell embryos, respectively. Because outer-inner cell polarity is established early in wild-type embryos,22 and because divisions at the 8-cell stage in wild-type are strictly aligned with this polarity axis, it is conceivable that the division defects in RPS5A>>bdl embryos reflect a loss of polarity. We addressed this question by imaging the inner WOX2pro:BOR1-mCitrine (ACE-W03) and outer WOX2pro:mCherry-NIP5;1 (ACE-W04) domain markers in RPS5A>>bdl and wild-type control (RPS5A>>Col) embryos. Despite characteristic defects in cell division orientation, no difference in BOR1-mCitrine and mCherry-NIP5;1 localization could be detected between RPS5A>>Col and RPS5A>>bdl embryos (Figures 2A–2D). As in wild-type embryos, the BOR1 marker is enriched at the inner cell membranes (Figures 2A and 2B) and mCherry-NIP5;1 to the outer membranes (Figures 2C and 2D) of defective 4-cell- and 8-cell-stage RPS5A>>bdl embryos. Thus, early outer-inner polarity axis establishment appears independent of transcriptional auxin response and the failure to divide asymmetrically in the mutant is likely not caused by global loss of polarity.

Asymmetric cell division in the zygote, in lateral root founder cells, and in meristemoid mother cells involve nuclear migration to the future division site, implying a strong association of division plane position with nuclear position.9,10,13,23 Compared to these systems, early embryonic cells have distinct cell geometry, and our previous observations on wild-type embryos suggested...
that the nucleus occupies a relatively large part of the cell, limiting its ability to move. A recent report proposed that nuclear position could constrain both symmetric and asymmetric division plane position in early Arabidopsis embryo cells. We explored this hypothesis in RPS5A>>bdl mutant embryo cells, which exhibit a consistent switch from asymmetric to symmetric division planes at the 16-cell stage. We also asked whether changes in the position of the nucleus correlate with the switch in division orientation in RPS5A>>bdl embryo cells.

We introduced the embryonic nuclear envelope marker WOX2pro:NUP54-GFP (ACE-W11) into the RPS5A-GAL4 background and crossed this line with wild-type or upstream activating sequence (UAS)-bdl to visualize the nuclear volume in RPS5A>>bdl and wild-type control embryos. We did not observe conspicuous differences in nuclear morphology between the two genotypes (Figure 2E). We next created nuclear and cell outline meshes by applying MorphoGraphX-based segmentation on the z stacks (Figure 2F) and defined cellular (CC) and nuclear centroid (NC) to calculate nuclear position relative to the centroid of the cell. Defining a central axis through the suspensor, we could measure absolute distance of NC to CC, as well as its displacement in longitudinal and radial directions (Figure 2G). For both 4-cell and 8-cell embryos, we could not find significant differences in nuclear position between wild-type and RPS5A>>bdl mutant embryos (Figure 2H). Interestingly, we found considerable variation in nuclear position even in wild-type cells, where the division plane is essentially invariable. These findings suggest that correlation of nuclear position with division plane position is not altered in the bdl mutant despite the switch to symmetric division. Additionally, these findings also imply that nuclear position may not be strongly connected to cell division orientation and is perhaps not a mechanism mediating its control in early embryos.

Given that minimal surface area and the cell centroid are defined by cellular shape, a switch to a different cell division plane in auxin-insensitive mutants could also be indirectly caused by altered cell shape. Segmentation analysis of wild-type embryos revealed that cells at the 4-cell stage exhibit higher cell volume and area compared to 8-cell stage (Figure 3A), corroborating our previous observation that early division transitions happen with minimal expansion. RPS5A>>bdl mutant embryos in comparison exhibit significantly larger cell volume and wall area at both 4-cell and 8-cell stages (Figure 3A), indicating that cell geometry is indeed affected in the bdl mutant. To determine whether the observed cell expansion is random or directed in either of the cellular directions, we measured circumferential, radial, and longitudinal cell lengths of the same embryonic cell volumes (Figure 3B). Although at 4-cell stage, cell length in RPS5A>>bdl embryos was slightly increased in comparison to wild-type in all measured directions, elongation in the longitudinal direction is dominant. At the 8-cell stage, the circumferential length is high in wild-type cells, while the radial and longitudinal lengths are similar to each other and lower (Figure 3C). In bdl embryos, however, longitudinal cell length is higher and closer to circumferential cell length compared to wild-type. To more directly measure cell shape, we calculated the ratio of radial
and circumferential cell lengths to the longitudinal cell length. At 4-cell stage, the ratios suggest minimal to no difference between wild-type and bdl embryos. In wild-type, 8-cell stage cells show higher radial and circumferential ratios compared to the 4-cell stage (Figure 3D), revealing obvious but significant cell shape changes during the 4- to 8-cell transition. In contrast, both ratios of bdl embryos at 8-cell stage are significantly lower in comparison to wild-type, indicating abnormal expansion defects in bdl embryos do lead to cell shape irregularities. Detailed analysis of cells in the upper (apical) and lower (basal) tier of the embryo revealed that shape defects in mutant cells are found in both tiers. In addition, the distinction in shape between apical and basal cells is reduced in RPS5A>>bdl embryos (Figures 3E and S1). Hence, this analysis shows that the altered cell division planes are preceded by changes in cell geometry, suggesting that the primary target of auxin control could be a process that controls cell shape.

Transcriptome analysis reveals altered cytoskeletal gene expression

To probe the genetic mechanisms underlying auxin-dependent cell expansion and division plane orientation, we performed transcriptome analysis, comparing manually isolated 8-cell wild-type and RPS5A>>bdl mutant embryos. Given that the molecular target of bdl is ARF-dependent transcriptional control, the immediate cellular pathways that are subject to auxin regulation should be apparent from the genes misregulated. We chose the 8-cell stage, as this represents the moment shortly before the switch in division orientation. Initial inspection of the RPS5A>>bdl transcriptome revealed the expected upregulation of BDL/IAA12 while other Aux/IAA genes were downregulated (Figure S2A), consistent with genome-wide dampening of auxin response. Additionally, 5 out of 11 YUC genes were upregulated in bdl embryos (Figure S2A), which shows that also auxin-dependent gene repression is inhibited in mutant embryos and validates the effectiveness of the inhibition of auxin response. After statistical analysis, we retained 421 up- and 414 downregulated genes in RPS5A>>bdl embryos (>2-fold difference; q value < 0.05; Table S1). Gene Ontology (GO) analysis did not identify obvious enrichment of functional categories. Nevertheless, among the highly misexpressed genes, we found several genes involved in cellular mechanisms, along with known developmental regulators. Here, we focus on 34 candidate genes that could be divided into three groups based on their ontology information and functional data from earlier studies (Figure 4A). The first group represents genes related to auxin signaling (IAA1, IAA9, and IAA30), biosynthesis (YUC1 and YUC8), and transport (PIN1, PIN4, LAX2, and NPY5). The second group includes transcription factors, of which most are known to be key regulators of development, including several known auxin response targets (e.g., TMO3, GATA20, WIP2, and TMO5). The third group contained genes known for their function in cytoskeletal organization and signaling, along with genes involved in cell wall composition and remodeling. A pectin methyl esterase (PME44), xyloglucan endotransglycosylase (XTH19), cellulase (CEL2), and an arabinogalactan protein (FLA12) were found downregulated in bdl embryos. All these are known for their roles in cell wall remodeling mechanisms during post-embryonic growth. We found significant downregulation of the ROP-activating guanine exchange factor, ROP-GEF5, along with ROP9, which belongs to type II sub-group of ROP gene family. The plant-specific small Rho guanosine triphosphatase (GTPase) switches, ROPs, are known for their function in tip-growing cells like pollen tube and root hair cells as well as interdigitating epidermal pavement cells by regulating actin-microtuble (MT) dynamics. Conversion from the inactive guanosine diphosphate (GDP)- to active GTP-bound
form of ROPs is triggered by ROP-GEFs. Additionally, two IQ67 domain (IQD) family genes IQD6 and IQD18 were also found downregulated in the RPSSA>>bdl background. IQD proteins interact with calmodulin (CaM) signaling modules and are proposed to mediate Ca²⁺-dependent regulation of MT organization and dynamics. IQD proteins are also emerging as key components in ROP signaling by regulating plasma membrane-MT interactions of the actin and MT cytoskeleton function in auxin-insensitive embryos.

**Auxin response controls cytoskeleton organization in the embryo**

The altered expression of a set of genes encoding regulators of actin and MT cytoskeleton function in auxin-insensitive embryos suggests that auxin response controls these two cytoskeletal structures. We have previously demonstrated that length and degree of MT polymerization is reduced in RPSSA>>bdl embryos, and modeling suggested this to contribute to choice of division plane. To address whether actin organization is also altered, we introduced a WOX2pro::LifeAct-tdTomato (ACE-W14) marker into the RPSSA>>bdl background. Previously, we reported thick F-actin bundles in early embryonic cells, which form arches around the nucleus (Figure 4B). These thick actin bundles were absent in RPSSA>>bdl cells (Figure S2B), and in addition, we observed loss of dense F-actin meshwork in mutant cells (Figure 4B). Quantification revealed a strong increase in cytosolic LifeAct-tdTomato signal in RPSSA>>bdl embryos (Figure S2C). Thus, in addition to the effects on the MT cytoskeleton, impaired auxin response causes a disruption of the actin cytoskeleton in the embryo. By inference, auxin controls the organization of both cytoskeletal structures.

**Auxin-dependent, MT-associated IQD6 functions in division control**

It is likely that the influence on cytoskeleton function that auxin exerts is mediated by the genes identified as being downregulated in RPSSA>>bdl embryos. Here, we focused on the IQD6 gene, which was strongly downregulated (Figure 4A). Previously, inhibition of auxin response on other developmental contexts had been shown to affect the expression of several IQD family members. Indeed, apart from IQD6 and IQD18, we also observed downregulation of several IQD family genes in the RPSSA>>bdl mutant background (Figure S3A). We first determined the subcellular localization of IQD6 protein, as well as its close homologs IQD7 and IQD8, by generating C-terminal fusions with YFP (IQDXpro:IQDX-sYFP). All three proteins exhibited filamentous localization near the cortical MT localization reported for embryos previously. We also observed downregulation of several IQD family genes in the RPSSA>>bdl mutant background (Figure S3A). We first determined the subcellular localization of IQD6 protein, as well as its close homologs IQD7 and IQD8, by generating C-terminal fusions with YFP (IQDXpro:IQDX-sYFP). All three proteins show broad accumulation in embryos and roots, with IQD6 and 7 showing a slight enrichment in the root vasculature (Figures 5A, S4A, and S4B).

To investigate the involvement of IQD6, 7, and 8 in division plane orientation, we analyzed the embryos of iqd678 triple mutants. This mutant was recently reported to have distorted cell division planes in roots, along with alterations in the organization of the MT cytoskeleton. Likewise, in 35% of the analyzed embryos (Figure S3B), the mutant shows a shift in division plane orientation during different stages, which varies from subtle to more severe defects (Figure S3C). The divisions leading to 8-cell embryos are symmetric and use the minimal surface area, similar to wild-type embryos (Figures S3D and S3E). In contrast, the divisions leading to 16-cell embryos show a high...
degree of division plane area variation, with values spanning across and beyond the relative areas of wild-type 8-cell and 16-cell stages (Figures 5D and 5E). Consequently, volume distribution ratios are also highly variable, from wild-type-like asymmetric divisions to highly symmetric divisions with volume ratios smaller than RPSSA>>bdl mutant embryos (Figures 1B and 5E). Nevertheless, the iqd678 mutant embryos do not show the severe bdl-like division phenotype (Figure 5D). These results suggest that IQD6–8 proteins are involved in cell division placement. However, the variability in division plane parameters indicate that this function is not absolute and may signify the involvement of other IQD proteins or additional components.

Both MT and actin cytoskeletons contribute to regulated division plane orientation

Disruption of MT and actin cytoskeleton is correlated with altered cell division planes in RPSSA>>bdl embryos, and this is in turn coupled to altered expression of genes encoding cytoskeletal regulators. It is thus likely that cytoskeleton organization contributes causally to division plane choice in embryo cells. If this were the case, one would expect direct interference with either cytoskeleton to alter cell division planes. Prolonged treatments with cytoskeleton-destabilizing or stabilizing drugs in embryos are not trivial and require in vitro culturing of seeds, which in itself can cause abnormalities.36 We therefore made use of genetic tools to depolymerize MT or F-actin filaments by expression of the PHS1Δ (MT)37 or SpvB (actin)38 proteins. Expression of these proteins was previously shown to be equivalent to treatment with MT- or actin-depolymerizing drugs and disrupts asymmetric radial expansion and polar migration of nuclei in lateral root founder cells.10 We generated fluorescently tagged versions of PHS1Δ (mNeonGreen-PHS1Δ) and SpvB (SpvB-mNeonGreen) and used GAL4-UAS two-component gene expression to drive their expression in embryos only after fertilization, driven by the RPSSA promoter. Expression of both proteins in embryos caused frequent changes in division planes. MT depolymerization through PHS1Δ expression led to defects in 95% of embryos (n = 122) and caused oblique divisions (Figures 6A and S5) that superficially resemble those induced by inhibition of auxin response. Depolymerizing actin through SpvB expression led to essentially indistinguishable defects in 85% of embryos (n = 165). Also, here, altered division planes are similar to those observed in RPSSA>>bdl and tir1afb12345 embryos (Figures 6A and S5). Defects were obvious at all stages analyzed (Figures 6A and S5). While wall area and cell volume ratio were more variable than in wild-type at 8-cell stage (Figures 6B and S6), there was no consistent switch to altered division plane. At 16-cell stage, the asymmetry and volume distribution among cells were also more variable than in wild-type (Figure 6B). In a small number of cells, division wall surface area was larger than in wild-type, correlating with a small population of cells with more asymmetric division (Figure 6B). At the same time, a larger fraction of cells show bdl-like orientation defects (Figure 6A), leading to smaller division wall surface area and thus overall lower average value in both RPSSA>>PHS1Δ and RPSSA>>SpvB embryos (Figure 6B). Consequently, division asymmetry was also reduced in both transgenic genotypes. Thus, while depolymerization of both cytoskeletons expectedly caused more pleiotropic defects in cell division plane orientation, these defects include the switch to smaller division wall surface area and loss of asymmetric division. We therefore conclude that regulation of the MT and actin cytoskeleton is critical for asymmetric cell division in the early Arabidopsis embryo and that auxin response may indeed regulate division orientation through its effects on the cytoskeleton.
DISCUSSION

Combining embryo-specific fluorescent cellular reporters with 3D imaging and cell segmentation, we analyzed the role of early polarity axis, nuclear position, cell shape, and auxin-mediated cytoskeleton dynamics in orienting the division plane in early embryos. Surprisingly, despite nearly invariant division planes, nucleus position was found to be variable, even in wild-type. Unless we missed a transient stabilization of nuclear position just prior to mitosis, this finding suggests that nuclear position is not instructive for positioning the division plane in early embryo cells. What is evident though is that changes in cell shape between wild-type and auxin-insensitive embryos correlate with altered division planes. This identifies the control of cell shape as a mediator of division plane choice (Figure 6C). While cell wall biology is complex and multifactorial, a key influence on cell shape is the deposition of cellulose fibers along tracks that are dictated by the CMT filaments. Thus, MTs dictate the pattern of cell wall fortification and thereby constrain and bias directional elongation, resulting in cell shape changes.39 Using markers for MT and actin, we show that the organization of both is subject to auxin-dependent regulation. This finding is consistent with their central role in post-embryonic division orientation control.39,41 We identified auxin-dependent genes with known functions in cytoskeleton reorganization. Characterization of IQD6 and its family members imply a significant role in orienting the division plane. While we did not explore other auxin-dependent genes here, several link to the processes identified to be critical to division orientation. First, a ROP11-IQD13 signaling module was found important for localized growth changes in the formation of xylem pits by organizing CMTs.29 ROPs are well known to regulate cytoskeleton dynamics during tip growth and in pavement cell interdigitation.42 CMT stability and polymerization dynamics regulate the PPB formation.30,44 Simulation studies of CMTs on segmented embryonic cell shapes identify transient auxin-mediated CMT stabilization as a plausible mechanism in division plane orientation.30 Thus, ROP-mediated cytoskeleton dynamics may play a critical role in keeping the homeostasis of CMTs and fine-tuning of the PPB for asymmetric orientation. Second, the identification of a set of cell-wall-related enzymes suggests that auxin regulation may also directly control wall biochemistry. Recent work has revealed the significance of pectin and xyloglucan in cell wall integrity and remodeling. Methyl esterification status of pectin determines the plasticity of the cell wall, and defects lead to severe phenotypes in post-embryonic tissues.45 These studies represent the wall reorganization effects at tissue and organ level, but our current knowledge about PME or XTH wall remodeling in confined cellular mechanisms like division plane orientation remains poor. The current study opens new avenues for answering these intriguing questions.

We show that outside-inside cell polarity establishment is independent of transcriptional auxin response. Thus, unless the outside-inside polarity system has multiple (auxin-dependent and independent) branches, auxin acts to control division downstream of this polarity axis. Because no regulators of this axis with function in the embryo have yet been identified, it is at present unknown how this polarity axis still biases the choice of the division plane, and it will be interesting to see how this interacts with the auxin-dependent control of cytoskeleton dynamics and cell shape. Recently, we identified a family of SOSEKI polarity proteins, of which two members are transcriptionally controlled by auxin response.46 Thus, at least part of the polarity system is dependent on auxin input. Misexpression of the SOSEKI1 protein causes oblique cell divisions, suggesting a link to the division orientation machinery. However, it is equally likely that SOSEKI1 affects the CMT or cell shape and only indirectly influences division plane. Further investigation of this protein family should help resolve how the different cell polarity systems are linked to division control in the embryo.

STAR METHODS

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SUPPLEMENTAL INFORMATION

Supplemental information can be found online at https://doi.org/10.1016/j.cub.2021.09.019.

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AUTHOR CONTRIBUTIONS

Conceptualization, P.V., T.d.Z., and D.W.; investigation, P.V., T.d.Z., S.S., C.-Y.L., and J.J.R.; resources, K.B.; formal analysis, P.V., T.d.Z., and S.S.; software and methodology, S.S. and R.S.S.; supervision, R.S.S. and D.W.; funding acquisition, P.V. and D.W.; writing—original draft, P.V.; writing—review & editing, P.V. and D.W. with input from all the authors.
DECLARATION OF INTERESTS

The authors declare no competing interests.

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# STAR METHODS

## KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Chemicals, peptides, and recombinant proteins** | | |
| SCRi Renaissance Stain 2200 | Renaissance Chemicals | SR2200 |
| Propidium Iodide | Sigma-Aldrich | Cat#P4170 |
| Paclitaxel, 99+%, ACROS Organics | Fisherscientific | Cat#33069-62-4 |
| Plant Agar | Duchefa | Cat#P1001.1000 |
| Phosphinotricin | Duchefa | Cat#P0159.1000 |
| Kanamycin (Sulfate Munohydrate) | Duchefa | Cat#K0126.0010 |
| **Critical commercial assays** | | |
| Ovation Pico WTA System V2 | Nugen | Cat#3302-12 |
| **Deposited data** | | |
| Microarray | This study | GEO: GSE165986 |

## Experimental models: organisms/strains

| Arabidopsis thaliana accession Col-0 | Widely distributed | N/A |
| Arabidopsis: tir1/atf6 hexuple mutant | Prigge et al.20 | N/A |
| Arabidopsis: RPS5Apro:GAL4-VP16 | Weijers et al.47 | RPSSA-GAL4 |
| Arabidopsis: UASpro:bdh | Weijers et al.48 | UAS-bdh |
| Arabidopsis: WOX2pro:BOR1-mCitrine | Liao and Weijers39 | ACE-W03 |
| Arabidopsis: WOX2pro:AtNUP54-GFP | Liao and Weijers39 | ACE-W11 |
| Arabidopsis: WOX2pro:Lifeact-tdtTomato | Liao and Weijers39 | ACE-W14 |
| Arabidopsis: IQD6pro:IQD6-sYFP | This study | N/A |
| Arabidopsis: IQD7pro:IQD7-sYFP | This study | N/A |
| Arabidopsis: IQD8pro:IQD8-sYFP | This study | N/A |
| Arabidopsis: UASpro:PHS1\ดาmNeonGreen | This study | N/A |
| Arabidopsis: UASpro:mNeonGreen-SpvB | This study | N/A |

## Oligonucleotides

For primers used see Table S1 | N/A | N/A |

## Recombinant DNA

| Plasmid: IQD6pro:IQD6-sYFP | This study | N/A |
| Plasmid: IQD7pro:IQD7-sYFP | This study | N/A |
| Plasmid: IQD8pro:IQD8-sYFP | This study | N/A |
| Plasmid: UASpro:PHS1\ดาmNeonGreen | This study | N/A |
| Plasmid: UASpro:mNeonGreen-SpvB | This study | N/A |

## Software and algorithms

| Fiji | Schindelin et al.49 | https://fiji.sc/ |
| MorphoGraphX | Barbier de Reuille et al.21 | https://www.mpipz.mpg.de/MorphoGraphX |
| R | R Development Core Team10 | https://www.r-project.org/ |

## RESOURCE AVAILABILITY

### Lead contact

Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Dolf Weijers (dolf.weijers@wur.nl).
Materials availability
DNA constructs and transgenic Arabidopsis seeds generated in this study are available from the lead contact, Dolf Weijers, upon request.

Data and code availability
The transcriptome data have been deposited in the NCBI Gene Expression Omnibus and are accessible through accession number GSE165986. The misregulated genes are listed in Table S1. Microscopy data reported in this paper will be shared by the lead contact upon request.

No new code has been generated in this study.

Any additional information required to reanalyze the data reported in this paper is available from the lead contact upon request.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Plant Material and Growth Conditions
Arabidopsis ecotype Columbia-0 (Col-0) was used for generating all the transgenic lines. Plants were grown under a constant temperature of 22°C with a 16-hr light/8-hr dark cycle in custom-built growth chambers with light intensity of 110 μE m⁻² s⁻¹. Arabidopsis seeds were surface sterilized using commercial bleach (4% Sodium hypochlorite) and 75% ethanol, and were subsequently plated on half-strength Murashige and Skoog (MS) medium with 0.7% plant agar (Duchefa, the Netherlands). After a 48-hour stratification and 10 days of growth on plates, seedlings were transferred to soil. tir1 afb hexaploid mutant seeds were provided by Mark Estelle and Michael Prigge (UCSD). ACE-W03 (WOX2pro:BOR1-mCitrine), ACE-W11 (WOX2pro:AtNUP54-GFP) and ACE-W14 (WOX2pro:Lifeact-tdTomato) were previously described. For all crosses RPS5A-GAL4 (RPS5Apro:mGAL4-VP16) was used as a female parent. For bdl embryo geometric analysis, F1 seeds of cross between RPS5A-GAL4 and Col-0 or UAS-bdl were used. For nuclear position, F1 seeds of cross between WOX2pro:AtNUP54-GFP (RPS5A-GAL4) and Col-0 or UAS-bdl were used. For early polarity analysis, F1 seeds of cross between WOX2pro:BOR1-mCitrine (RPS5A-GAL4) and Col-0 or UAS-bdl were used. For F-actin organization, F1 seeds of cross between WOX2pro:Lifeact-tdTomato (RPS5A-GAL4) and Col-0 or UAS-bdl were used. For analysis of CMT F1 seeds of cross between UASpro:PHS1Δp-mNeonGreen and Col-0 or RPS5A-GAL4 were used.

For analysis of F-actin F1 seeds of cross between UASpro:mNeonGreen-SpvB and Col-0 or RPS5A-GAL4 were used. Seeds of wild-type (Col-0) and T-DNA insertion lines for IQD6 (At1g17480, iqd6: SALK_107689), IQD7 (At1g17480, iqd7: SALK_025224) and IQD8 (At1g72670, iqd8: SALK_107689) were obtained from the Nottingham Arabidopsis Stock Center. All lines were backcrossed at least once with Col-0 and subsequently iqd6, iqd7, and iqd8 were crossed among themselves to generate iqd678 triple mutant.

METHOD DETAILS

Molecular cloning
Plasmids were cloned based on previously described ligation-independent cloning methods and vectors. Whole genomic IQD-sYFP fusions were prepared by cloning up to 3kb of promoter including downstream genomic region up to the stop codon into the pPLV117, containing a super Yellow Fluorescent Protein (sYFP). To generate UASpro:PHS1Δp-mNeonGreen and UASpro:mNeonGreen-SpvB plasmids, PHS1Δp-mNeonGreen and mNeonGreen-SpvB sequences were made by overlapping PCR and introduced into HpaI linearized pPLV32. All oligonucleotides used in this study are listed in primer Table S2. IQD-sYFP fusions, UASpro:PHS1Δp-mNeonGreen and UASpro:mNeonGreen-SpvB were transformed into the Col-0. ACE plasmids were transformed into homozygous RPS5A-GAL4 (RPS5Apro:GAL4-VP16) driver line.

Generation and selection of transgenic plants
Transgenic plants were generated by Agrobacterium-mediated floral dip transformation of constructs (see previous section) into backgrounds indicated. Transgenic individuals were selected based on resistance to the appropriate antibiotic or herbicide (15 mg/L phosphinotricin; 50 mg/L kanamycin) on 0.5xMS media, and transplanted to fresh, unsupplemented plates or soil after resistance became apparent. Transgenic lines were analyzed in T2, T3 or T4 generation.

Microscopy and image analysis
Embryos were stained by the modified Pseudo-Schiff propidium iodide (mPS-PI) staining method described in Yoshida et al. with the following modification: An extra treatment with 1% SDS and 0.2 M NaOH for 10 minutes at 37°C was added after fixation. The stained ovules/ embryos were mounted in a drop of chloral hydrate in a well generated by SecureSeal round imaging spacers (20mm, Thermofisher) and observed by confocal microscopy taking 2 stack images. A series of 2D confocal images were recorded at 0.1 μm intervals using a Leica TCS SP5II confocal laser scanning microscope with a 63 x NA = 1.20 water-immersion objective with pinhole set to 1.0 airy unit. PI was excited using a diode laser with excitation at 561 nm and detection at 600-700 nm.

Embryo samples were prepared as described in Liao and Weijers. Images for qualitative purpose were acquired in 8-bit format, images for segmentation were acquired in 16-bit format. Images were acquired using a Leica TCS SP5II confocal laser scanning microscope.
For segmentation, in MophoGraphX (MGX), 21 confocal image stacks (TIF) were Gaussian blurred using sigma value 0.6. 3D cell segmentation and nuclear position measurements interfered from the perinuclear actin arches. All image processes and measurements were conducted via Fiji. Cleavage signal intensity was documented and defined as the cytosolic Lifeact-tdTomato signal. 4 to 5 optical sections above the nucleus were used for the selection of ROIs. The optical sections containing the nucleus were not selected for the measurement to avoid interference from the perinuclear actin arches. All image processes and measurements were conducted via Fiji.

**3D cell segmentation and nuclear position measurements**

For segmentation, in MophoGraphX (MGX), 21 confocal image stacks (TIF) were Gaussian blurred using sigma value 0.6 μm, subsequently we applied the ITK watershed auto-seeding with level threshold value in the range 300–1500 and default smoothing levels. Segmented bitmap stacks were manually corrected for oversegmentation errors within MGX by fusing together multiple labels into the single cells, which were represented using a combination of the select and paint bucket tools in MGX. Then, we approximated the segmented cells by creating triangulated surface meshes using marching cubes 3D with cube size of 1. Nuclear measurements were performed on segmented meshes created using the same segmentation method described above using the nuclei marker channel. Cell and nucleus centroid positions were determined in MGX by calculating the center of gravity of their triangulated surface meshes. Organic central directions were determined in the same way as described in the 3D Cell Atlas Add-on for MGX by manually placing a straight line through the embryo using the “Bezier line” in MGX. For each cell then 3 directions relative to this central line were calculated: a longitudinal direction that is identical with the direction defined by the central line, a radial direction that was defined by the cell centroid and its closest point on the central line, and a circumferential direction that was defined by the cross product of the previous two directions. To calculate the distances between cell centroid and nucleus centroid along the longitudinal and radial direction, the scalar product of the vector defined by the centroids and the vector of the respective direction was computed.

**3D cell morphology measurements**

Cell sizes along longitudinal, radial and circumferential directions were computed in MophoGraphX by first performing a Principal Component Analysis (PCA) on the voxels of segmented cells. Then the component of the PCA’s tensor along the axis of interest was computed.

**Shortest division plane estimation and comparison to actual division plane**

To compute the relative division plane area, we used the following pipeline in MorphoGraphX which was adapted from Barbier de Reuille et al. First, we calculated the center of actual division plane by calculating the center of gravity of the triangulated surface meshes of the shared wall between two neighboring daughter cells. Then the daughter cells of recently divided cells in segmented meshes were merged. The actual division plane was approximated as a flat wall by computing the principal components of the vertices that were located at the shared border of the two daughter cells. After we simulated a division using this flat wall to determine the surface area of the real division wall (A_real), then the mean areas of the top 0.1% shortest (A_min) and longest division planes (A_max) in merged cells were determined by sampling of > 10000 division directions uniformly spread on the cell volume, going through the center of the actual division wall. Finally, we computed $\hat{A} = (A_{real} - A_{min}) / (A_{max} - A_{min})$, where $\hat{A}$ is the relative cell wall area, A_min the area of the shortest sampled division planes, A_max the largest sampled division planes, and A_real the area of the flat approximation of the real cell wall.

**Embryo isolation and transcriptome analysis**

Ovules were collected from ~60 siliques using vacuum extraction. Siliques were stuck to double-sided tape and sliced open using a needle. Open siliques were submerged in 1x PhosphateBuffered Saline (PBS) buffer and ovules were collected using a vacuum pump through 50 μm filters. Collected ovules were then transferred to isolation buffer (1x First Strand Buffer (FSB; Invitrogen), 1mM Dithiothreitol (DTT), 4% RNaseLater, MQ), and volume was reduced to ~20 μL. Embryo isolation was performed according to Raissig et al. with the following adaptations. A Zeiss ConfoCor 1 inverted microscope (Carl Zeiss Microscopy GmbH, Jena, Germany) together with an Eppendorf Transferman 4r micromanipulator (Eppendorf AG) and VacuTip II microcapillaries (Eppendorf) were used to isolate about 40–50 washed embryos in 50 μL isolation buffer.

RNA was amplified using the Ovation Pico WTA System V2 (NuGEN, CA, USA), labeled with the ENCORE Biotin Module (NuGEN) and hybridized to Arabidopsis Gene 1.1 ST arrays (Affymetrix, CA, USA) according to the manufacturers protocol. Microarray analysis was performed using the MADMAX pipeline and a custom CDF file (MBNI CustomCDF version 19.0.0). Here, all expression values were (quantile) normalized by the Robust multi-array average algorithm (RMA). Probe sets were redefined using current genome information and re-organized according to TAIR10 gene definitions. Linear models and an intensity-based moderated t statistic.
approach\textsuperscript{60,61} were used to identify differentially expressed genes (probe sets). P values were corrected for multiple testing using an optimized false discovery rate (FDR) approach.\textsuperscript{62}

QUANTIFICATION AND STATISTICAL ANALYSIS

Microsoft Excel was used to analyze the data. Plots-of-Data web app was used to plot measured data points, which is based on R.\textsuperscript{63} All error bars are represented as median ± 95% confidence intervals for each sample and were calculated by percentile bootstrap method provided by the Plots-of-Data web app. Sample sizes for each experiment are depicted in figure legend.
Supplemental Information

Auxin-dependent control of cytoskeleton and cell shape regulates division orientation in the *Arabidopsis* embryo

Prasad Vaddepalli, Thijs de Zeeuw, Sören Strauss, Katharina Bürstenbinder, Che-Yang Liao, João Jacob Ramalho, Richard S. Smith, and Dolf Weijers
Figure S1: 3D cell shape analysis of wild-type and bdl 8-cell embryos, Related to Figure 3. Cells were split into two groups (apical, basal) based on their position. (A) Average radial, circumferential and longitudinal cell lengths (in μm) are shown. Their ratios are shown in (B) to quantify cell shape changes.
Figure S2: Differential expression of selected genes and quantification of actin, Related to Figure 4.

(A) Differential expression of YUC and AUX/IAA genes in RPS5A>>bdl 8-cell mutant embryos. Note that 5 out of 11 YUC genes (YUC1,3,4,8,9) are upregulated in RPS5A>>bdl background compared to RPS5A>>Col control embryos. Many of the Aux/IAAs are downregulated (IAA1,2,4,8,9,13,18,19,20,26,27,30) in RPS5A>>bdl embryos except BDL (IAA12), which is highly upregulated. Those Aux/IAA’s whose expression is not altered have very low absolute expression levels, and can essentially be considered “not expressed” in the embryo. (B and C) Quantification of Lifeact-tdTomato (ACE-W14) reporter in wild-type (RPS5A>>Col) and RPS5A>>bdl embryos. Quantification of (B) perinuclear actin arches and (C) Average cytoplasmic signal intensity in RPS5A>>bdl 8-cell embryos compared to RPS5A>>Col control 8-cell embryos.
Figure S3: IQD gene family misexpression in RPS5A>>bdl embryos and iqd678 embryo phenotype, Related to Figure 5.

(A) Phylogenetic tree of all Arabidopsis IQD genes, rooted to IQD33, combined with their misexpression in 8-cell bdl (RPS5A>>bdl) embryos. Fold-change values are given for expression levels of genes in RPS5A>>bdl mutant embryos, relative to wild-type (RPS5A>>col). Yellow boxes indicate subclades previously shown to be misregulated in auxin-related datasets S1-S4. (B) Quantification of skewed division planes in iqd678 early embryos. Quantification is based on visual inspection of at least 250 individual chloral hydrate-cleared embryos. (C) iqd678 embryos can show either a subtle or severe phenotype (Stained using modified Pseudo-Schiff propidium iodide (mPS-PI) method). Scale bars: 5 μm.
Figure S4: IQD subcellular protein localization in roots and embryos, Related to Figure 5.
(A) IQD6, -7, and -8 are visualized using $p_{IQDX}$::$IQDX$-$s$YFP reporter lines. Scale bars: 5 µm.
(B) IQD subcellular protein localization in 5-day old Arabidopsis roots. IQD6-, 7-, and 8 are visualized using a $p_{IQDX}$::$IQDX$-$s$YFP reporter line merged with membrane visualisation using Propidium Iodide (PI) staining. Scale bars: 30 µm. (C) Details of roots with arrowheads indicate observed localization of protein in preprophase bands (PPB; blue), spindles (white) and phragmoplasts (magenta). Scale bars: 15 µm. (D and E) 2D embryo phenotype for the $iqd678$ mutant.
Figure S5: 2D embryo phenotype for RPS5A>>PHS1A and RPS5A>>SpvB- mutant embryos, related to Figure 6.

Embryos exhibit high variation of division plane defects among samples. Arrows indicate bdl-like division plane defects. Stained with SCRI Renaissance Stain 2200 (SR2200).
Figure S6: 3D phenotype of RPS5A>>PHS1A and RPS5A>>SpvB- mutant embryos with volumetric measurements, related to Figure 6.

Mesh colour per cell corresponds to cellular volume (in μm$^3$) indicated in the colour scale. Samples used for geometric analysis in Figure 4B.
| Primer               | Sequence                                                                 |
|---------------------|--------------------------------------------------------------------------|
| IQD6/AT2G26180_F    | TAGTTGGAATGGGTCCCAAAAAAGTGAACAGA                                       |
| IQD6/AT2G26180_R    | TTATGGAATTGGAATGGTTTGCACCTCCATCTGCCTGGGTAGTGA                           |
| IQD7/AT1G17480_F    | TAGTTGGAATGGGTGCCAAAACCGACACTAAA                                       |
| IQD7/AT1G17480_R    | TTATGGAATTGGAATGGGTCCACCTCGGCTTGCTTTGG                               |
| IQD8/AT1G72670_F    | TAGTTGGAATGGGTCCAAAAGGAAATAATGGGAGTCTG                                 |
| IQD8/AT1G72670_R    | TTATGGAATTGGAATGGGTCCACCTACGGCTTTTG                                 |
| mNeonGreen-SpvB _F  | TTCTAGTTGGAATGGGTATGGTGAGCGAAAAGGCGAG                                 |
| mNeonGreen-SpvB _R  | AGATCCACACATCCGCCCTACACGTCCATG                                       |
| SpvB_F              | CCGATCAGGTGGATCTGGAGGCGATGGAGGGTAGTAATTCATCT                            |
| SpvB_R              | ATCCTATGGAATTGGTACTATTCATAGTGAGTAACCTC                                |
| PHS1ΔP _F           | TTCTAGTTGGAATGGTATGGTGACTAGTGGCAGG                                    |
| PHS1ΔP _R           | CAGATCCACCTTCATCGCCATACGGAGCAGCTTGTAATC                               |
| PHS1ΔP-mNeonGreen_F | GATCAGGTGGATCTGGAGGCATGGTGAGCAAGGGCGAG                                 |
| PHS1ΔP-mNeonGreen_R | GATCCTTATGGAATTGGTACCTTTGACAGCTCGTCC                                  |

**Table S2:** Oligonucleotides used in this study are listed, related to STAR Methods.
Supplemental references:

S1. Schlereth, A., Möller, B., Liu, W., Kientz, M., Flipse, J., Rademacher, E.H., Schmid, M., Jürgens, G., and Weijers, D. (2010). MONOPTEROS controls embryonic root initiation by regulating a mobile transcription factor. Nature 464, 913–917.

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