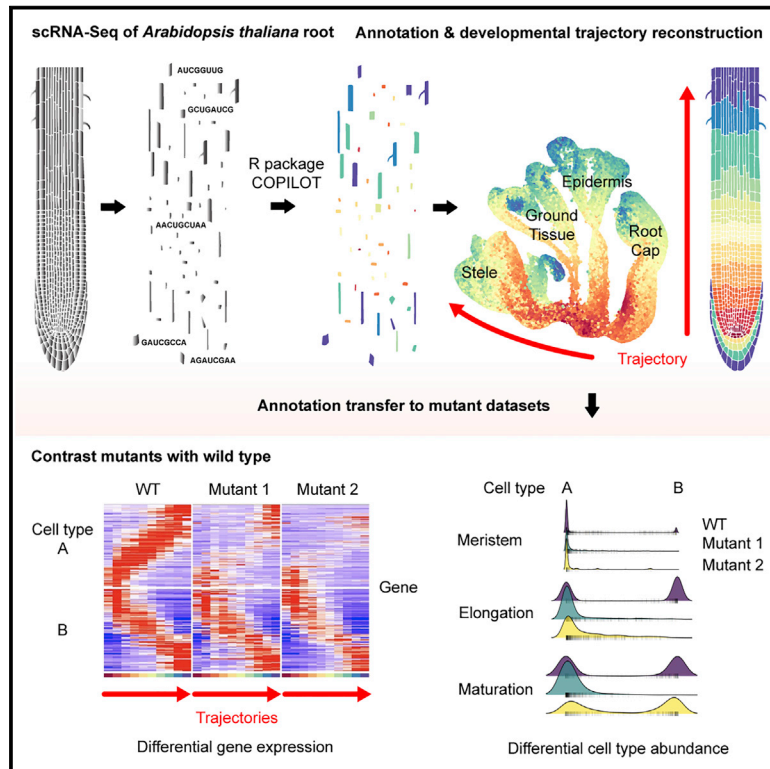


Developmental Cell

A single-cell *Arabidopsis* root atlas reveals developmental trajectories in wild-type and cell identity mutants

Graphical abstract



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In brief

How do transcriptional networks regulate organ development? Using scRNA-seq, Shahan and Hsu et al. produced an *Arabidopsis* root atlas, revealing gradual gene expression changes underlying differentiation of cell types and candidate regulators of cell fate. The atlas enabled interpretation of smaller scRNA-seq datasets and revealed new phenotypes in developmental mutants.

Highlights

- scRNA-seq of >110,000 cells produced a comprehensive *Arabidopsis* root atlas
- Gradual, overlapping waves of gene expression underlie development of all cell types
- Developmental trajectories enable visualization of cell specification events
- scRNA-seq of the *scarecrow* mutant reveals a cell identity change occurs over time



Resource

A single-cell *Arabidopsis* root atlas reveals developmental trajectories in wild-type and cell identity mutants

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SUMMARY

In all multicellular organisms, transcriptional networks orchestrate organ development. The *Arabidopsis* root, with its simple structure and indeterminate growth, is an ideal model for investigating the spatiotemporal transcriptional signatures underlying developmental trajectories. To map gene expression dynamics across root cell types and developmental time, we built a comprehensive, organ-scale atlas at single-cell resolution. In addition to estimating developmental progressions in pseudotime, we employed the mathematical concept of optimal transport to infer developmental trajectories and identify their underlying regulators. To demonstrate the utility of the atlas to interpret new datasets, we profiled mutants for two key transcriptional regulators at single-cell resolution, *shortroot* and *scarecrow*. We report transcriptomic and *in vivo* evidence for tissue trans-differentiation underlying a mixed cell identity phenotype in *scarecrow*. Our results support the atlas as a rich community resource for unraveling the transcriptional programs that specify and maintain cell identity to regulate spatiotemporal organ development.

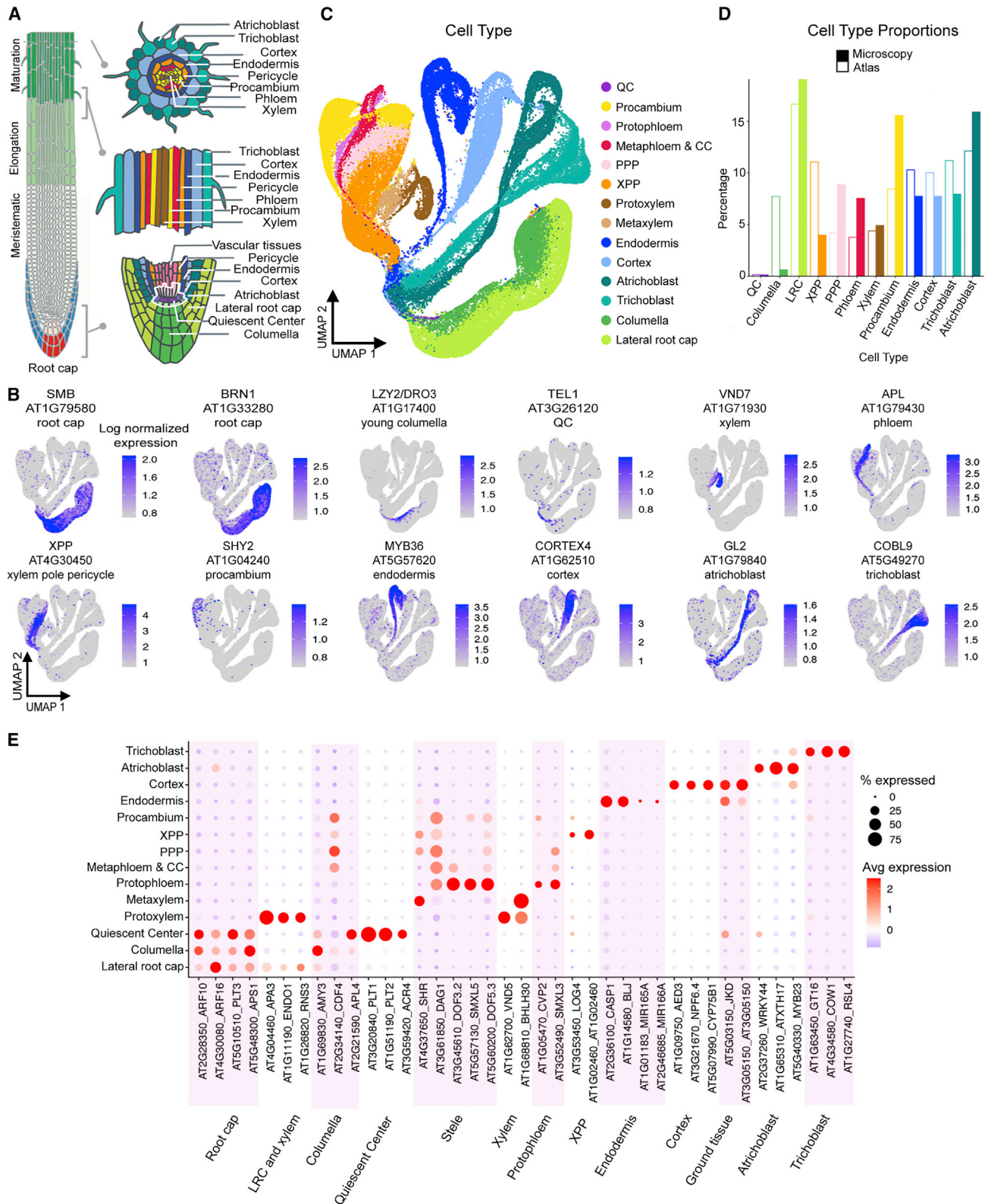
INTRODUCTION

Precisely controlled transcriptional networks specify cell identity, relate positional information, and regulate tissue maturation (Drapek et al., 2017). Defining how these networks orchestrate organ development and function requires detailed knowledge of spatiotemporal gene expression patterns. However, in animal models such as the zebrafish embryo, cells migrate during development and thus present a challenge for cell lineage tracing and subsequent inference of gene expression dynamics (Farrell et al., 2018). The immobile cells and organization of the *Arabidopsis thaliana* root simplify cell lineage tracing and facilitate the study of spatiotemporal organ development (Dolan et al., 1993; Figure 1A). Cell types are arranged in concentric layers around a central vasculature. Cell lineages are ordered longitudinally along a temporal developmental axis, with the oldest cells closest to the shoot and the youngest cells adjacent to the stem cell niche at the root tip. With each new cell division at

the root tip, older cells are displaced shootward from the stem cell niche. Thus, root anatomy simplifies interrogation of the developmental trajectories from stem cell to differentiated tissue (Efroni and Birnbaum, 2016; McFaline-Figueroa et al., 2020).

The *Arabidopsis* root is a tractable model organ with established markers for most cell types as well as expression profiles for morphologically defined developmental stages (Birnbaum et al., 2003; Brady et al., 2007a; Li et al., 2016). Recently, pioneering studies applied droplet-based single-cell RNA sequencing (scRNA-seq) to the *Arabidopsis* root and demonstrated the utility of this technology to identify new cell-type markers, examine gene expression dynamics across pseudotime, and identify regulators that control cell-type-specific responses to environmental conditions (Denyer et al., 2019; Jean-Baptiste et al., 2019; Ryu et al., 2019; Shulse et al., 2019; Zhang et al., 2019; Wendrich et al., 2020). These reports also established foundational principles for root scRNA-seq, including the successful capture of all major cell types from samples





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prepared from whole roots and the utility of known markers and gene expression profiles to accurately annotate major cell types. However, none of these first-generation atlases combined more than 12,500 cells and only [Wendrich et al. \(2020\)](#) inferred developmental progressions for more than three cell types. Further, each atlas is enriched for a subset of cell types or developmental stages at the expense of others ([Figure S1](#)). Thus, there is currently no comprehensive *Arabidopsis* root atlas that captures a finely resolved spectrum of developmental states for all major cell types.

By contrast, recent developmental studies using animal or human samples profiled hundreds of thousands ([Schiebinger et al., 2019](#)) or even millions ([Cao et al., 2019](#)) of cells and high temporal resolution was achieved by densely sampling time points across development ([Briggs et al., 2018](#); [Farrell et al., 2018](#); [Schiebinger et al., 2019](#); [Massri et al., 2021](#)). High pseudotemporal resolution from increased cell numbers provides greater statistical power and enables identification of a finely resolved order of transcriptional events, which is important for considering causal models of gene regulation ([Schiebinger et al., 2019](#)). For the *Arabidopsis* root, in which all cell types are represented at all developmental stages, greater pseudotemporal resolution across development will be gained with an atlas that integrates more cells for all cell types and developmental zones.

Here, we present a primary root gene expression atlas with an order of magnitude more cells than previous *Arabidopsis* datasets. Given the continuous nature of cell states represented in our data, we developed a largely cluster-agnostic annotation approach to avoid bias associated with choosing a clustering resolution. In addition to estimating pseudotime progressions for all cell types, we demonstrate the first application of an optimal transport-based method, StationaryOT, to reconstruct developmental trajectories from plant scRNA-seq data. Cell fate probabilities calculated by StationaryOT shed light on how the fate acquisition of each cell type relates to all other major root cell types. Regressions applied to the cell fate probabilities and gene expression data identified known transcription factors (TFs) involved in cell identity and differentiation. Finally, we tested the ability of the atlas to inform new datasets and demonstrated the power of scRNA-seq to identify new developmental phenotypes by profiling two cell identity mutants, *shortroot* and *scarecrow*.

RESULTS

Integration of over 110,000 cells produces an organ-scale atlas

To build an atlas, we used the 10X Genomics scRNA-seq platform to profile over 96,000 root cells. We harvested 0.5 cm of tissue from 5- to 7-day-old primary root tips across thirteen sets of

independently grown wild-type (WT) seedlings ([Data S1](#)). The transcriptional profiles of all samples were highly correlated, suggesting that batch effects such as differences in plant age are unlikely to substantively affect downstream analyses ([Figure S1](#)). Gene expression matrices calculated by kallisto ([Bray et al., 2016](#)) and bustools ([Melsted et al., 2019](#)) served as input to Cell preproCessing Pipeline kaLlistO busTools (COPILOT), our pre-processing software, which incorporates detection and removal of low-quality cells ([Data S1](#); [STAR Methods](#)).

To add additional depth and assess lab-to-lab data variability, we selected three published root scRNA-seq datasets ([Denyer et al., 2019](#); [Ryu et al., 2019](#)) to combine with data generated in this study ([Data S1](#)). After excluding genes affected by protoplasting (the process of dissociating plant cells from their cell walls; [Denyer et al., 2019](#)) we integrated 110,427 cells into an organ-scale atlas ([Figure S1](#); [Data S1](#); [STAR Methods](#)). A median of 2,768 genes were detected per cell with 24,997 total genes detected, representing 90% of the coding genes in the *Arabidopsis* genome.

Cell annotation places tissues in known developmental contexts

Inspection of marker genes indicated that all major cell and tissue types are discernible as discrete topological features in 2D uniform manifold approximation and projection (UMAP) space ([Figure 1B](#)). To infer precise cell-type annotations, we combined the information from four independent approaches ([STAR Methods](#); [Figures S1–S3](#); [Data S1, S2, and S3](#)) and assigned each cell to one of fourteen cell types ([Figure 1C](#)) and to one of seven developmental stages ([Figure 2A](#)) in a largely cluster-agnostic fashion.

We first mapped cells to 3D root geometry locations ([Schmidt et al., 2014](#)) using novoSpaRc ([Nitzan et al., 2019](#)), an algorithm that reconstructs the locations of single cells in space based on scRNA-seq data ([Data S1](#); [STAR Methods](#)). Second, we used SEMITONES ([Vlot et al., 2020](#)), an algorithm that identifies enriched features in single-cell data without prior clustering, to estimate the enrichment of marker gene expression in cell neighborhoods. Third, we calculated the correlation coefficient of each cell's expression profile to published gene expression profiles of root cell types isolated with fluorescent reporters ([Brady et al., 2007a](#); [Li et al., 2016](#)). Finally, we used an information-theoretic approach to compute index of cell identity (ICI) scores for each cell ([Birbaum and Kussell, 2011](#); [Efroni et al., 2015](#)) ([Data S3](#)). The ICI score is quantitative and represents the relative contribution of cell identities as determined from a reference expression profile dataset. Combining these approaches allowed the expression profile of each cell to inform the boundaries between cell types and developmental stages.

(B) Expression of known cell-type markers. The color scale for each plot represents log normalized, corrected UMI counts for the indicated gene.

(C) UMAP with cell-type labels. The crossing over or apparent mixture between some cell types, e.g., trichoblast and atrichoblast, is a result of 2D projection and absent in 3D ([Video S1](#)).

(D) The proportion of each cell-type group in the atlas is comparable with *in vivo* cell-type proportions ([Cartwright et al., 2009](#)).

(E) Cell-type expression for 40 genes, the spatial expression profiles of which have been previously characterized. Dot size represents the percentage of cells in which each gene is expressed (% expressed). Dot colors indicate the average scaled expression of each gene in each cell-type group with warmer colors indicating higher expression levels.

CC, companion cell; QC, quiescent center; PPP, phloem pole pericycle; XPP, xylem pole pericycle; LRC, lateral root cap. See also [Figures S1–S6](#), [Datasets S1–S3](#), [Data S1](#), and [Video S1](#).

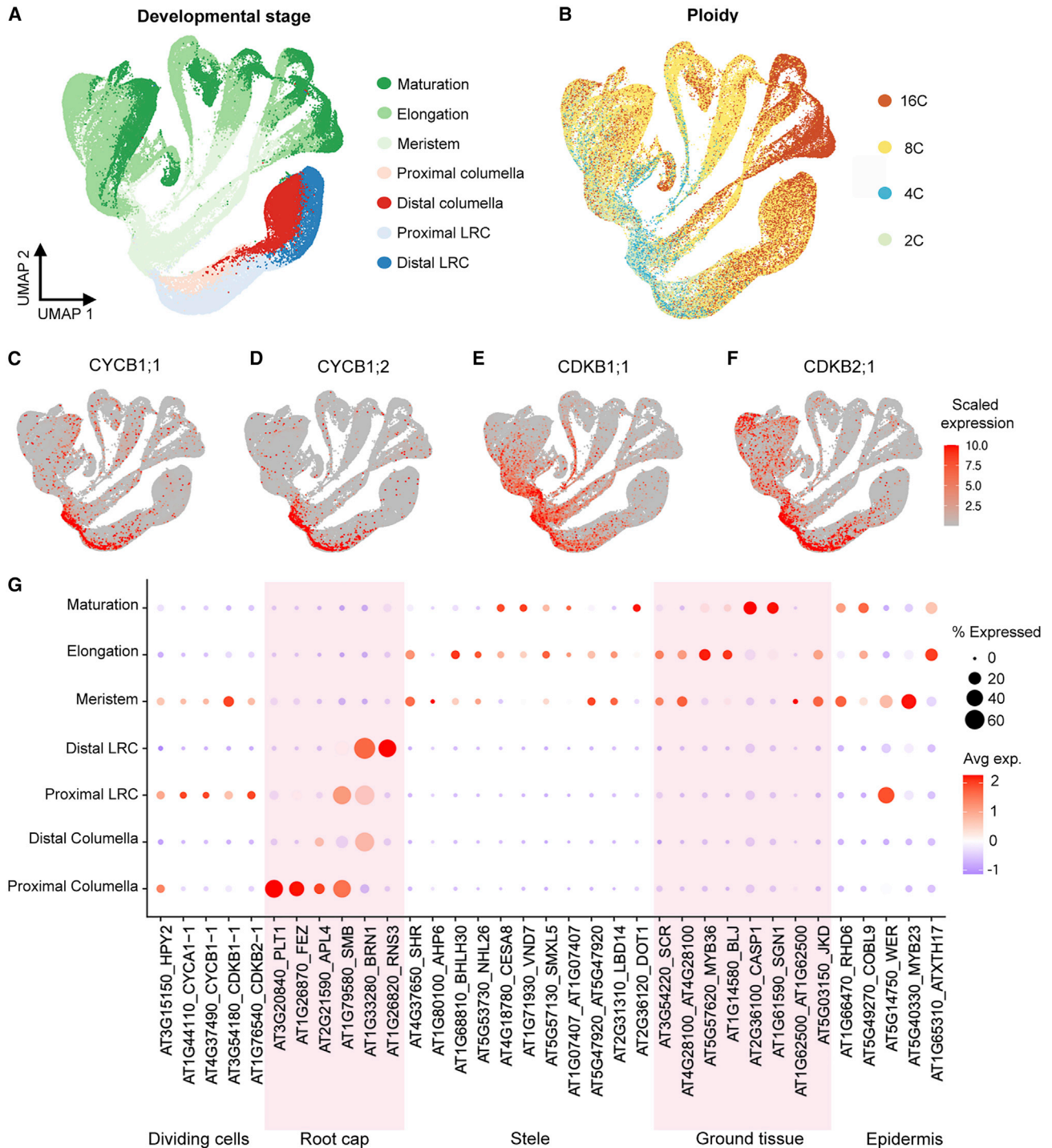


Figure 2. Expression profiles of known genes support the atlas developmental stage annotations

(A) UMAP with developmental stage annotations. LRC, lateral root cap.

(B) UMAP with cell ploidy annotations based on gene expression profiles from [Bhosale et al. \(2018\)](#).

(C–F) Scaled expression ([STAR Methods](#)) of four previously characterized cyclin genes CYCB1;1 (C), CYCB1;2 (D), CDKB1;1 (E), and CDKB2;1 (F) ([Ishida et al., 2009](#)).

(G) Developmental stage expression profiles for 35 genes expressed across the four major root tissue types. Dot size represents the percentage of cells in which each gene is expressed (% expressed). Dot colors indicate the average scaled expression of each gene in each developmental stage group with warmer colors indicating higher expression levels. Root cap: lateral root cap and columella. See also [Figures S1–S6](#) and [Datasets S1 and S2](#).

The resulting atlas ordination consists of cells organized within continuous branches corresponding to four major root tissues (Dolan et al., 1993), each connected to a central group of cells (Figure 1C). Lateral root cap (LRC) and columella cells comprise the root cap and form a single branch. Trichoblast (hair) and atrichoblast (non-hair) cells constitute the epidermis and form a second major branch. Cortex and endodermis cells, which together make up the ground tissue, form a third branch. Finally, the phloem, xylem, procambium, and pericycle cell types are present in the stele tissue and form a fourth branch. Based on marker genes (Data S1), we distinguished additional cell types within the phloem (metaphloem and companion cells; proto-phloem), xylem (protoxylem and metaxylem), and pericycle (xylem pole and phloem pole pericycle). However, we note that fewer validated markers were available for these subtypes. Surprisingly, the ground tissue and epidermis cell types show a clear “sub-branching” topology at the tips of the main branches on the UMAP (Figure 1C). These bifurcations may reflect a developmental phenomenon since they are unlikely to reflect technical artifacts such as differences in protoplasting-induced gene expression signatures (Figure S4).

Overall, atlas cell-type proportions are comparable to both microscopy data (Cartwright et al., 2009; Figure 1D) and previously published root scRNA-seq datasets. Expression profiles of previously characterized genes (not used in the annotation process) also support the accuracy of the annotation (Figure 1E). Differential expression analyses across all cell-type groups (STAR Methods) identified cell-type-specific genes that may be useful for the construction of fluorescent reporter lines (Data S1; Figure S5).

We assigned developmental stage annotations to vascular, epidermal, and ground tissue cell types by comparing each cell transcriptome with gene expression profiles of manually dissected root tissue segments corresponding to meristematic, elongation, and maturation zones (Brady et al., 2007a). Based on these annotations, young cells of the proximal meristem are at the base of each major branch followed by distal meristematic, elongating, and finally mature cells at the tips (Figure 2A). To assign developmental stages to cells in the root cap, we calculated the spatial distance for each cell to the nearest quiescent center (QC) cell using the imputed geometry from novoSpaRc (STAR Methods).

To assess the overall accuracy of the developmental stage annotations, we examined expression patterns of previously characterized genes. First, we annotated the atlas with gene expression profiles associated with DNA endoploidy levels (Figure 2B) (Bhosale et al., 2018; STAR Methods). In agreement with the annotation, the expression of genes associated with increasing ploidy is correlated with increasing maturation. Additionally, the expression of four G2/M phase cell-cycle genes supports the meristematic zone annotation and indicates proximal versus distal root cap cells (Figures 2C–2F). The cyclins CYCB1;1 and CYCB1;2 are expressed in the proximal meristem while CDKB1;1 and CDKB2;1 are expressed in both proximal and distal meristematic cells (Ishida et al., 2009). Lastly, developmental stage expression profiles of known genes agree with published *in vivo* characterizations (Figure 2G).

Overall, the atlas annotations suggest that the combined transcriptome data accurately describe relationships between and

within individual cell types. Similar to previous *Arabidopsis* root atlas UMAP and tSNE plots, older cells from each tissue type radiate from a central group of young cells. However, the integration of greater cell numbers captures more cell states along developmental time and therefore suggests a continuous progression of differentiation for all major root cell types.

Developmental progression can be inferred across individual tissue types

To analyze developmental progression in more detail, we started from a simple pseudotime analysis within annotated cell lineages. We subdivided the atlas into four tissue/lineage groups based on the stem cell of origin (Dolan et al., 1993) and quantified cell state progression using two methodologically distinct, non-graph-based tools: CytoTRACE (Gulati et al., 2020) and scVelo (Bergen et al., 2020). CytoTRACE uses gene diversity to estimate pseudotime while scVelo is based on the concept of RNA velocity. The results from both methods were strongly correlated (Data S4; STAR Methods), suggesting that they reflect true biological signal. We therefore averaged the pseudotime estimations into a “consensus pseudotime” annotation for each tissue (Figures 3 and 4). Overall, the pseudotime estimations reflect biological knowledge. For example, the consensus time annotation for the ground tissue corresponds with the developmental stage annotation and with expression of known endodermis and cortex markers (Figures 3A–3F). As expected, given the 0.5 cm length of harvested root tissue, scaled expression (STAR Methods) of *SCARECROW* (*SCR*), *MYB36*, and *CASPARIAN STRIP MEMBRANE DOMAIN PROTEIN 1* (*CASP1*) represent markers for endodermis cells spanning the meristematic zone to early maturation zone. Expression of *JACKDAW* (*JKD*), a ground tissue marker, as well as cortex-specific markers *CORTEX* (AT1G09750) and *NPF6.4* (AT3G21670) also match the expected profiles. Examples of newly identified genes with expression profiles specific to a subset of the developmental progression are shown for cortex (Figure 3F). Differential expression analyses generated by partitioning the pseudotime ordering into ten groups (T0 to T9) identified a gradual progression of genes dynamically expressed during cortex and endodermis differentiation (Data S4), including previously characterized developmental regulators (Figure 3G).

Similarly, differential expression analyses across ten pseudotime bins show gradual, overlapping waves of gene expression for stele, epidermis + LRC, and columella cells (Figure 4). In agreement with previous work on the root meristem (Wendrich et al., 2017), these results suggest that gradual changes in gene expression also underlie differentiation in the elongation and maturation zones. Gradual, overlapping gene expression dynamics across development are also supported by a dearth of cell-type-specific markers specific to a particular developmental zone (Figure S6). Interestingly, there are two distinct groups of genes along the columella pseudotime progression (Figure 4J), consistent with a rapid change in transcription that could reflect the differentiation of cells immediately after stem cell division (Hong et al., 2015). Also of interest was the lack of pericycle cells in the two most mature pseudotime bins (Figure 4J). This agrees with previous observations that the pericycle matures more slowly than other cell types and retains meristematic characteristics (Beekman and De Smet, 2014).

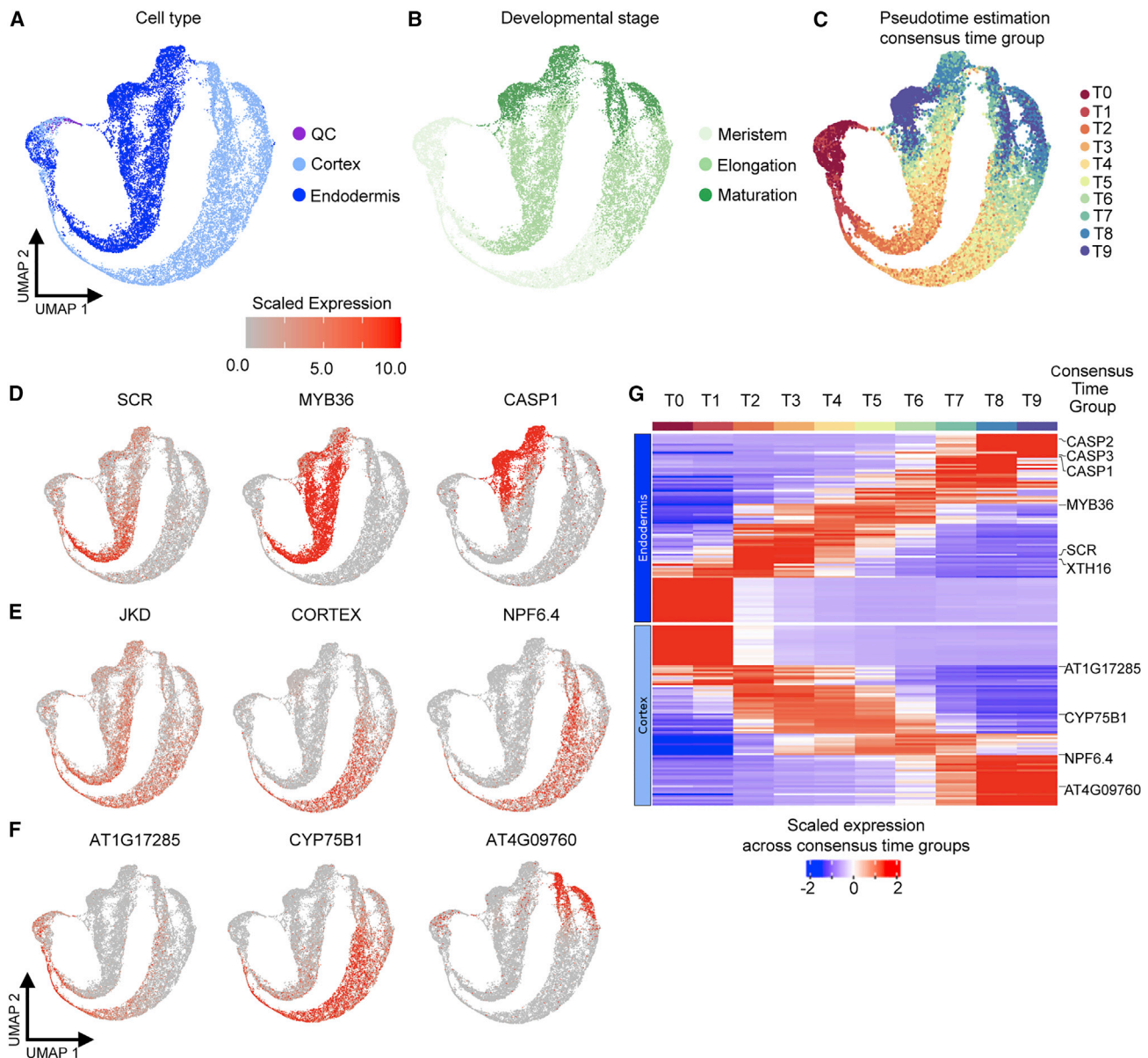


Figure 3. Pseudotime estimates reflect the dynamics of ground tissue differentiation

(A) Endodermis and cortex-annotated cells (ground tissue) were extracted from the atlas and re-embedded in a 2D UMAP. QC cells were included to help anchor pseudotime estimations.

(B) Ground tissue cells annotated with developmental stage labels.

(C) Ground tissue cells annotated with consensus pseudotime group labels. T0 denotes the youngest cells.

(D) Scaled expression patterns (STAR Methods) of known endodermis markers.

(E) Scaled expression of known ground tissue (JKD) and cortex markers.

(F) Newly identified cortex-expressed genes are candidates for marker development.

(G) Scaled expression of 90 and 94 non-redundant, differentially expressed genes across consensus pseudotime groups for cortex and endodermis, respectively. Warmer colors denote higher expression. Although thousands of differentially expressed genes were identified across pseudotime, only the most strongly differentially expressed genes for each of the ten pseudotime bins were plotted for simplicity. See also Data S4.

Optimal transport analysis identifies developmental trajectories

Because pseudotime inference indicates that root cell types mature at different rates, we used an optimal transport (OT)-based method to infer developmental trajectories across the entire atlas.

This method was initially developed as a way to move large quantities of earth with minimal work (Monge, 1781). More recently, OT was used to infer developmental trajectories from animal and human scRNA-seq data (Schiebinger et al., 2019; Marjanovic et al., 2020; Massri et al., 2021) but has yet to be applied to plant data.

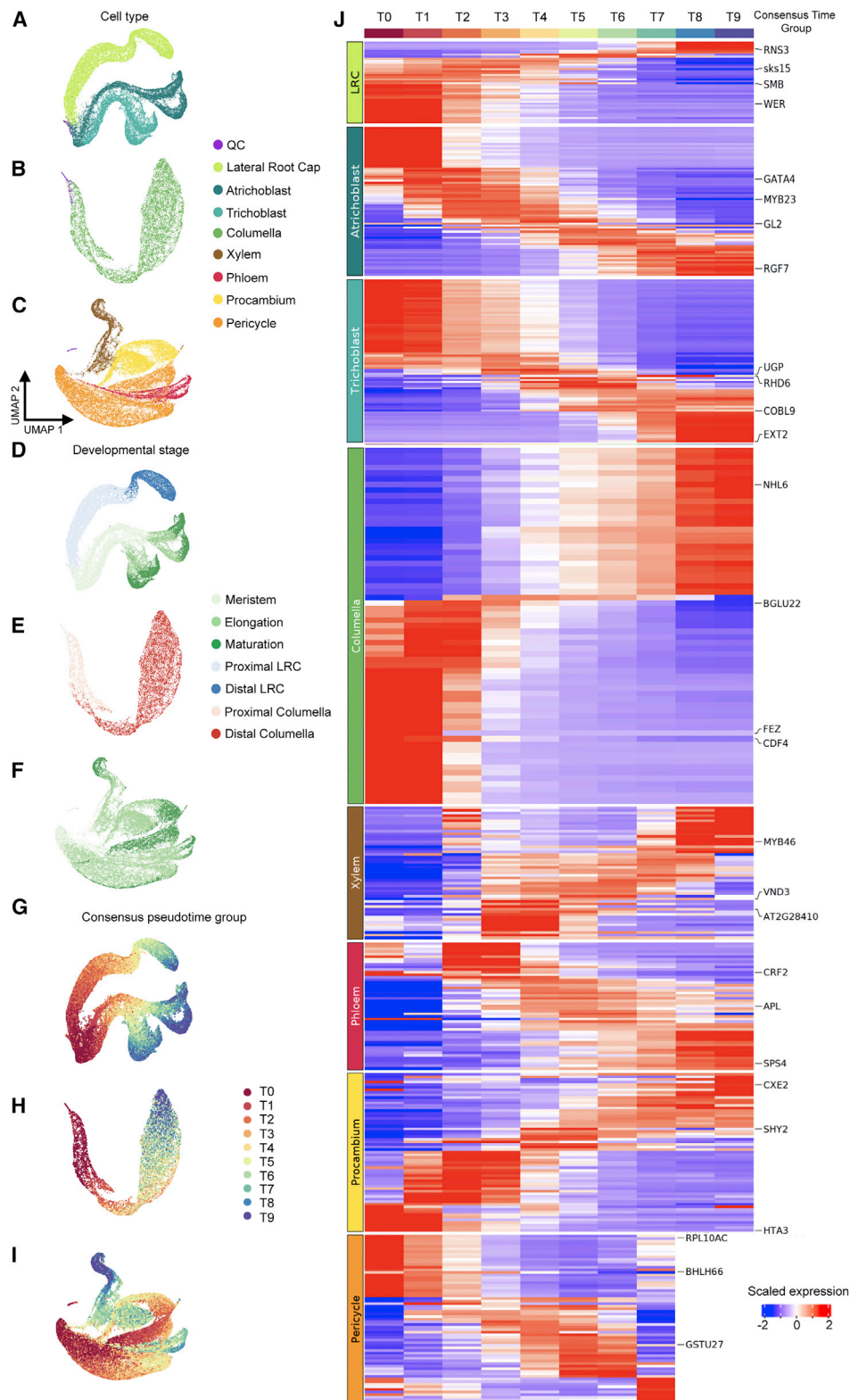


Figure 4. Pseudotime progressions indicate gradual gene expression changes underlie development across tissues and developmental zones

(A–C) Cells annotated as trichoblast, atrichoblast, and lateral root cap (A), columella (B), and stele (C) were extracted from the atlas and re-embedded in individual UMAPs.

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OT connects cells from one static snapshot to their putative ancestors at earlier developmental stages and to their descendants at later developmental stages (Zhang et al., 2021). This allows developmental trajectories of individual cells or entire lineages to be followed through pseudotime. Compared with separating cells by lineage annotation and analyzing pseudotemporal trends within each lineage, OT allows us to probe further back in earlier pseudotime, where lineage-annotations are less reliable. Consequently, this allows us to analyze fate specification events.

The 0.5 cm portion of the root harvested for the atlas can be thought of as a system in equilibrium: cell divisions in the meristem create new cells, which are balanced against the flux of cells exiting this region. We applied stationary OT analysis (StationaryOT, Figure 5A) (Zhang et al., 2021), which leverages estimates of cellular growth rates to infer trajectories for systems in equilibrium. We used the consensus pseudotime to define groups of cells that represent terminal destinations (i.e., the “fates”; STAR Methods) and estimated growth rates for individual cell types based on time-lapse imaging data of dividing cells (Rahni and Birnbaum, 2019). Using these parameters, StationaryOT calculates a vector of fate probabilities for each cell in the atlas, i.e., the likelihood that a given cell will eventually give rise to a mature cell of a particular cell type. Individual fate probabilities can be visualized on the atlas UMAP coordinates and agree with our cell-type annotations, as shown for endodermis (Figure 5B). The maximum fate probability for each cell, which indicates the most likely ultimate cell lineage, agrees with our lineage-annotations (Figures 5C and S7), and cells appear to gradually become more biased toward specific fates at later pseudotimes (Figure 5C). Taken together, these results suggest that the developmental trajectories inferred by StationaryOT, which are largely independent of the atlas annotations and do not require segmentation of the atlas into constituent lineages, reflect existing biological knowledge for differentiation of each cell and tissue type.

Differentiation events can be visualized by projecting multiple fate probabilities in barycentric coordinates as “triangle plots” (STAR Methods). Contrary to pseudotime inference methods, which are applied to individual tissues or cell lineages, these visualizations can be used to interrogate how fate acquisition of each cell type relates to all other cell types. To explore the divergence of endodermis and cortex identities, we designated a vertex of the triangle for each of these fates with the third vertex representing all other possible fates. Cells were then plotted according to their relative probabilities. The position of meristematic cells in the triangle interior indicates lower cortex or endodermis fate probabilities at earlier developmental stages (Figure 5D). Mature cells are grouped at the cortex and endodermis vertices, which indicate 100% cortex or endodermal fate probabilities, respectively (Figure 5F; STAR Methods). Plotting the expression of known endodermis markers indicates that endodermis fate probabilities increase with maturation as expected (Figure 5G). Interestingly, in the elongation zone, endodermis cells are already strongly fated while cortex fate appears indeterminate (Figure 5E). This could reflect the putative “ground

state” of the cortex for which the ground tissue was named and suggests that elongating cortex cells have the potential to acquire different fates (Esau, 1953; Cui, 2015).

In another example, the fate probabilities for trichoblast and atrichoblast, which together form the epidermis, are more similar to each other than either is to LRC cells, although all derive from the same stem cell (Figure 5H). Similarly, columella root cap cell fates are distinct from all other fates except LRC (Figure 5I). Plotting cells by developmental zone annotations (Figures 5J–5L) indicates that atrichoblast and trichoblast cell fates are indeterminate in the meristem with some fluidity in the elongation zone, which agrees with previous observations that epidermal cell fate is not fixed in young cells (Berger et al., 1998a; Ryu et al., 2019).

For stele cell types, plotting cells according to fate probabilities reflects the distinct identities of xylem and phloem, both compared with each other and with procambium and pericycle cells (Figures 5M and 5N). This is visualized on tetrahedron plots by, for example, the concentration of xylem cells on the side of the triangle between the “other” and xylem vertices, indicating that the cells have higher fate probabilities for xylem than for phloem or any other cell type (Figure 5N). By contrast, procambium and pericycle fates appear to be fluid (Figures 5M, 5O–5Q), similar to the fluidity between atrichoblast and trichoblast fates.

OT analysis facilitates identification of developmental regulators

To identify TFs with expression patterns predictive of fate specification probabilities for each cell type, we applied L1-regularized linear regression (i.e., the Lasso) (Data S5; Figure S7). Among top ranked genes were numerous known regulators with positive coefficients, indicating a positive influence on a given cell lineage (STAR Methods; Data S5). Examples include: (1) *MYB36* and *SCR* for meristematic and elongation endodermis, (2) *JKD* for meristematic and elongation cortex, (3) *GLABRA2*, *MYB23*, and *CAPRICE* for meristematic and elongation atrichoblast, and (4) *RHD6* (*BHLH83*) for meristematic trichoblast. The re-discovery of known regulators for all major root cell types as top candidates supports the utility of the atlas itself as well as the StationaryOT approach to identify TFs with key roles in cell fate specification. Many of the genes identified by the regressions are unstudied and represent a rich resource for functional characterization.

scRNA-seq reveals differentiation pathways of cell identity mutants

In addition to identifying new candidate regulators, scRNA-seq allows us to ask how known regulators control tissue and organ development. In the root, the TFs *SHORTROOT* (*SHR*) and *SCR* function in a transcriptional regulatory complex and are essential for stem cell niche maintenance and tissue patterning (Benfey et al., 1993; Di Laurenzio et al., 1996). Using annotation label transfer from the atlas to inform new datasets (Stuart et al.,

(D–F) UMAPs for epidermis plus lateral root cap (D), columella (E), and stele (F) annotated by developmental stage.

(G–I) UMAPs for epidermis plus lateral root cap (G), columella (H), and stele (I) annotated by consensus time groups.

(J) Scaled expression of the top ten non-redundant, most highly differentially expressed genes across consensus pseudotime groups for each cell type. See also Data S4.

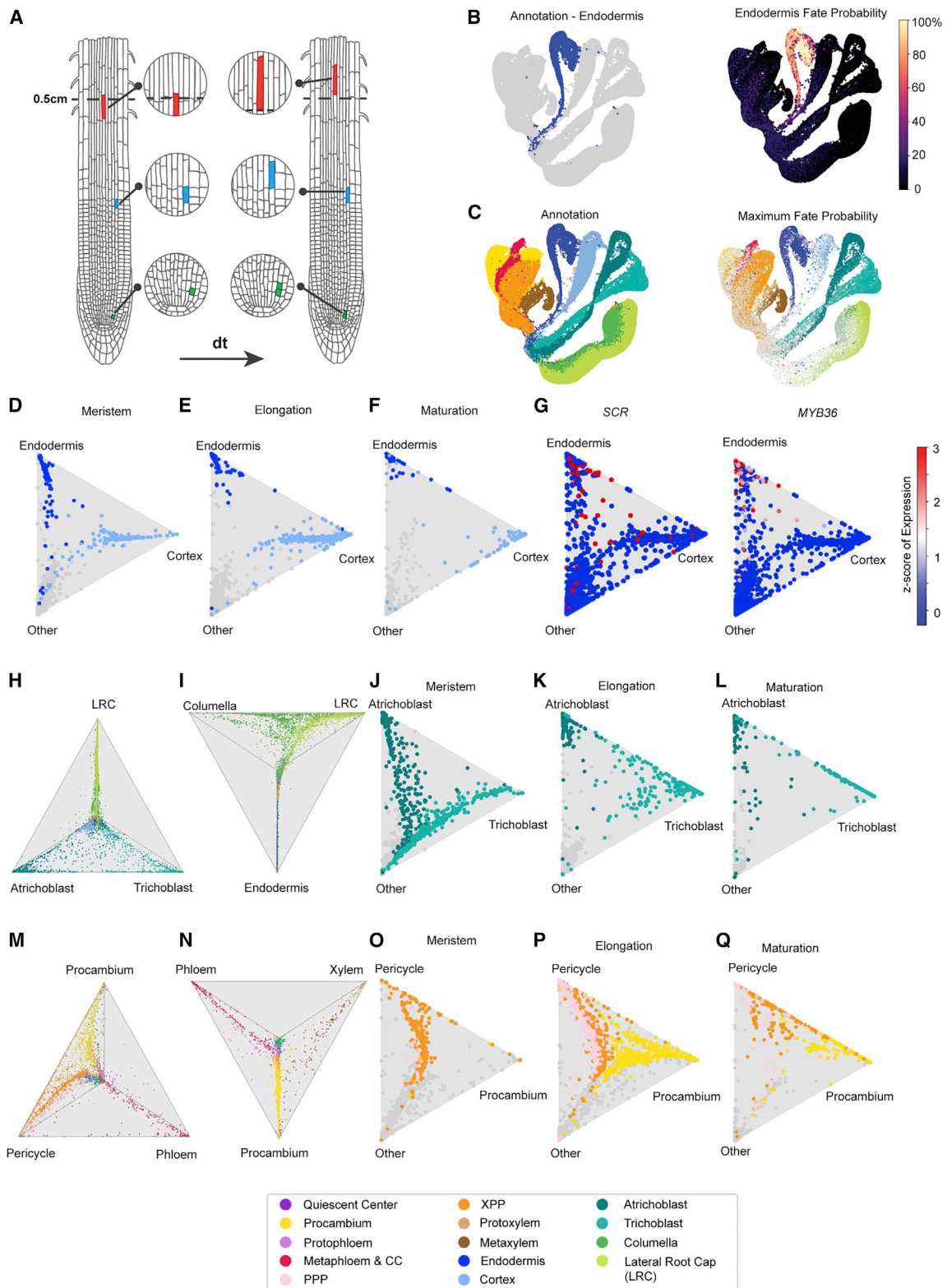


Figure 5. Optimal transport identifies developmental trajectories

(A) The root tip, denoted here as the 0.5 cm harvested for scRNA-seq, remains in equilibrium over a time period of duration dt . Individual cells progress through developmental stages, including dividing (green, transit amplifying divisions following stem cell divisions), enlarging (blue, elongation zone), and exiting the region of interest (red, early maturation zone).

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2019), we asked how the loss of *SHR* or *SCR* function affects tissue composition as well as cell identity and differentiation.

Both *shr* and *scr* mutants lack the asymmetric cell division that patterns the ground tissue, resulting in a single mutant tissue layer instead of the cortex and endodermis cell layers. Previous detection of tissue-specific markers and morphologies revealed that the mutant layer has cortex-like attributes in *shr* (Benfey et al., 1993) but a mixture of cortex and endodermis characteristics in *scr* (Di Laurenzio et al., 1996). These phenotypes were reflected in the scRNA-seq data given the significant reduction of cells expressing endodermal markers in both *shr* and *scr* (Figures 6A–6C). A second striking observation was the decrease in protoxylem cell abundance in both mutants and the decrease of protophloem and metaphloem abundance in *shr* (Figure 6C), consistent with reports of defects in *shr* and *scr* stele development (Levesque et al., 2006; Carlsbecker et al., 2010; Yu et al., 2010; Cui et al., 2011; Kim et al., 2020). In both mutants, we also identified a significant reduction in the abundance of xylem pole and phloem pole pericycle cells (Figure 6C). This is surprising given that there are cells located in the radial pericycle position in both mutants (Kim et al., 2020; Di Laurenzio et al., 1996). However, in *Arabidopsis*, lateral roots are formed from xylem pole pericycle cells (Beekman and De Smet, 2014), and lateral root development is altered in the *shr* mutant (Lucas et al., 2011). This observation previously led to the hypothesis that *shr* cells may differentiate into a state that cannot support normal lateral root formation (Lucas et al., 2011). Taken together, these results indicate a putative loss of pericycle identity in *shr*.

scRNA-seq suggests trans-differentiation of the *scr* mutant layer

We next asked how individual cells contribute to the reported mixed identity of the *scr* mutant layer (Di Laurenzio et al., 1996). One hypothesis is that cells acquire an endodermis or cortex identity early in development and the mutant layer is a heterogeneous mixture of the two cell types along the entire cell file. Alternatively, each cell may have a mixture of cortex and endodermis attributes. A third hypothesis is that cells acquire one identity early in development and subsequently change their fate. To distinguish among these possibilities, we used StationaryOT to calculate *scr* cell fate probabilities. *scr* cortex and endodermis cells exist on a continuum between cortex

and endodermis fates, as indicated by the cells aligned on the side of the triangle plot between the cortex and endodermis vertices (Figure 6D). This reflects the probabilities of both endodermis and cortex fates for these cells. In the *shr* dataset, although some cells are annotated as endodermis, the lack of cells near the endodermis vertex coupled with low confidence scores following label transfer via Seurat (Figure 6E) suggests that few if any *shr* cells are endodermis-like. Similar to *shr* but unlike WT, *scr* endodermis cells do not show a progression from the central part of the triangle toward the endodermis vertex. This suggests that *scr* cells may not gradually acquire endodermis identity from an undifferentiated state.

To further explore the developmental progression of the *scr* mutant layer, we extracted cortex and endodermis-annotated cells from the *scr* dataset. We asked if the proportion of cells with each cell-type annotation changes according to developmental zone. We observed that most meristematic and elongating *scr* cells are confidently classified as cortex, though a subset of cells with low cortex prediction scores is evident in the elongation zone (Figure 6F). Differentiating *scr* cells, however, are confidently annotated as either cortex or endodermis, though some cells seem to have attributes of both. By contrast, nearly all *shr* mutant layer cells are confidently annotated as cortex (Figure 6F). In agreement with these results, consensus pseudotime annotation labels transferred from the atlas suggest that the youngest cells of the *scr* mutant layer are primarily cortex-like while endodermis identity is most evident in older cells (Figures 7A–7C). By contrast, cortex identity is predominant in all developmental states for *shr* mutant layer cells (Figures 7D–7F). Together, these results support the hypothesis that *scr* mutant layer cells are cortex-like in the early stages of development but acquire attributes of endodermal identity as they age.

To test our hypothesis *in vivo*, we asked if spatial expression patterns of known cortex and endodermis markers are altered in the *scr* mutant layer. In WT roots, transcriptional reporters for MYB36, an endodermis marker, and AT1G09750 (CORTEX), a cortex marker, are expressed in the elongation zones of their respective cell types (Figures 3D and 3E; Liberman et al., 2015; Lee et al., 2006). The MYB36 reporter is also expressed in the meristematic zone of the endodermis. However, scRNA-seq data suggest that these expression patterns are altered in the *scr* mutant layer: CORTEX expression is reduced to the early

(B) Endodermis fate probability (right) agrees with endodermis annotations (left), visualized on the UMAP.

(C) All fate probabilities are visualized on the UMAP (right). Cells are colored according to the lineage of maximum fate probability and cells fade to gray as the fate specification becomes less determined (i.e., as the maximum fate probability decreases).

(D–Q) StationaryOT fate probabilities reflect known developmental relationships and, in some cases, fate fluidity between cell types. For each plot, the dataset was down-sampled to 10,000 cells to facilitate visualization.

(D–F) Triangle plots with cells plotted according to cortex, endodermis, and all other fate probabilities. Cells annotated as cortex and endodermis are colored light and dark blue, respectively, with all other cells in gray. The three plots show cells from meristem (D), elongation (E), and maturation (F) developmental zones.

(G) Increasing endodermis fate probabilities agree with developmental stage annotations and with expression patterns of *SCARECROW* (*SCR*) and *MYB36*. The legend shows z scores of gene expression, where a score of 1 is one standard deviation above mean expression.

(H and I) Cells are arranged on tetrahedron plots according to cell fate probabilities for atrichoblast, trichoblast, and LRC (H) and columella, LRC, and endodermis (I). The top vertex of each face of the tetrahedron plots (looking down) contains all other cell-type fates besides the three labeled at each of the remaining vertices.

(J–L) Cells are plotted according to atrichoblast, trichoblast, and all other fate probabilities. Cells annotated as atrichoblast and trichoblast are colored accordingly with all other cells in gray. The three plots show cells from meristem (J), elongation (K), and maturation (L) developmental zones.

(M and N) Tetrahedron plots representing procambium, pericycle, and phloem (M) and phloem, xylem, and procambium (N) cell fate probabilities. Xylem, phloem, and pericycle terminal fates from pseudotime estimates were used for StationaryOT, but stele cells are colored here according to annotated subtypes.

(O–Q) Cells are plotted according to pericycle, procambium, and all other fate probabilities. The three plots show cells from meristem (O), elongation (P), and maturation (Q) developmental zones. See also Figure S7.

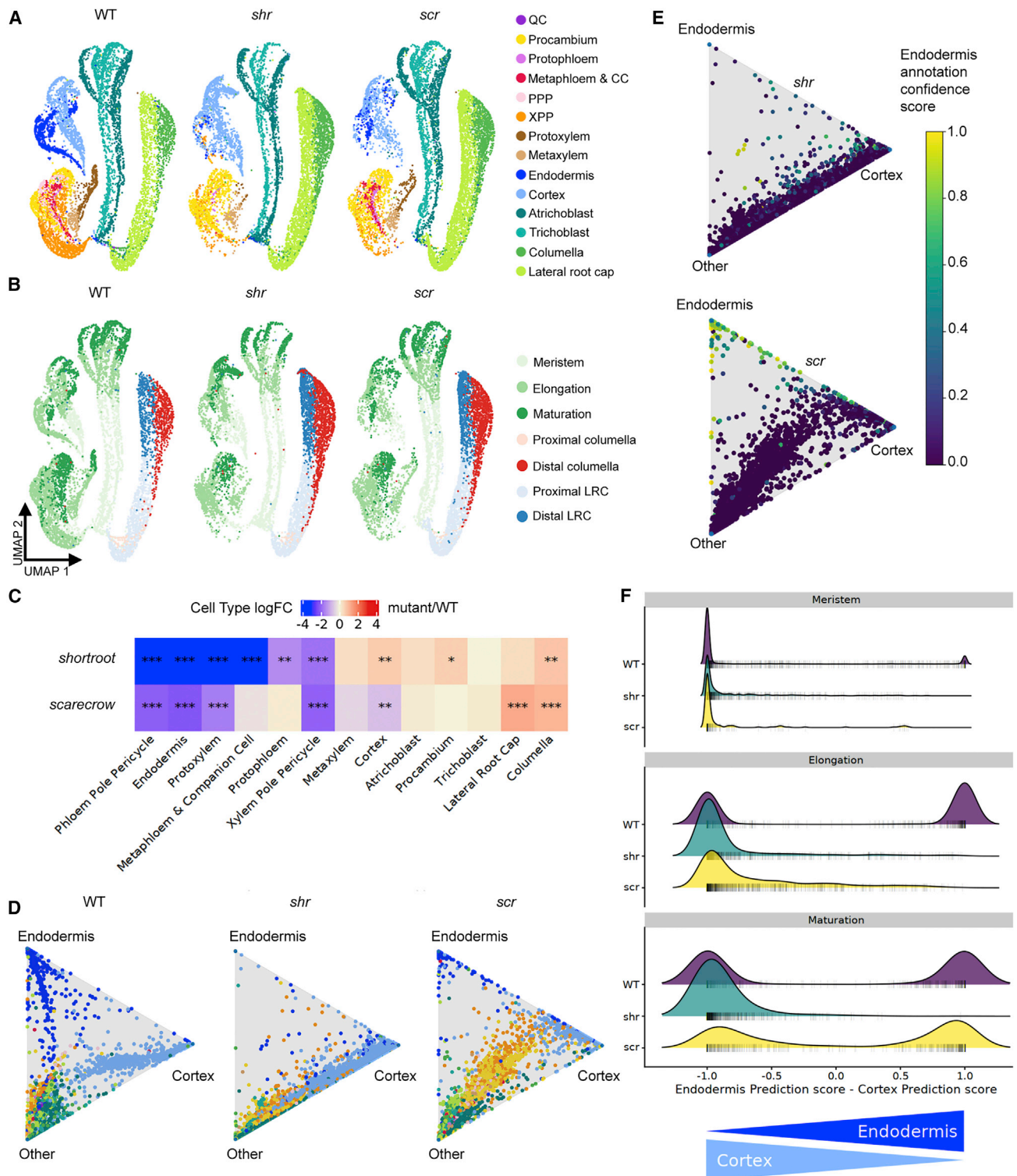


Figure 6. Atlas informs cell-type abundance and identity changes in *shr* and *scr* mutants

(A) UMAPs with cell-type annotations representing WT integrated with *shr* and *scr*. Data from each genotype were down-sampled to 10,000 cells to facilitate comparison.

(B) UMAPs from A but labeled with developmental stage annotation.

(C) Differential abundance analysis using the full integrated WT, *shr*, and *scr* dataset reports significant changes in per label cell-type abundance between mutants and WT. ***False discovery rate (FDR) < 0.001; **FDR < 0.01; *FDR < 0.05.

(D) Triangle plots illustrating cell fate probabilities calculated by StationaryOT. Cell-type color legend is the same as (A).

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elongation zone while MYB36 is expressed in older cells of the elongation and maturation zones but not in the meristem (Figures 7G and 7H).

Consistent with the scRNA-seq observations, expression from a MYB36 transcriptional reporter was visible only in the late elongation and maturation zones of the *scr* mutant layer while signal from a CORTEX transcriptional reporter was diminished in the elongation and maturation zones (Figures 7I–7P). Additionally, a transcriptional reporter for the meristematic cortex, CO2, was previously shown to be robustly expressed in young mutant layer cells closest to the QC in *scr-4* (Heidstra et al., 2004). Taken together, the *in vivo* expression patterns of MYB36, CORTEX, and CO2 reporters validate developmental observations made from scRNA-seq data and suggest that young *scr* mutant layer cells are cortex-like while the identity of older cells changes to more endodermis-like. Although endodermal identity has been considered independent of *SCR* and the existence of *SCR*-independent regulation of MYB36 has previously been proposed (Drapek et al., 2018), our results indicate that *SCR* is required in meristematic and early elongation cells for MYB36 expression and endodermal identity.

DISCUSSION

Observations made from WT and mutant data lay the foundation to address fundamental questions regarding common versus shared developmental regulatory programs between cell types, cell identity transitions, and the roles of neighboring cells in determining cell identity. Building organ-scale gene expression maps is also essential to drive technological innovation such as reprogramming cell identity and inducing phenotypic changes via cell-type-specific gene editing. To address these goals, we built a comprehensive root scRNA-seq atlas, developed an iterative pipeline to annotate each cell individually, and developed COPILOT, a species-agnostic quality control software for scRNA-seq data. An interactive web interface is available for the atlas at <https://phytozome-next.jgi.doe.gov/tools/scrna/>.

The resolution of developmental progression represented in the atlas provides an opportunity to ask how cell fate specification and stabilization differ between cell types, especially those that arise from divisions of the same stem cell. For example, genes uniquely expressed in the cortex or endodermis early in development for each cell type may include new regulators of cell-type specification in the ground tissue. Further, StationaryOT allows insight into transcriptional similarities across cell-type fate specification. An intriguing question is how and why cell types that arise from the same stem cell, such as procambium and phloem, are more transcriptionally distinct than cell types which arise from different stem cells, e.g., procambium and pericycle.

Transcriptional regulators of tissue patterning, cell identity specification, and differentiation have previously been identified for each root tissue. However, we have by no means discovered all regulators, and we have limited understanding of what connects known gene regulatory networks (GRNs) operating at different developmental stages in individual cell types (Drapek et al., 2017). The regression we applied to StationaryOT and gene expression data identified a number of uncharacterized genes as candidate regulators of cell fate. Although we highlighted candidates predicted to push cells toward a given lineage, the analysis also identified genes for each cell type that do not favor the lineage. These genes will be interesting to perturb and test for phenotypes with approaches such as cell-type-specific overexpression. Given the applicability of StationaryOT to the full atlas, the candidates may also include TFs that coordinate developmental processes across cell and tissue types.

For future studies, the atlas represents a rich resource to infer GRNs underlying the differentiation of each cell type with tools such as CellOracle (Kamimoto et al., 2020). The atlas data can also be compared or combined with data from other modalities to examine gene regulatory relationships and narrow down the candidate TFs that regulate cell fate decisions. For example, GRNs inferred from the atlas data could be compared with DAP-seq data (O'Malley et al., 2016) to determine if TFs of interest bind to regulatory regions of predicted downstream genes. Another promising avenue to identify transcriptional regulators controlling cell fate and differentiation is the combination of chromatin accessibility (scATAC-seq) and scRNA-seq data (Stuart and Satija, 2019; Rautenstrauch et al., 2022), the feasibility of which has been demonstrated for *Arabidopsis* and rice roots (Dorrity et al., 2021; Farmer et al., 2021; Zhang et al., 2021).

Beyond WT root development, the atlas enables interrogation of cell identity and tissue composition changes in a mutant context. The putative trans-differentiation from cortex to endodermis identity in the *scr* mutant layer represents a new system with which to investigate transcriptional changes underlying cell identity transitions. In regeneration studies, plant cells show a widespread ability to acquire new fates (Efroni, 2018), which raises questions such as how do cells “forget” their old fate and are there unstable transitional states required for identity transitions? To date, there are few transcriptome-level datasets describing cell identity changes in plants, although such transitions represent important developmental processes including pericycle cells undergoing identity changes during lateral root formation (Wangenheim et al., 2016; Gala et al., 2021). The *scr* scRNA-seq data will allow us to probe questions about these transitional states, such as do cells express heterogeneous mixtures of cortex and endodermal identity and do cortex cells “de-differentiate” prior to expression of endodermal markers?

(E) Triangle plots show cells arranged according to endodermis, cortex, and all other fate probabilities for *shr* (top) and *scr* (bottom) as calculated by StationaryOT. Each dot represents one cell. Dots are colored by endodermis annotation confidence scores after label transfer from the WT atlas by Seurat. Zero and one are the lowest and highest confidence scores, respectively.

(F) Data density plot of the cortex classification score subtracted from the endodermis classification score for each cell, plotted by developmental stage. On the x axis, a value of 1 indicates confident endodermal classification while a value of -1 indicates confident cortex classification. The annotation of each *scr* and *shr* cell was assigned using a weighted vote classifier based on reference cell labels from the atlas (Stuart et al., 2019). Cell-type classification scores range from zero (lowest confidence) to one (highest confidence). Absolute cell numbers are represented by the shaded bars. See also [Datasets S1](#) and [S2](#), [Data S1](#).

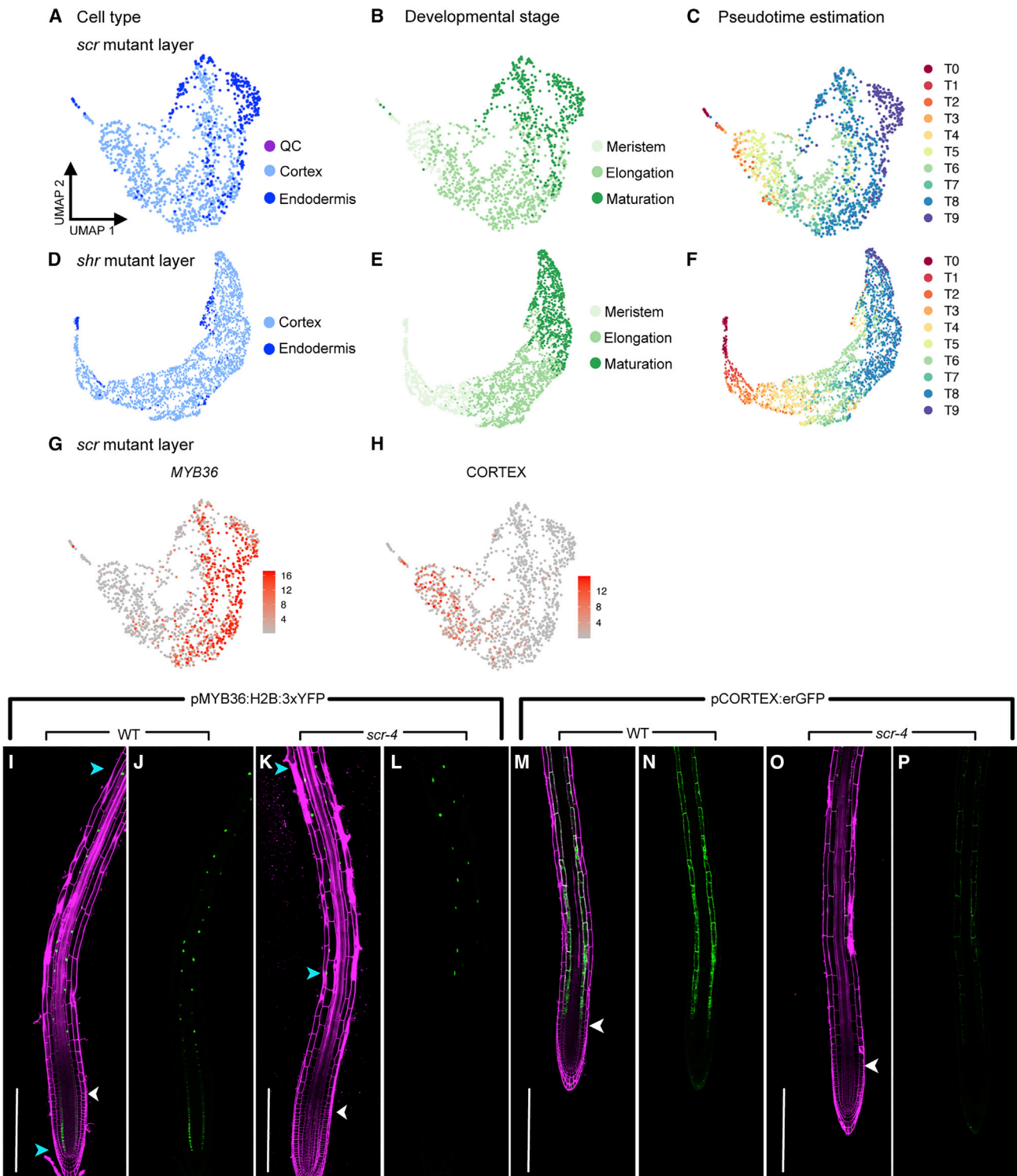


Figure 7. Spatial expression patterns of MYB36 and CORTEX transcriptional reporters are consistent with cortex to endodermis fate transition in the *scarecrow* mutant layer

(A) Cortex, endodermis, and QC cells extracted from the *scr* dataset and re-embedded in a UMAP.

(B) Developmental stage annotation labels were transferred from the WT atlas to the *scr* mutant layer cells.

(C) Consensus pseudotime group annotation labels were transferred from WT ground tissue to *scr*. Warmer to cooler colors represent the developmental progression from youngest to oldest cells, respectively.

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To facilitate the utility of the atlas as a community resource, we produced comprehensive tutorials and toy datasets to demonstrate how the atlas annotation labels can be transferred to new datasets. In addition to analyzing mutants, the atlas can guide interpretation of scRNA-seq data from plants responding to environmental stress, as well as data from crop species for which comprehensive root cell-type markers are unavailable.

Limitations of the study

We relied only on transcriptional profiles to determine a cell's identity and developmental state, which excludes other information such as proteomic profiles. We note that the atlas developmental stage annotation is based on correlation with microarray data from tissue segments hand-dissected according to morphological markers. The boundaries between developmental zones in the atlas may not correlate precisely with root morphology due to variability between roots and between individuals in interpreting the markers.

STAR★METHODS

Detailed methods are provided in the online version of this paper and include the following:

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SUPPLEMENTAL INFORMATION

Supplemental information can be found online at <https://doi.org/10.1016/j.devcel.2022.01.008>.

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AUTHOR CONTRIBUTIONS

R.S., C.-W.H., T.M.N., B.J.C., P.N.B., and U.O. conceptualized the experiments. R.S., T.M.N., and I.W.T. generated the scRNA-seq data. R.S., C.-W.H., T.M.N., B.J.C., A.H.C.V., L.G., S.Z., A.A., and G.S. analyzed the data. R.S. performed confocal imaging. R.S. wrote the manuscript with input from all authors. G.S., P.N.B., and U.O. supervised the experiments and analyses.

DECLARATION OF INTERESTS

P.N.B. is a member of the *Developmental Cell* advisory board and is the co-founder and Chair of the Scientific Advisory Board of Hi Fidelity Genetics, a company that works on crop root growth.

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(D) Cortex and endodermis cells were extracted from the *shr* dataset and re-embedded in a UMAP. (E and F) As for *scr*, developmental stage (E) and consensus time group annotation labels (F) were transferred from the WT atlas to *shr* mutant layer cells. (G and H) Scaled expression of *MYB36* (G) and *AT1G09750* (CORTEX reporter; H) in cells of the *scr* mutant layer. (I–L) pMYB36:H2B:3x Yellow Fluorescent Protein (YFP) reporter in WT (I, J) and *scr-4* (K, L) showing loss of meristem and elongation zone expression in *scr* mutant. Blue arrowheads mark the longitudinal location of the first and last cells in the image with visible YFP. (M–P) pCORTEX:erGFP reporter in *scr-4*/pCORTEX:erGFP F2 progeny with WT ground tissue phenotype (M and N) and *scr-4* mutant layer phenotype (O and P) showing reduced expression of cortex marker as cells mature in the mutant. Red and green channel overlay images (I, K, M, and O) are propidium iodide-stained roots (magenta) and YFP or GFP signal. Green channel images (J, L, N, and P) are YFP or GFP alone. Scale bars, 200 μ m. White arrowheads mark the beginning of the elongation zone. See also [Data S1](#).

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STAR★METHODS

KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---|--|--------------------------|
| Bacterial and virus strains | | |
| N/A | N/A | N/A |
| Biological samples | | |
| <i>Arabidopsis thaliana</i> ecotype Columbia-0 | N/A | N/A |
| <i>Arabidopsis</i> : pCORTEX:erGFP | Lee et al. (2006) | N/A |
| <i>Arabidopsis</i> : pMYB36:H2B:3xYFP | Drapek et al. (2018) | N/A |
| <i>Arabidopsis</i> : scarecrow-4 | Fukaki et al. (1998) | ABRC stock number CS6505 |
| <i>Arabidopsis</i> : shortroot-2 | Levesque et al. (2006) | ABRC stock number CS2972 |
| Chemicals, peptides, and recombinant proteins | | |
| Linsmaier and Skoog medium | Caisson Labs | Product number LSP03-1LT |
| Cellulase ONOZUKA R-10 | GoldBio | Cat#C8001.0005 |
| Pectolyase | Sigma | Cat#P3026 |
| Bovine Serum Albumin | Sigma | Cat#A3912 |
| β-mercaptoethanol | Sigma | Cat#M6250 |
| Mannitol | Sigma | Cat#SLBV3117 |
| MES | Sigma | Cat#6120 |
| KCl | Fisher Scientific | Cat#AM9640G |
| CaCl ₂ | Sigma | Cat#21115 |
| Propidium iodide | Sigma | Cat#P4170 |
| Basta (Glufosinate ammonium) | Fisher Scientific | Cat#J66186-MD |
| Critical commercial assays | | |
| Chromium Single Cell Controller | 10X Genomics | Product Code 120263 |
| Chromium i7 Multiplex Kit | 10X Genomics | Product Code 120262 |
| Chromium Single Cell 3' GEM Library & Gel Bead Kit v3 | 10X Genomics | Product Code 1000092 |
| Chromium Chip B Single Cell Kit | 10X Genomics | Product Code 1000074 |
| DNA High Sensitivity Bioanalyzer Kit | Agilent | Cat#5067-4626 |
| Deposited data | | |
| Single Cell mRNA Sequencing data | This study | GEO: GSE152766 |
| Single Cell RNA-Seq wild type <i>Arabidopsis</i> root cells - sc_1 | This study | GEO: GSM4625993 |
| Single Cell RNA-Seq wild type <i>Arabidopsis</i> root cells - sc_9_at | This study | GEO: GSM4625994 |
| Single Cell RNA-Seq wild type <i>Arabidopsis</i> root cells - sc_10_at | This study | GEO: GSM4625995 |
| Single Cell RNA-Seq wild type <i>Arabidopsis</i> root cells - sc_11 | This study | GEO: GSM4625996 |
| Single Cell RNA-Seq wild type <i>Arabidopsis</i> root cells - sc_12 | This study | GEO: GSM4625997 |
| Single Cell RNA-Seq wild type <i>Arabidopsis</i> root cells - sc_20 | This study | GEO: GSM4625998 |
| Single Cell RNA-Seq wild type <i>Arabidopsis</i> root cells - sc_21 | This study | GEO: GSM4625999 |
| Single Cell RNA-Seq <i>scr</i> mutant <i>Arabidopsis</i> root cells - sc_25 | This study | GEO: GSM4626000 |

(Continued on next page)

Continued

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---|------------|-----------------|
| Single Cell RNA-Seq wild type <i>Arabidopsis</i> root cells - sc_30 | This study | GEO: GSM4626001 |
| Single Cell RNA-Seq wild type <i>Arabidopsis</i> root cells - sc_31 | This study | GEO: GSM4626002 |
| Single Cell RNA-Seq <i>scr</i> mutant <i>Arabidopsis</i> root cells - sc_36 | This study | GEO: GSM4626003 |
| Single Cell RNA-Seq wild type <i>Arabidopsis</i> root cells - sc_37 | This study | GEO: GSM4626004 |
| Single Cell RNA-Seq wild type <i>Arabidopsis</i> root cells - sc_40 | This study | GEO: GSM4626005 |
| Single Cell RNA-Seq wild type <i>Arabidopsis</i> root cells - sc_51 | This study | GEO: GSM4626006 |
| Single Cell RNA-Seq <i>shr</i> mutant <i>Arabidopsis</i> root cells - sc_52 | This study | GEO: GSM4626007 |
| Single Cell RNA-Seq <i>shr</i> mutant <i>Arabidopsis</i> root cells - sc_53 | This study | GEO: GSM4626008 |
| Single Cell RNA-Seq wild type <i>Arabidopsis</i> root cells - col0 | This study | GEO: GSM4626009 |
| Single Cell RNA-Seq wild type <i>Arabidopsis</i> root cells - tnw1 | This study | GEO: GSM4626010 |
| Single Cell RNA-Seq wild type <i>Arabidopsis</i> root cells - tnw2 | This study | GEO: GSM4626011 |

Experimental models: Organisms/strains

Arabidopsis thaliana

Oligonucleotides

| | | |
|---------------------------|----------------------|--|
| TCTCCATACCTCAAACCTCCTCC | N/A | F genotyping primer for <i>shortroot-2</i> |
| TTGCTCTCCGTCTACTGC | N/A | R genotyping primer for <i>shortroot-2</i> |
| CTTATCCATTCCTCAAACCTCTATT | Fukaki et al. (1998) | F genotyping primer for <i>scarecrow-4</i> . Amplifies mutant allele |
| TGGTGATCGGTAGAGAATT | Fukaki et al. (1998) | R genotyping primer for <i>scarecrow-4</i> |
| TTATCCATTCCTCAAACCTCAGT | Fukaki et al. (1998) | F genotyping primer for <i>scarecrow-4</i> . Amplifies WT allele |

Recombinant DNA

| | | |
|-----|-----|-----|
| N/A | N/A | N/A |
|-----|-----|-----|

Software and algorithms

| | | |
|--------------------------|--|---|
| Cell Ranger v3.1.0 | 10X Genomics | https://support.10xgenomics.com/single-cell-gene-expression/software/pipelines/latest/installation |
| scKB | This Study | https://github.com/ohlerlab/scKB |
| COPILOT | This Study | https://github.com/ohlerlab/COPILOT |
| Seurat v3.1.5 | Stuart et al. (2019); Butler et al. (2018) | https://satijalab.org/seurat/ |
| iRoCS | Schmidt et al. (2014) | https://lmb.informatik.uni-freiburg.de/resources/opensource/iRoCS/ |
| novoSpaRc | Nitzan et al. (2019) | https://github.com/rajewsky-lab/novosparc |
| SEMITONES | Vlot et al. (2020) | github.com/ohlerlab/SEMITONES |
| Trimmomatic v0.39.0 | Bolger et al. (2014) | http://www.usadellab.org/cms/?page=trimmomatic |
| FastQC v0.11.8 | Andrews (2010) | https://www.bioinformatics.babraham.ac.uk/projects/fastqc/ |
| STAR v2.7.1a & v2.7.2b | Dobin and Gingeras (2016) | https://github.com/alexdobin/STAR |
| DESeq2 v1.24.0 & v1.26.0 | Love et al. (2014) | https://bioconductor.org/packages/release/bioc/html/DESeq2.html |

(Continued on next page)

| Continued | | |
|---|---|---|
| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
| BBTools | Joint Genome Institute | https://jgi.doe.gov/data-and-tools/bbtools/ |
| gcrma v2.58.0 | Gentry et al. (2017) | https://www.bioconductor.org/packages/release/bioc/html/gcrma.html |
| FSQN v0.0.1 | Franks et al. (2018) | https://github.com/jenniferfranks/FSQN/ |
| gprofiler2 v0.2.1 | Kolberg et al. (2020) | https://cran.r-project.org/web/packages/gprofiler2/index.html |
| CytoTRACE v0.1.0 | Gulati et al. (2020) | https://cytotrace.stanford.edu/ |
| scVelo v0.1.25 | Bergen et al. (2020) | https://scvelo.readthedocs.io/installation/ |
| ComplexHeatmap v2.10.0 | Gu et al. (2016) | https://bioconductor.org/packages/release/bioc/html/ComplexHeatmap.html |
| StationaryOT | Zhang et al. (2021) | |
| EdgeR v3.36.0 | Robinson et al. (2010); McCarthy et al. (2012) | https://bioconductor.org/packages/release/bioc/html/edgeR.html |
| Original Codes | | |
| https://doi.org/10.5281/zenodo.5775932 | This study | https://zenodo.org/badge/latestdoi/421176705 |

RESOURCE AVAILABILITY

Lead contact

- Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Philip N. Benfey (philip.benfey@duke.edu).

Materials availability

Seeds for the *scr-4/pCORTEX:erGFP* and *scr-4/pMYB36:H2B:3xYFP* lines are available from Philip N. Benfey upon request.

Data and code availability

- Single-cell RNA-seq data have been deposited at GEO: GSE152766 and are publicly available as of the date of publication. Accession numbers are listed in the [key resources table](#). Microscopy data reported in this paper will be shared by the lead contact upon request.
- All original code has been deposited at Zenodo and is publicly available as of the date of publication. DOIs are listed in the [key resources table](#).
- 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

Seeds from wild type *Arabidopsis thaliana* ecotype Columbia (Col-0), *shortroot-2* (Col-0; Arabidopsis Biological Resource Center (ABRC) stock number CS2972), and *scarecrow-4* (Landsberg background; ABRC stock number CS6505; we backcrossed to Col-0 > 5 times) were surface sterilized with a 50% (v/v) bleach, 0.05% (v/v) Tween-20 solution for 10 minutes and subsequently stratified for 48 hours at 4°C. Seeds were sown at a density of ~150-300 seeds/row on 1X Linsmaier and Skoog (LSP03-1LT, Caisson Labs; pH 5.7), 1% sucrose media covered by 100 μm nylon mesh. Plates were placed vertically in a Percival chamber programmed to 16h light, 8h dark conditions at 22°C.

METHOD DETAILS

Protoplast isolation and scRNA-seq

Five days after sowing, 1,000-3,500 primary roots/sample were cut ~0.5 cm from the root tip and placed in a 35 mm-diameter dish containing a 70 μm cell strainer and 4.5 mL enzyme solution (1.25% [w/v] cellulase [ONOZUKA R-10, GoldBio], 0.1% Pectolyase [Sigma], 0.4 M mannitol, 20 mM MES (pH 5.7), 20 mM KCl, 10 mM CaCl₂, 0.1% bovine serum albumin, and 0.000194% (v/v) -mercaptoethanol). Roots were harvested 3-4 hours after the lights were illuminated in the growth chamber set to long day conditions. After digestion at 25°C for 1 hour at 85 rpm on an orbital shaker with occasional stirring, the cell solution was filtered twice through 40 μm cell strainers and centrifuged for 5 minutes at 500 x g in a swinging bucket centrifuge. Subsequently, the pellet was resuspended with 1 mL washing solution (0.4 M mannitol, 20 mM MES (pH 5.7), 20 mM KCl, 10 mM CaCl₂, 0.1% bovine serum albumin,

and 0.000194% (v/v) -mercaptoethanol) and centrifuged for 3 minutes at 500 x g. The pellet was resuspended with washing solution to a final concentration of ~1000 cells/ μ L. The protoplast suspension was then loaded onto microfluidic chips (10X Genomics) with v3 chemistry to capture either 5,000 or 10,000 cells/sample. Cells were barcoded with a Chromium Controller (10X Genomics). mRNA was reverse transcribed and Illumina libraries were constructed for sequencing with reagents from a 3' Gene Expression v3 kit (10X Genomics) according to the manufacturer's instructions. cDNA and final library quality were assessed with a Bioanalyzer High Sensitivity DNA Chip (Agilent). Sequencing was performed with a NovaSeq 6000 instrument (Illumina) to produce 100bp paired end reads.

Transgenic lines

Plants homozygous for the *scr-4* allele (Fukaki et al., 1998) were crossed with previously published pCORTEX:erGFP (Lee et al., 2006) and pMYB36:H2B:3xYFP (Drapek et al., 2018) transcriptional reporter lines. F2 generation seedlings were imaged at 5 days old. Individuals homozygous for the *scr-4* allele were identified by the presence of a mutant layer. pCORTEX:erGFP/*scr-4* seedlings were grown on 1X MS plates with 10 μ g/mL BASTA to confirm presence of the reporter construct prior to imaging.

Microscopy and image processing

Roots were stained with 10 mg/ml propidium iodide (PI) for 1 minute and imaged with a Zeiss 880 confocal using a x40 objective. The following are excitation (ex) and emission (em) parameters. PI: ex: 561 nm; em: 600-650 nm; YFP: ex: 488 nm, em: 530-560 nm; GFP: ex: 488 nm; em: 500-550 nm. Median longitudinal sections were chosen for each image and representative images are shown. All image analyses were performed in ImageJ. The minimum signal for each channel was adjusted by measuring the intensity histogram of the background and removing the mean plus two standard deviations from the signal. Brightness was adjusted for each channel to maximize the range of display. When GFP or YFP signals from two images are directly compared, the maximum brightness was adjusted identically for each image.

QUANTIFICATION AND STATISTICAL ANALYSIS

scRNA-seq data pre-processing

FASTQ files were generated from Illumina BCL files with Cell Ranger (v3.1.0) mkfastq (10X Genomics). Subsequently, gene-by-cell raw count matrices of spliced and un-spliced transcripts were generated using kallisto (Bray et al., 2016) (v0.46.2) and bustools (Melsted et al., 2019) (v0.40.0) as well as R packages BUSpaRse (Moses and Pachter, 2020) (v1.1.3) and BSgenome (v1.54.0; Pagès, 2020). The pipeline is summarized on our scKB GitHub repository (<https://github.com/ohlerlab/scKB>). Reads were aligned to the *Arabidopsis* genome BSgenome object ("BSgenome.Athaliana.TAIR.TAIR9") with TAIR10 gene annotation file. Samples sc_9 and sc_10 (Data S1) contained a mixture of *Arabidopsis* and rice (*Oryza sativa* X. Kitaake) root protoplasts. We mapped the reads to a concatenated version of the *Arabidopsis* TAIR10 and rice MSU7 genomes and retained only the reads which specifically mapped to the *Arabidopsis* genome. The matrices of spliced and un-spliced counts were combined into a total count matrix. Genes with no counts in any cell were removed. Cells were filtered based on the following. First, putative dying cells were identified based on the enrichment of mitochondrial gene expression (> 5% of the total UMI counts) and the mode of the putative dying cells' count distribution was treated as the initial boundary to separate cells into two groups representing low and high-quality cells. Second, expression profile references were built for both low and high-quality cells by taking the average of log-normalized counts. Third, the whole distribution of low-quality cells was recovered by comparing the Pearson correlation coefficient of each high-quality cell to the two references. In other words, if cells in the high-quality group have higher correlation to the low-quality cell profile than the high-quality cell profile, then those cells would be re-annotated as low quality. COPILOT offers functionality that allows iterative filtering until there are no cells more similar to the low-quality cell expression profile than the high-quality cell expression profile. However, in cases where the count distributions of high-quality cells and low-quality cells are not clearly separated, iterative filtering would result in over-filtering, which removes many cells that should be retained as high-quality cells. Therefore, to avoid over-filtering, we forced the algorithm to perform the cell filtering procedure only once. Finally, the low-quality cells and cells enriched in mitochondrial expression were removed along with the top 1% of high-quality cells in terms of total UMI counts in order to address any issues associated with outliers. In other words, after iterative filtering and removing cells having enriched mitochondrial expression, cells are further filtered for outliers. We used the top 1% of cells in terms of total UMI counts as a cut-off. Putative doublets were removed using DoubletFinder (McGinnis et al., 2019) with default parameters according to the estimated doublet rate (10X Genomics Chromium Single Cell 3' Reagent Kit User Guide (v3 Chemistry)). This pre-processing pipeline is available as an R package, COPILOT (<https://github.com/ohlerlab/COPILOT>), with a jupyter notebook tutorial. In downstream analyses, we did not consider mitochondrial, chloroplast, or known protoplasting-affected genes (Denyer et al., 2019) (\log_2 fold-change ≥ 2 or ≤ -2 after protoplasting). These exclusions were biologically motivated with the goal to minimize noise that may affect dimensionality reduction or clustering. e.g., chloroplast development is repressed in roots and protoplasting causes stress-related gene expression changes.

Normalization and dimensionality reduction

Using Seurat version 3.1.5, data were normalized using the SCTransform method (Hafemeister and Satija, 2019) followed by principal component analysis (PCA) and non-linear dimensionality reduction using UMAP. Fifty principal components were calculated using the RunPCA function with parameters "approx" set to FALSE. UMAP embedding was generated by RunUMAP function using all 50 principal components with parameters n_neighbors = 30, min_dist = 0.3, umap.method = "umap-learn", metric = "correlation". All

steps are incorporated into the COPILOT R package and a jupyter notebook demonstrating the analysis is provided (<https://github.com/ohlerlab/COPILOT>).

Integration of Seurat objects

Data were integrated following the Seurat reference-based integration pipeline (Stuart et al., 2019; Butler et al., 2018). The sample with the highest median UMI/gene per cell and number of genes detected was chosen as the reference (sample name: sc_12; Data S1). Overall, 16 WT replicates were used to build the atlas, including three previously published samples (Data S1). A jupyter notebook demonstrating the integration process is available on Github (<https://github.com/ohlerlab/COPILOT>).

Plotted gene expression values

'Log-normalized' indicates expression values extracted from the slot 'data' of a Seurat object's 'SCT' assay, which contains the log-normalized, 'corrected' counts produced by the SCTransform function (Hafemeister and Satija, 2019). 'Scaled Expression' indicates batch-corrected, log-normalized values extracted from the slot 'data' of a Seurat object's 'integrated' assay. These values are scaled such that any value above 10 is set to 10 (Stuart et al., 2019). However, the integrated assay only contains genes that are shared among all the samples that are integrated, which excluded some genes of interest. Therefore, given that the observed batch effect among our samples is small (Figure S1), we chose to make several plots with expression values from the 'data' slot of a Seurat object's 'SCT' assay.

Cell type and developmental stage annotation

The atlas annotation is based on comparison to published whole-transcriptome profiles (Brady et al., 2007a; Li et al., 2016) of root cells isolated from reporter lines as well as known markers (Data S1) that have been previously validated and show specific local expression on the atlas UMAP. We combined four annotation methods, described below.

Annotation based on spatial mapping

We built a 3D root geometry reference based on confocal image stacks published with the interactive *Arabidopsis* root analysis tool iRoCS (Schmidt et al., 2014). The x, y and z confocal image coordination (in micrometers) of each cell's centroid was manually documented as a location in 3D geometry followed by labeling of cell type, developmental zone, and distance from QC (in number of cells). The 3D root geometry records 3,957 cell locations covering 0.2 cm from the primary root tip (Figure S2 and Data S1). A subset containing 50,000 atlas cells was mapped to the 3D root geometry using novoSpaRc (Nitzan et al., 2019) with default parameters and binarized spatial expression profiles of 49 markers based on published images of transcriptional reporters or in situ hybridizations (Data S1). These markers serve as anchors that bridge the scRNA-seq data to the root geometry. The mapping accuracy was estimated by performing left-one-out cross validation over 100 times. Average Pearson correlation of 0.7 was achieved between predictions of the mapped model and ground truth. The mapping information of each cell from the scRNA atlas to a location was extracted, and each cell was annotated according to its mapped location. Distal root cap refers to root cap cells located at the two outermost cell layers of root cap while proximal root cap cells include root cap cell layers closer to QC. This classification is based on the observation that the cells mapped to the outermost cell layer share the same top markers (Columella: AT3G61930; Lateral root cap: AT1G33280) with the cells mapped to the second outermost layer. The cell layers closer to QC share the same sets of markers as well (Columella: AT2G04025, AT1G78520; Lateral root cap: AT1G79580). AT3G61930 is treated as a marker for proximal and distal columella in the annotation method in the next section.

Marker annotation

The enrichment scores of known cell type-specific markers (De Rybel et al., 2013; Schürholz et al., 2018; Muñoz et al., 2008; Menand et al., 2007; Bonke et al., 2003; Clay and Nelson, 2005; Lee and Schiefelbein, 2002; Brady et al., 2007b; Kamiya et al., 2016; Huang et al., 2017; Miyashima et al., 2019; Matsuzaki et al., 2010; Wallner et al., 2017; Ishida et al., 2009; Taniguchi et al., 2017; Kamiya et al., 2015; Aida et al., 2004; Kubo et al., 2005; Lee and Schiefelbein, 1999) (Data S1) were calculated for each cell in the atlas using SEMITONES (Vlot et al., 2020; github.com/ohlerlab/SEMITONES). SEMITONES uses cluster/reference-free, rank based statistics to calculate the significance of local enrichment of gene expression based on a distance between cells. Dimension reduction was performed on the raw cell-by-gene matrix and used to estimate the distance among cells to save computational resources. We chose UMAP to reduce dimensions, and distance among cells in the UMAP space was estimated via a radial basis function over the Euclidean distance (RBF kernel) metric. The size of a cell neighborhood was determined by setting the parameter "gamma" to 0.8. A gene is considered significantly enriched with respect to a cell if its enrichment score is more than 5 standard deviations away from the mean of the permutation null distribution. This permutation null distribution is obtained by applying enrichment scoring to 100 times permuted expression vectors. Cells were then annotated with a cell type label according to which significantly enriched marker had the highest enrichment score.

To complement the SEMITONES annotation approach, marker gene expression z-scores were calculated for a second marker annotation that depends on hard-clustering. In this approach, clusters were first defined using the Seurat FindClusters function by setting an extremely high modularity parameter ($res = 500$), which results in 3,034 clusters that only have tens of cells each. These finely-resolved clusters were then annotated by comparing the average marker gene z-scores. Cells that were annotated with the same cell identity by the SEMITONES and z-score approaches were considered confidently annotated. This combination was

particularly useful to annotate very young cells at the base of the UMAP because it incorporates high resolution from the z-score approach with low noise from the SEMITONES annotation.

Correlation annotation

Prior to scRNA-seq sample integration, Pearson correlation coefficient was calculated between each cell and whole-transcriptome reference expression profiles for cell types and developmental zones. We used bulk RNA-seq data (Li et al., 2016) previously generated for 14 cell types isolated with FACS. Further, we compared each cell in the atlas to ATH1 microarray data generated for thirteen cell types and thirteen tissue segments hand-dissected along the longitudinal axis of the root (Brady et al., 2007a). Each expression profile was built by first aligning the quality-filtered FASTQ reads, which are processed by Trimmomatic (Bolger et al., 2014) (v0.39) with default parameters and quality-checked by FastQC (Andrews, 2010) (v0.11.8), to the TAIR10 genome using STAR (Dobin and Gingeras, 2016) (v2.7.1a) with default parameters. Then, count normalization was carried out with DESeq and vst function in R package DESeq2 (Love et al., 2014) (v1.24.0) with default parameters. 181 genes that are highly variable across cell types in both RNA-seq and microarray data were kept, while 500 highly variable genes across 3 developmental zones and 809 highly variable genes across 13 developmental sections were selected, respectively. The SCTranform log-normalized counts in each cell and DESeq2 normalized counts in each expression profile were used to calculate Pearson correlation coefficient. Each cell was labeled with the cell type and developmental zone with which it had the highest correlation coefficient. We defined a high confidence annotation as correlation coefficient > 0.6.

Index of Cell Identity (ICI) calculation

Another method to infer cell identity was an Index of Cell Identity (ICI)-based classification approach (Efroni et al., 2015). We identified 13 datasets (Birnbaum et al., 2003; Brady et al., 2007a; Li et al., 2016; Lee et al., 2006; Nawy et al., 2005; Clark et al., 2019; Dinneny et al., 2008; Gifford et al., 2008; Bargmann et al., 2013; Yadav et al., 2014; Birnbaum and Yuan, 2015) consisting of cell-type specific gene expression profiles (RNA-seq or ATH1 Microarray) for the 18 cell types considered for this atlas (Figure S3; Data S3). RNA-seq data was preprocessed by adapter- and quality-trimming raw FASTAQ reads using the BBDuk tool (BBTools suite; sourceforge.net/projects/bbmap/), using adapter sequences found in the adapters.fa resource within bbtools, and parameters, k=23, mink=11, hdist=1, ktrim=r, and qtrim=10. Trimmed reads were mapped with the STAR (Dobin and Gingeras, 2016) utility (v2.7.2b) using default parameters with counts per gene quantified using the quantMode GeneCounts parameter. Read counts were then processed using the DESeq2 R package (Love et al., 2014) (v1.26.0), using a design matrix that treats datasets generated with the same marker:GFP construct as replicates, by running the estimateSizeFactors, estimateDispersions, and the vst functions to model gene expression. Microarray expression datasets were processed using the gcrma (Gentry et al., 2017) R package (v2.58.0). RNA-seq and microarray expression datasets were then harmonized using the FSN (Franks et al., 2018) R package (v0.0.1) to model the RNA-seq gene expression distributions using the microarray data as a reference. FSN-processed data from both the combined ATH1 and RNaseq datasets, as well as the DESeq2-processed RNaseq datasets alone, were then used to build two ICI specificity score (spec) tables (using the same methodology as described by Efroni and colleagues (2015), binning expression of each gene into 10 bins, with a minimum background bin set to 3). Markers were identified from this spec table, using a total information level of 50, and normalized, scaled expression of all identified markers was examined in all original datasets. Based on how well correlated each dataset was with its associated datasets of the same cell type, some datasets were filtered out. After dataset filtering, the final spec tables were re-calculated with the same parameters. The spec tables were then used (with an information level of 50) to compute ICI scores, p-values (using the permutation procedure described previously by Efroni et al., 2015) for all 18 cell types for cells in the atlas, using the log-transformed data values in the SCT assay of each individual dataset's Seurat object. For each cell, the highest-scoring cell type (from either the combined ATH1/RNaseq or RNaseq only spec tables) was assigned as the ICI-derived annotation. We defined a high confidence annotation as adjusted p-value < 0.01.

Combination of annotation methods

Final cell type annotations were assigned by combining information from the four annotation approaches. For procambium, metaxylem, and protoxylem cell types, which lack bulk RNA-seq or microarray references, we used only spatial mapping and marker annotation methods. For the remaining cells, if a cell had the same label from at least two of the four annotation methods, then it was annotated as such. Otherwise, the cell was temporarily treated as un-annotated during the first final annotation step. In the second step, we leveraged information from Seurat by clustering with a low modularity parameter (res = 0.5) to further prune out noise. The resulting annotation ("consensus annotation") represents the most confidently annotated cells. We built new reference expression profiles for each cell type by taking the average of the expression values for cells in the consensus annotation. All cells were then re-annotated using the correlation-based approach by comparison to these newly built references. The annotation of QC cells was performed separately since the correlation-based approach results in cells annotated as QC but that are enriched in expression of cell-cycle genes (Data S1), which does not agree with the low cell division activity of the QC. In an alternative method, we identified 158 QC cells (~0.1 % of the atlas, which is similar to QC cell type proportions from microscopy data (Cartwright et al., 2009) that have high averaged z-scores of validated QC markers and low averaged z-scores of cell-cycle genes in the SEMITONES-defined neighborhood with enriched expression of QC markers. Finally, we performed another round of denoising by clustering to obtain the final annotation.

To assign a developmental stage annotation to each cell, we used an approach similar to that described for cell type annotation, during which we used microarray-based whole-transcriptome profiles from thirteen root longitudinal sections as reference expression profiles (Brady et al., 2007a). Sections meristem 1-6, elongation 7 and 8, and maturation 9-12 correspond to the atlas meristem, elongation, and maturation labels, respectively. In practice, cell type and developmental stage annotations were performed simultaneously, meaning that the newly built references described in previous sections refer to the combination of developmental stage and cell type. A jupyter notebook demonstrating the annotation process is available from Github (<https://github.com/ohlerlab/COPILOT>).

Ploidy annotation

We assigned each cell a ploidy label based on correlation with four published bulk RNA-seq profiles (Bhosale et al., 2018) (Figure 2B; Data S1). A jupyter notebook demonstrating the annotation process is available from Github (<https://github.com/ohlerlab/COPILOT>).

Differentially expressed genes

To identify cell type and cell type + developmental stage marker genes, we used the default Wilcoxon test available from the Seurat FindMarkers function on scaled expression. Significant markers for cell type + developmental stage were selected based on the following criteria: adjusted p-value < 0.05, average log fold change > 3, and pct.diff > 0.4, where pct.diff is defined as the difference of gene percentage expression between the cluster considered and the rest of the cells. Genes that were identified as markers for multiple cell types were reassigned to the cell type with the highest average log fold change and pct.diff. Marker specificity was estimated by percentage expression in cells that do not belong to the cluster considered. The expression pattern of marker genes was also verified with Seurat's dot plot tool.

In addition to cluster-dependent differential expression analysis implemented in Seurat, the cluster-agnostic tool SEMITONES was used to search for cell type + developmental stage marker genes *de novo* based on scaled expression. Reference cells for each cell type + developmental stage were chosen by searching for cells with the highest average similarity based on a similarity matrix calculated via the RBF kernel on 50 UMAP dimensions.

In DE analyses along pseudotime bins, we used the Seurat FindMarkers function to first prefilter features using a log₂ fold-change threshold of 1 and a minimum percentage difference in expression of 0.25. We then performed differential expression testing for each combination of cell type and pseudotime bin using the receiver operating characteristic (ROC) test implemented in Seurat FindMarkers. A classifier was built for each gene based on the ability of that gene's expression level to distinguish between two groups of cells. The first group of cells corresponds to the pseudotime bin of interest within a particular cell type whereas the second group is the remaining cells within the trajectory for that tissue. Classification power based on Area under the ROC Curve (AUC) was used to estimate the performance of the classifier. An AUC value of 1 indicates increased expression values in the first group that can perfectly distinguish the two groupings, whereas an AUC of 0.5 indicates that the gene has no predictive power to distinguish the groups. Only markers with an AUC greater than 0.75 were retained for downstream analysis. We rank ordered markers based on AUC, percentage difference, and fold-change.

Bifurcation patterns on atlas UMAP

To examine bifurcation patterns within cell lineages, ground tissue and epidermis sub-branches were labeled based on clusters identified with Seurat (modularity parameter $res = 0.5$) (Figure S4). Gene ontology analysis was conducted on identified DE genes using R package "gprofiler2" (Kolberg et al., 2020).

Pseudotime estimation

Pseudotimes were inferred with the R package CytoTRACE (Gulati et al., 2020) (v0.1.0) and Python-based scVelo (Bergen et al., 2020) (v0.1.25). We opted not to use graph-based tools given their dependency on the selection of dimensional reduction embeddings and parameters. The batch-corrected and scaled ('integrated' assay in Seurat object) expression values were used as input for CytoTRACE and scVelo. Instead of using the default scaled expression values which were centered at 0 and capped at 10, all the negative values were treated as no expression and the values were floored at 0. The ratio of spliced and un-spliced transcripts of each gene and cell was calculated using raw counts. The ratio was then multiplied by the batch-corrected non-negative expression count matrix to generate the corresponding spliced and unspliced count matrices, which serve as input for scVelo. Latent time was then estimated by running pp.moments function with parameter, $n_pcs = 50$, $n_neighbors = 100$ and tl.velocity function with mode set to "dynamical" in scVelo, while CytoTRACE was implemented with default parameters.

A consensus pseudotime was derived by taking the average of CytoTRACE and scVelo-inferred latent time. Consensus time was estimated for each tissue/lineage independently to address differences in maturation rates. The consensus time for QC cells were then averaged and all the cells in the trajectory were divided into ten evenly sized groups (T0-T9) each containing the same number of cells. We chose ten bins after examining the data annotated according to correlation with microarray data from twelve manually dissected longitudinal tissue sections (Brady et al., 2007a). Of the twelve section labels, we found that two (Meristem-section 6 and Maturation-section12) were outliers and had fewer cells than the other ten sections. We therefore chose ten bins to more evenly spread the cells across all bins. A jupyter notebook demonstrating how results from the two tools were combined is provided under the GitHub repository for COPILOT (<https://github.com/ohlerlab/COPILOT>).

Genes dynamically expressed across pseudotime

We applied the approach described under ‘Differentially expressed genes’ to identify genes that vary along the developmental progression of each tissue type. We used the combination of cell type and consensus time group (10 groups ranging from T0 to T9) as identity of interest among which differential expression analysis was performed. Spearman’s correlation of each marker with consensus time was considered as an additional metric to aid in selecting genes that vary along the gradient of differentiation. Ten genes were selected for each cell type and consensus time group combination. Genes were arranged according to their highest rank along consensus time. Pseudo-bulk expression profiles within each consensus time group were calculated for each gene and row scaled expression values were then displayed using ComplexHeatmap in R (Gu et al., 2016) (v2.10.0). All code used to identify and plot genes differentially expressed across pseudotime is available as a jupyter notebook on GitHub (<https://github.com/ohlerlab/COPILOT>).

Computing trajectories with StationaryOT

Daily growth rates were estimated from imaging data of the growing meristem over a period of up to a week (Rahni and Birnbaum, 2019). Using these growth rates and examining the proportion of cells in each developmental stage, we estimated that roughly 5% of the cells in each lineage would be replaced in a 6-hour period. We selected the top 5% most differentiated cells from each lineage as sinks, as defined by pseudotime. We applied StationaryOT using entropic regularization with the regularization parameter set to $\varepsilon = 0.025$, and the cost matrix normalized to have unit mean. However, we found the results to be robust using quadratic regularization and varying the time and degree of regularization by a factor of two. Due to computational limitations, the dataset was partitioned into 10 subsets and StationaryOT was applied to each subset. This was repeated 10 times with random partitions to account for sampling error. Cell-by-cell averaging was performed on the computed fates to create a set of consensus fates. Between a single subset in a partition and the consensus fates in the full atlas, 97% of cells shared the same maximum fate type and the maximum fate values had a correlation of 0.96. Accounting for all fate values, rather than just maximums, the correlation rose to 0.99.

Visualizing fate probabilities

StationaryOT assigns a vector of fate probabilities to each cell. Up to three fates are visualized at a time (e.g., endodermis, cortex, and other being the sum of the remaining fates) using barycentric coordinates to represent 3-dimensional probability vectors in a two-dimensional triangle plot. A corner of the triangle is associated with each of these possible fates, and cells are positioned according to their relative probabilities as follows:

Let a, b, c denote the vertices of the triangle in R^2 and let $p = (p_1, p_2, p_3)$ denote the probability vector we wish to visualize. The components of p are used as coefficients in a convex combination of the vertices. In other words, the probability vector p is mapped to $p_1a + p_2b + p_3c \in R^2$. Note that $p_1 + p_2 + p_3 = 1$, so each probability vector is mapped to a point inside the triangle. Cells perfectly fated to obtain a single fate are positioned exactly at the corresponding vertex, while cells with indeterminate fates are positioned in the interior of the triangle. The very center of the triangle corresponds to cells that are equally likely to choose any of the three fates, and cells along an edge have zero chance of reaching the opposite vertex.

Lasso regression

To identify genes that play roles in lineage determination, we applied Lasso regression to gene expression data and fate data from StationaryOT. This analysis was applied to cells in each developmental stage, and then the full dataset. As a result, we obtained lists of genes with possible lineage determining roles for each stage. Lasso is a linear regression method with an L^1 regularization term to control sparsity (Tibshirani, 1996). We applied Lasso to gene expression matrices E_s , for each developmental stage (meristem, elongation, maturation) and the full atlas. For the regressions, we restricted E_s to only contain expression data from transcription factors (Pruneda-Paz et al., 2014). Note that cells from the root cap were assigned meristematic, elongation, and maturation stage labels according to correlation annotation with bulk RNA-seq datasets generated from hand dissected tissue. This was done to create a fit that is applicable to all cell types, and hence is more selective in its component genes. The regression was performed on a lineage-by-lineage basis against $f_{s,L}$, fate probabilities for cells from stages, to a lineage L . In this setting, the objective function for Lasso is:

$$\frac{1}{2n} \|f_{s,L} - E_s w\|_2^2 + \alpha \|w\|_1$$

Here, n is the number of cells, w is a vector of regression coefficients, and α is a regularization coefficient. To determine an optimal α for each regression, $\alpha_{opt, s, L}$, that balanced sparsity and predictive power, we tested a range of α for each stage and lineage. We created a graph of R^2 versus the number of non-zero coefficients for each fit. We then chose $\alpha_{opt, s, L}$ by selecting the value of α corresponding to the knee point of the graph, which was determined using the Kneedle algorithm (Satopaa et al., 2011). An example of these graphs for the meristematic zone are found in Figure S7. The linear_model.Lasso function from the Python package scikit-learn was used as the solver for the regressions (<https://scikit-learn.org/stable/about.html#citing-scikit-learn>). The regression assigns a coefficient w_i to each gene i , which determines its predicted impact on lineage determination. Specifically, the coefficient for a gene is a prediction of how much a unit change in that gene’s atlas expression values (after normalization and integration) affects a cell’s probability of achieving that lineage. Coefficients can be either positive or negative: a positive coefficient for a gene implies that up-regulation of that gene favors the given lineage, while a negative coefficient implies that down-regulation favors the lineage.

The magnitude of these regression coefficients can be used to rank genes in terms of lineage determining capacity, providing candidates for further investigation. Gene lists for each stage and lineage are included in [Data S5](#).

shortroot and scarecrow mutant analysis

Annotations were transferred from the atlas to two *scr-4* biological replicates, two *shr-2* biological replicates, and five wild type biological replicates that were grown and processed together with the mutants (WT samples sc20, 21, 30, 31, and 51). Label transfer was performed following the Seurat pipeline. A jupyter notebook tutorial is available on Github (github.com/Hsu-Che-Wei/COPILLOT). Mutant and WT data were integrated following the Seurat reference-based integration pipeline ([Stuart et al., 2019](#); [Butler et al., 2018](#)).

Cell identity differential abundance

We used differential abundance analysis to examine which cell types were enriched or depleted in *shr* or *scr* compared to WT ([Amezquita et al., 2020](#)). First, we quantified the number of cells assigned to each label on a per sample basis. We then used the EdgeR package ([Robinson et al., 2010](#); [McCarthy et al., 2012](#)) to fit a negative binomial generalized linear model in which the counts represent cells per label. Normalization was conducted according to the number of cells per sample. Separate contrasts were used to compare *shr* versus WT or *scr* versus WT, each with a blocking factor to account for any potential batch effects between different experimental runs. Differences in abundance were tested using the function `glmQLFTest`. P-values were adjusted for multiple testing according to [Benjamini and Hochberg \(1995\)](#) and cell type labels with a false discovery rate less than 0.05 were considered significantly altered. We then used ComplexHeatmap ([Gu et al., 2016](#)) in R to plot the \log_2 fold-change estimates (mutant/WT) from EdgeR.

ADDITIONAL RESOURCES

Data deposition: <https://www.ncbi.nlm.nih.gov/geo/>

Interactive web browser for the atlas: <https://phytozome-next.jgi.doe.gov/tools/scrna/>