

# Single-Cell Resolution of Lineage Trajectories in the *Arabidopsis* Stomatal Lineage and Developing Leaf

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## SUMMARY

Dynamic cell states underlie flexible developmental programs, such as with the stomatal lineage of the *Arabidopsis* epidermis. Initial stages of the lineage feature asynchronous and indeterminate divisions modulated by environmental cues, enabling cell fate flexibility to generate the requisite density and pattern of stomata for a given environment. It remains unclear, however, how flexibility of cell fates is controlled. Here, we uncovered distinct models of cell state differentiation within *Arabidopsis* leaf tissue by leveraging single-cell transcriptomics and molecular genetics. Our findings resolved underlying heterogeneity within cell states of the flexible epidermal stomatal lineage, which appear to exist along a continuum, with progressive cell specification. Beyond the early stages of the lineage, we discovered that the core transcriptional regulator SPEECHLESS is required for cell fate commitment to yield stomatal guard cells. Overall, our work has refined the stomatal lineage paradigm and uncovered progressive cell state decisions along lineage trajectories in developing leaves.

## INTRODUCTION

Cellular fate specification and differentiation are core features of developmental programs in multicellular organisms. Molecular genetics established the classical view of cell fate commitments as discrete and sequential stages, but recent technological advances with single-cell RNA sequencing (scRNA-seq) have revealed ways in which cell state decisions are continuous and heterogeneous. Cell states may indeed be now defined by gene expression profiles that mark states as putatively dynamic instances of a given cell transition or identity, corresponding to cell function and lineage relationships (Morris, 2019; Sagar and Grün, 2020; Wagner et al., 2016). While remarkable progress has been made to elucidate cell states within a range of tissue and animal systems (Karaïskos et al., 2017; Schaum et al., 2018; Sebé-Pedrós et al., 2018; Siebert et al., 2019; Tintori et al., 2016; Wagner et al., 2018), much is to be learned from state dynamics in plants. In addition to exhibiting many differentiated cell types with no counterparts in animals, plants display extreme biological flexibility, longevity, and

regenerative capacity. Multicellularity, and therefore developmental processes, arose independently in plants and animals, though there are many cases of convergence in the deployment of specific molecular regulators and developmental strategies.

Recent applications of scRNA-seq to plants include detailed analyses of regeneration and meiosis utilizing hundreds of highly selected cells (Efroni et al., 2016; Nelms and Walbot, 2019). Conversely, atlases comprised of tens of thousands of cells primarily focus on the *Arabidopsis thaliana* (herein referred to as *Arabidopsis*) root, and they provide refined insight to well-characterized cell type-specific lineage progressions (Denyer et al., 2019; Ryu et al., 2019; Shaham et al., 2020; Shulse et al., 2019; Zhang et al., 2019). While these pioneering studies have validated scRNA-seq approaches for resolving unidirectional differentiation processes, they have not focused on flexible and indeterminate

features of plant development. One exemplary developmental model for cell fate specification and flexible lineage progression is the *Arabidopsis* epidermal stomatal lineage (Pillitteri and Torii, 2012). Stomata are cellular valves consisting of two sister “guard cells” that modulate the aperture of a pore between them to facilitate atmosphere gas exchange. Guard cells are one of the final products of a multipotent stem cell lineage that gives rise to the leaf epidermis. Initial stages of the lineage feature asynchronous and indeterminate divisions modulated by extrinsic environmental cues, enabling cell fate flexibility to generate the requisite density and pattern of stomata for a given environment (Hetherington and Woodward, 2003). Given that environmental cues integrate with underlying genetic programs, it is thought that core regulators serve dynamic roles within cell states to tune cell specification and differentiation (Lau et al., 2018; Vatén et al., 2018), though it is unclear how the flexibility of cell fates is controlled.

Here, we leveraged scRNA-seq and molecular genetics to examine dynamic developmental states of the stomatal lineage in the context of the growing *Arabidopsis* leaf. Examining stomatal production and lineage progression within a broader organ-level context provides insight into cellular programs that coordinate to build a functional organ. scRNA-seq identified distinct models of cell state differentiation within leaf tissue, revealing cell states along with putative regulators across the mesophyll, vasculature, and epidermis. Our atlas of leaf development serves as a useful resource to pursue creative questions and generate hypotheses, as well as inspire comparative analyses with other organs or species, with the goal to better define attributes of tissue development both within an organism and between species. Applied to the question of developmental flexibility, scRNA-seq revealed that flexible cell states on the epidermal landscape may exist along a continuum, with progressive cell specification. This model suggests that obligate core transcriptional regulators and signaling cascades control a range of cellular programs within lineage continuums. One such critical regulator, the transcription factor SPEECHLESS (SPCH), is known to integrate environmental fluctuations in the early stages of the lineage. However, our scRNA-seq and subsequent functional perturbations indicated that SPCH serves a broadened role that enforces cell specification and commitment to generate stomata. Collectively, our findings uncover progressive cell state decisions along lineage trajectories in developing leaves.

## RESULTS

### Complementary scRNA-seq approaches to resolve underlying cell heterogeneity

To leverage our understanding of flexible divisions within the epidermal stomatal lineage, while also surveying developmental processes within inner tissue mesophyll and vasculature, we used complementary single-cell RNA sequencing (scRNA-seq) approaches (Table S1) with 10 days-post-germination (dpG) seedlings. Overall, we sequenced transcriptomes from ~18,000 cells with droplet-based 10X Genomics and ~500 cells with plate-based Smart-seq2. This approach allowed us to capture a range of cell states, gene coverage, and cytometry data. In all cases, prior

to sequencing, whole leaf tissue was broken down by enzyme-mediated protoplasting followed by fluorescence-activated cell sorting (FACS), which enriched for rare stomatal lineage cells that are challenging to capture from whole tissue. We ultimately resolved underlying heterogeneity among cell states in the developing leaf. Here, we define cell states as instances of putative cell transitions or types (e.g. specific precursor or mature cell) and thereby encompass the range of possibilities for what it means to be a cell along a lineage trajectory (Morris, 2019). Insight into cell states enables us to build models of differentiation programs delineated by predicted lineage trajectories.

### Distinct models of cell state differentiation within leaf tissue

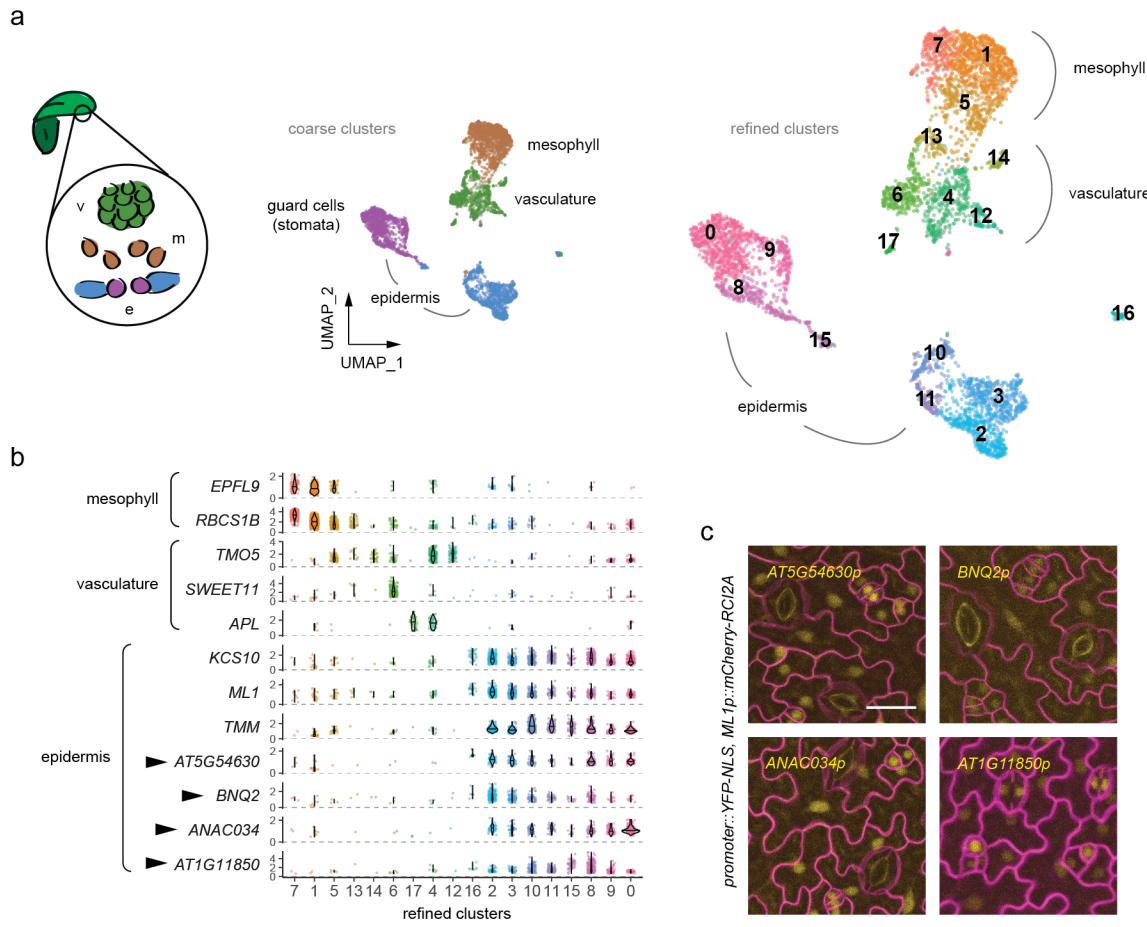
We first profiled cell states from seedlings that expressed the transgene *ML1p::YFP-RCI2A*, in which YFP is fused to the transmembrane domain-containing protein RCI2A and expressed under the predominantly epidermal homeobox transcription factor *MERISTEM LAYER 1 (ML1)* (Lu et al., 1996; Nylander et al., 2001; Roeder et al., 2010). It should be noted that the *ML1* gene promoter is also lowly active in internal tissue cells (Iida et al., 2019), and previous studies showed that FACS-enrichment with this promoter isolates mesophyll cells among other cell types (Adrian et al., 2015; Tian et al., 2019; Yadav et al., 2014). Thus, while our approach enriched epidermal cells at the expense of the more abundant mesophyll cells, the sorted pool of cells was expected to contain a mix of epidermal and inner-tissue cells that would enable comparative analyses. Single cells were captured by droplet-based 10X Genomics microfluidics (Zheng et al., 2017) for downstream library preparation and sequencing. We built our gene expression matrix with Cell Ranger and used the Seurat v3 analysis pipeline (Butler et al., 2018) to obtain transcriptomes for 5,021 cells, with a median of 1,870 detected genes and 5,026 unique molecular identifiers (UMIs) per cell (Figure S1). Our scRNA-seq atlas of the developing leaf offers a substantial number of detected genes and cell expression profiles to uncover putative developmental programs.

Graph-based unsupervised clustering and differential expression analysis revealed potential cell states within mesophyll, vasculature, and epidermal tissues (Figure 1A, Table S5). Clusters of cells were visualized on a Uniform Manifold Approximation and Projection (UMAP) plot, wherein local and global graph structure is preserved to represent the similarity of cell transcriptomes within and between clusters (Becht et al., 2018; McInnes et al., 2018). While the analysis can be performed at distinct resolutions to yield broader or refined clusters (Figure S2, Figure 1A), we primarily focused on a refined resolution to enable examination of putative cell states. At the developmental stage profiled, there are proliferating cells in all aerial tissues of the seedling. The transcriptional signature of cell cycle phases can override and confound cell state assignment (Butler et al., 2018), so we performed a regression to correct for this and enable comparisons of related modules within different tissues (Table S4, Figure S1).

We began by broadly defining coarse cell clusters that correspond to mesophyll, vasculature, and epidermal cells, based on the expression of well-characterized, tissue-specific players (Figure 1A-B). Similar to findings in *Arabidopsis* roots (Shulse et

al., 2019), many genes with known roles in leaf development were enriched in our differential expression analysis, but were not necessarily the most enriched genes (Table S5). Mesophyll cells were demarcated by expression of the secretory peptide gene *STOMAGEN* (also known as *EPIDERMAL PATTERNING FACTOR LIKE 9, EPFL9*) (Hunt et al., 2010; Kondo et al., 2010; Sugano et al., 2010) and the photosynthetic enzyme RuBisCO subunit gene *RBCS1B*, among other genes encoding for proteins involved in photosynthesis (Sawchuk et al., 2008). Vasculature cells were demarcated by genes encoding regulators of the xylem and phloem, respective water and sugar transporting tissues. These included the xylem bHLH transcriptional regulator gene *TARGET OF MONOPTEROS 5 (TMO5)* (De Rybel et al., 2013), along with the phloem sugar transporter gene *SWEET11* (Chen et al., 2012) and MYB transcriptional regulator gene *ALTERED*

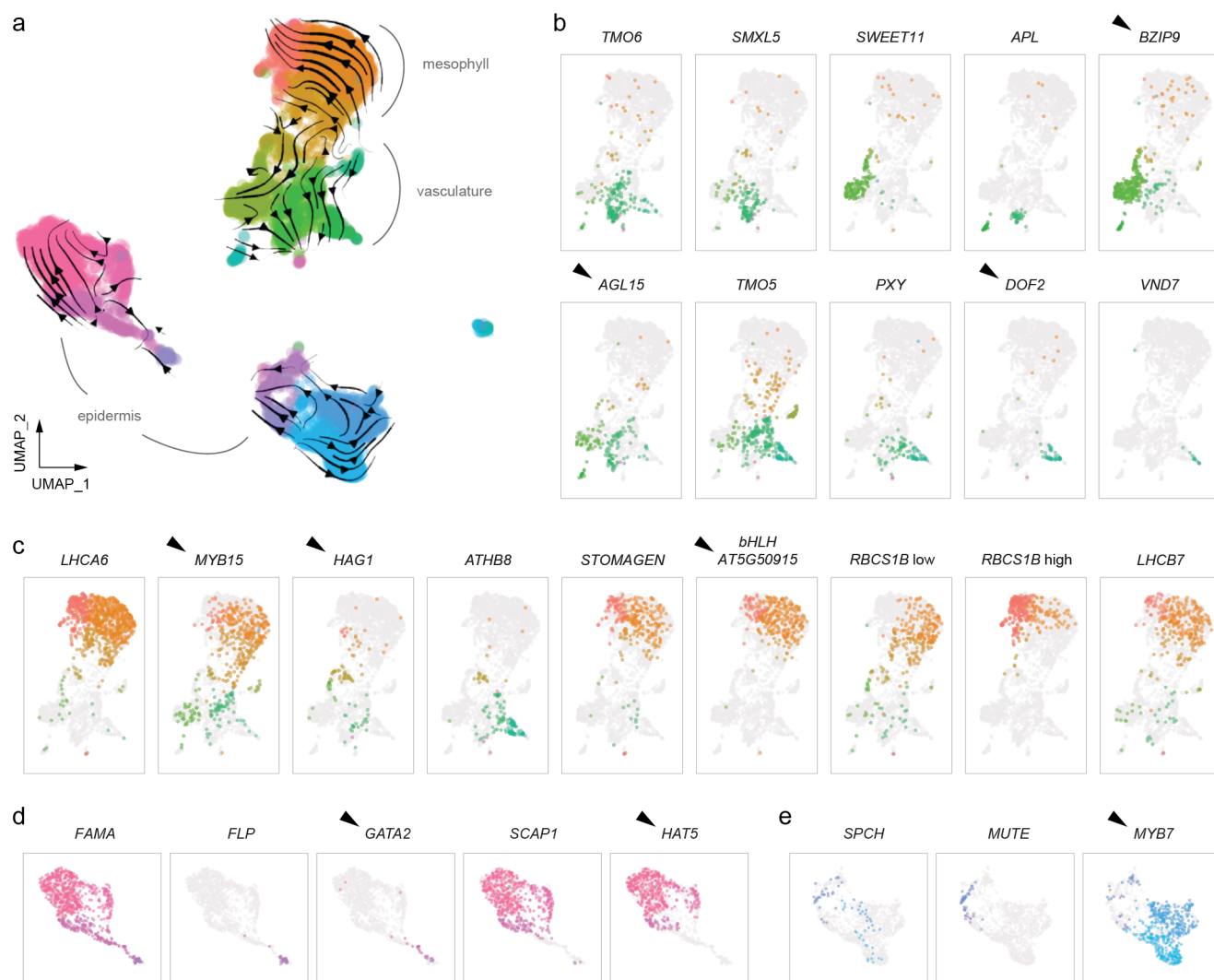
*PHLOEM DEVELOPMENT (APL)* (Bonke et al., 2003). Epidermal cells were demarcated by broad expression of the homeobox transcriptional regulator gene *ML1*, along with genes encoding the cuticle biosynthesis enzyme 3-KETOACYL-COA SYNTHASE 10 (*KCS9/FDH*) (Pruitt et al., 2000; Yephremov et al., 1999) and the stomatal lineage leucine-rich repeat (LRR) receptor-like *TOO MANY MOUTHS (TMM)* (Nadeau and Sack, 2002). We found that mesophyll and vasculature cell identities lie adjacent to each other within our developing leaf atlas, while younger epidermal cells are separated from differentiated guard cells. To further validate our atlas, we generated in planta *YFP-NLS* reporters with promoters from a selection of differentially expressed genes (Figure 1B-C, Figure S3), which provided additional support for our epidermal cluster assignments.



**Figure 1. scRNA-seq atlas of the developing leaf.** (a) Schematic of a leaf cross-section (left, m: mesophyll, v: vasculature, e: epidermis) is color-coded to match the UMAP of developing leaf tissue from scRNA-seq (10X Genomics v2, 5,021 cells displayed) with 10 days-post-germination (dpg) *ML1p::YFP-RCI2A* seedlings. With a coarse resolution (left, resolution 0.1), major clusters from the mesophyll, vasculature, and epidermis tissue layers are distinguishable. Fine resolution clusters (right, resolution 1) of putative cell states within our UMAP atlas underscore the possibility of capturing state transitions within lineages. Colors and respective number labels represent distinct clusters from graph-based clustering, and number labels designate the order from the largest to smallest cluster (0 to 17, respectively), based on the number of cells per cluster. (b) Violin plots depict expression profiles of known and predicted tissue-specific players within resolution 1 clusters, colored accordingly. Arrowheads correspond to expression profiles of genes with unappreciated roles in the epidermis, for which reporters were generated in (c). (c) Confocal images of promoter-driven *YFP-NLS* reporters (yellow) corroborate predicted profiles from our developing leaf UMAP atlas. Cell outlines (magenta) are visualized with *ML1::mCherry-RCI2A*, and the abaxial epidermis was imaged from first true leaves of 10 dpg seedlings. All images were taken at the same magnification (scale bar: 20  $\mu$ m).

To elucidate potential cell states within the mesophyll, vasculature, and epidermis, we inferred transcriptional dynamics with a steady-state deterministic model using scVelo (Figure 2A) (Bergen et al., 2019; La Manno et al., 2018). Briefly, the approach predicts future cell states by leveraging ratios of nascent (unspliced) and mature (spliced) RNA transcripts. In our sequencing data, we detected reads of unspliced transcripts from secondary priming events during 10X Genomics library construction (5% of total reads: ~3,500 unspliced reads/cell with ~70,000 total reads per cell). We therefore were able to identify

putative, directed transitions between cell states in our atlas, which were corroborated by previously described expressed genes (Figure 2B-E), even though the scVelo model relies on rarely detected transcripts and cell states. We also identified enriched components of signaling pathways and transcriptional control that define these transitions and states in the mesophyll, vasculature, and epidermis (Table S5). Our atlas thus has the potential to yield insight into developmental lineage trajectories that have eluded previous analyses.



**Figure 2. Models of cell state differentiation within leaf tissue.** (a) Lineage inference models by scVelo layered onto the UMAP of the developing leaf scRNA-seq atlas. Size of arrows indicate the degree of gene expression changes within predicted, directed transitions from one cell state to another. Colors represent graph-based clustering with resolution 1, and grey brackets demarcate major clusters from the mesophyll, vasculature, and epidermis. (b-e) UMAP insets of the major clusters that correspond to different tissues; which include (b) vasculature, (c) mesophyll, (d) stomatal guard cells, and (e) young epidermal cells. Expression of respective dynamic lineage genes is highlighted by resolution 1 cluster colors. Two insets depicting *RBCS1B* expression distinguish 'low' versus 'high' gene expression within mesophyll cells. Cells with 'low' and 'high' expression were selected from the bottom and top 25%, respectively. Representative, previously uncharacterized, potential regulators are designated by arrowheads.

Genetic programs that drive specification and differentiation of vascular xylem and phloem tissues are well defined in root, stem, and hypocotyl tissue (Hellmann et al., 2018), and thus provide a useful framework to evaluate the scVelo lineage inference trajectory. Using known markers, we identified procambial meristematic cells, vasculature precursor cells, and maturing phloem and xylem cells (Figure 2B). The developing leaf vasculature exhibited a similar trajectory as in other tissues; expression of the transcriptional regulator genes *SUPPRESSOR OF MAX2-LIKE 5* (*SMXL5*) and *TARGET OF MONOPTEROS 6* (*TMO6/DOF5.3*) (Wallner et al., 2017) demarcated differentiating procambium and phloem, while the sugar transporter gene *SWEET11* (Chen et al., 2012) defined potential companion and parenchyma cells, and the MYB family transcriptional regulator gene *APL* was expressed in mature phloem cells, downstream of the predicted onset of *SMXL5* and *SWEET11*. The bHLH transcriptional regulator gene *TMO5* was expressed in the procambium and overlapped with *SMXL5* expression in our atlas. It was also expressed in the xylem, along with the receptor-like kinase gene *PHLOEM INTERCALATED WITH XYLEM* (*PXY*) (Fisher and Turner, 2007). Downstream of *TMO5* and *PXY* expression in our lineage inference, the transcriptional regulator gene *VASCULAR RELATED NAC-DOMAIN PROTEIN 7* (*VND7*) (Kubo et al., 2005) was restricted to differentiating xylem cells with activated secondary wall biosynthesis programs. Our differential expression analysis identified several putative transcription factors with unappreciated roles in the vasculature (e.g. *BASIC LEUCINE ZIPPER 9*, *BZIP9*; *AGAMOUS-LIKE 15*, *AGL15*; and *DOF ZINC FINGER PROTEIN 2*, *DOF2*) (Table S5). A comparison with root scRNA-seq data (Ryu et al., 2019) indicated similar restriction of expression to phloem or xylem cell types. Further analysis is necessary to define the function of these genes, though as a general strategy for prioritizing potential lineage regulators, there is weight in the consistent association of a gene with common cell types from different tissues. Conversely, identification of regulators specific to leaf vasculature cells, compared with the root and other tissues, may point to respective divergent developmental processes.

Given that we were able to elucidate vasculature cell states, we next asked whether our developing leaf atlas could provide insight into mesophyll differentiation. Mesophyll function in photosynthesis is extensively studied, but less is known about cell differentiation states in this considerably less accessible and tractable inner leaf tissue. Trajectory inference indicated that distinct cell fate decisions distinguish the mesophyll and vasculature (Figure 2C). Clusters of cells from the mesophyll and vasculature appear bridged by putative ground meristem and procambium cells that give rise to differentiating cells within each tissue. For instance, onset of the light-harvesting complex gene *LHCA6* and the homeobox transcription factor gene *ATHB8*, previously characterized progenitor markers (Sawchuk et al., 2008; Scarpella et al., 2004), was detected in ground cells that are committed to mesophyll and vasculature development, respectively. We thus leveraged differentially expressed genes (Table S5) to identify enriched putative regulators of the ground meristem, which yielded the transcription factor genes *MYB15* and *HAG1/MYB28*. Conversely, analysis of putative mesophyll differentiation regulators identified the bHLH transcription factor

gene *AT5G50915*, that also overlapped with the secretory peptide gene *STOMAGEN/EPFL9* involved with early epidermal-mesophyll tissue coordination to control stomatal density (Hunt et al., 2010; Kondo et al., 2010; Sugano et al., 2010). Interestingly, we may be able to distinguish the adaxial palisade and abaxial spongy mesophyll, based on respective divergent expression of the light-harvesting complex gene *LHCb7* and the RuBisCO subunit gene *RBCS1B* (Sawchuk et al., 2008), though there is substantial overlap between these expression profiles. One feature of *RBCS1B* expression, as with other highly expressed genes, was that it also enabled us to distinguish a quantitative gradient within our scRNA-seq data, with low expression detected in cells predicted to be precursors of those with higher expression. Through new perspectives on establishment and maintenance of cell states, our atlas may extend our understanding of palisade and spongy mesophyll differentiation.

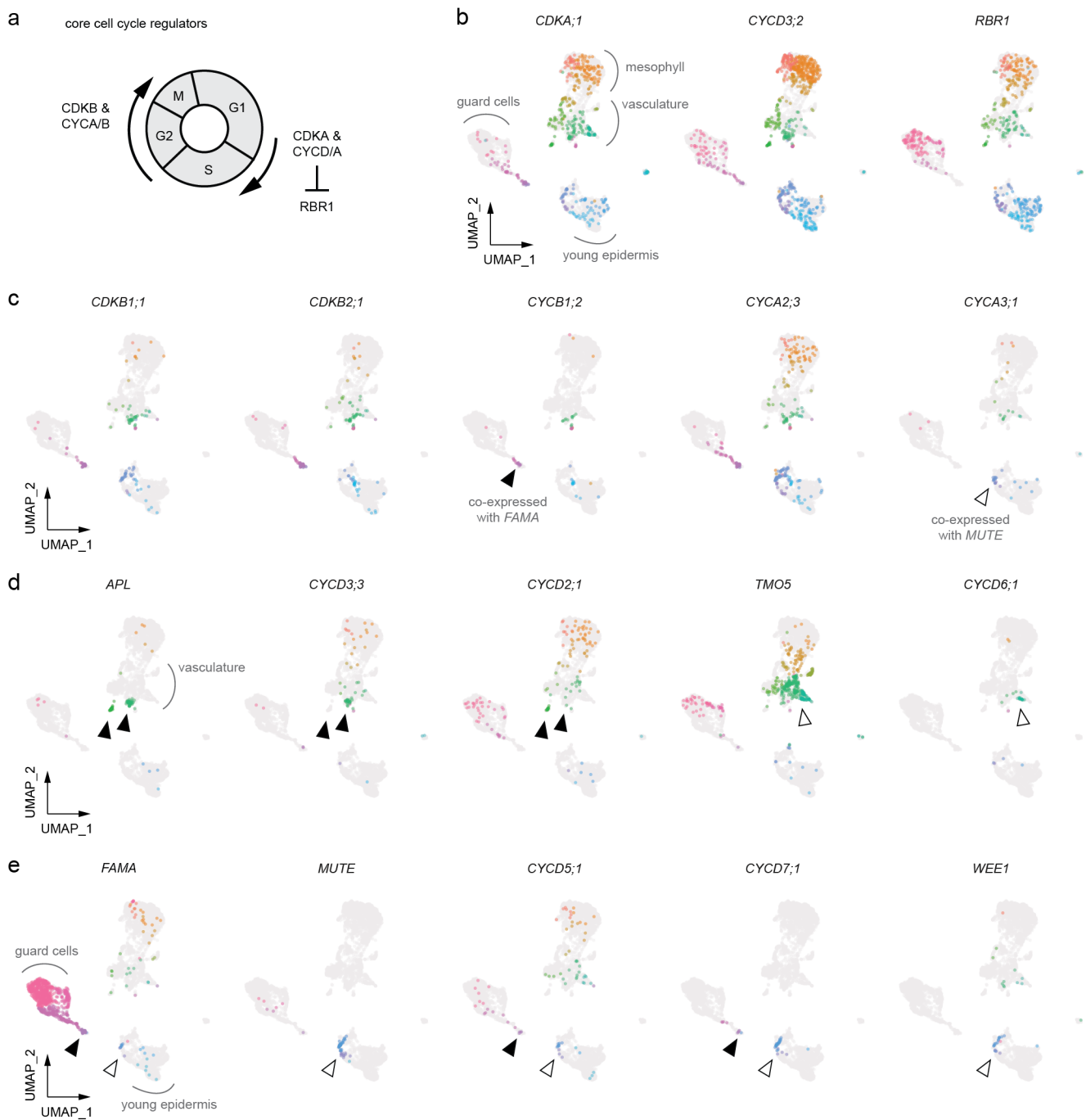
Lineage inference also provided insight into directed developmental transitions within the epidermis. Epidermal cells were divided into two main clusters in our atlas. One cluster featured expression of the bHLH transcriptional regulator gene *FAMA*, which indicated that it represents differentiating stomatal guard cells (Figure 2D). Within this cluster, the MYB transcription factor gene *FOUR LIPS* (*FLP/MYB124*) exhibited much more restricted expression, as previous reports link *FLP* to the earliest stages of guard cell differentiation (Lai et al., 2005). Restricted to the opposite region within the cluster was the Dof-type transcriptional regulator gene *STOMATAL CARPENTER 1* (*SCAP1*) (Negi et al., 2013), indicating that these are mature guard cells. With *FAMA*, *FLP*, and *SCAP1* gene profiles as guides of differentiation states, differential gene expression analysis identified additional putative regulators, such as the transcription factors *GATA2* and *HAT5* that respectively mark the beginning and end of guard cell differentiation.

The second epidermal cluster in our atlas represented young epidermal and stomatal lineage cells (Figure 2E). A notable feature of this cluster was its diverging trajectory inference (depicted as deviating arrows on the UMAP, Figure 2A) modeling multiple cell fate decisions. A finer examination of these states that includes additional scRNA-seq data follows below (Figure 4B), but at this resolution, we observed relatively broad expression of the bHLH transcriptional regulator gene *SPCH* (MacAlister et al., 2007; Pillitteri et al., 2007) in early stomatal lineage stem cells. Its parologue *MUTE* (MacAlister et al., 2007; Pillitteri et al., 2007) displayed much more restricted expression in cells committing to stomatal guard cell differentiation, and accordingly, these cells in the young epidermal cluster were oriented toward the guard cell cluster. Opposed to *MUTE*, cells expressed the transcription factor gene *MYB7*. *MYB7* has been associated with production of secondary metabolite phenylpropanoids (Albert, 2015) and may define cells progressing toward the alternative pavement cell fate. Genes encoding cuticular wax biosynthesis and defense-related factors were co-expressed with *MYB7* (Table S5), and overall there were a substantial number of putative regulators dynamically expressed in patterns traversing the young epidermal clusters. Thus, in contrast with previous RNA-seq studies that were biased by bulk identification of cells expressing specific markers (e.g. *SPCH*, *MUTE*, and *FAMA*) (Adrian et al., 2015), leveraging the

broadly expressed *ML1* gene promoter with scRNA-seq enabled us to capture cells that take alternative trajectories to generate either stomata or pavement cells. We thus resolved heterogeneous transcriptional changes and distinct models of cell state differentiation on the epidermis of developing leaves.

An emergent signature from our atlas may mark the spatial origin of cells. All tissues in the leaf display adaxial (top) and abaxial (bottom) polarity (Kidner and Timmermans, 2010), which can be seen with the vasculature xylem and phloem, palisade and spongy mesophyll, and the adaxial and abaxial epidermis. Overlaying expression of the core regulators, including members

of the *HD-ZIP III*, *KANADI* and *YABBY* gene families (Figure S4), enabled us to detect expected patterns of the leaf polarity axis in internal tissues. We also found that we have potentially obtained epidermal cells from each side of the leaf. Because expression of such polarity genes is highest in leaf primordia (Ram et al., 2020), we lose the ability to more confidently assign location in maturing leaf cells. Nonetheless, we found a substantial number of guard cells expressing adaxial or abaxial signatures (Table S6), which may provide insight into physiological processes prioritized by stomata originating from these different surfaces of the leaf (Mott, 2007).



**Figure 3. Cell cycle regulators exhibit distinct, yet overlapping expression profiles. (legend on next page)**

**Figure 3. Cell cycle regulators exhibit distinct, yet overlapping expression profiles.** (a) Schematic of the cell cycle highlights the regulatory roles of cyclin-dependent kinases (CDKs) and cyclins (CYCs), as well as the G1/S-phase checkpoint regulator RETINOBLASTOMA-RELATED PROTEIN 1 (RBR1). (b-e) UMAPs depict gene expression profiles of cell cycle regulators within the developing leaf atlas. Colors represent graph-based clustering with resolution 1. (b) Core regulators of G1/S phase. (c) CDKB/CYCB markers of G2/M phase, along with distinct expression profiles of CYCAs. While CYCA2;3 expression is relatively broad, CYCA3;1 is likely expressed only in G1/S phase, along with the epidermal transcriptional regulator MUTE. (d) Co-expression of CYCD regulators with the phloem transcriptional regulator APL (black arrowheads) and the procambium/xylem transcriptional regulator TMO5 (white arrowhead). (e) Co-expression of CYCDs and WEE1 with the epidermal transcription factors FAMA (black arrowhead) and MUTE (white arrowhead).

### Cell cycle regulators exhibit distinct, yet overlapping expression profiles

We next asked whether comparisons between leaf tissues can serve to define distinct and overlapping features of common regulatory modules. We hypothesized that common modules integrate with and are controlled by different tissue-specific contexts. One such critical, conserved module is the cell cycle. Because the expression signature for G1/S or G2/M phases can overwhelm other features of cell identity, we originally regressed out the cell cycle signature from our analysis (Figure S1). However, plant genomes often encode several core cyclin-dependent kinase (CDK)/cyclin regulators, which can exhibit lineage-specific expression and function (Adrian et al., 2015; Gutierrez, 2016; Han et al., 2018; Sozzani et al., 2010; Weimer et al., 2018). An active area of research aims to understand how cell cycle and differentiation processes are coordinated.

In the leaf atlas, we found that core regulators exhibit distinct, yet overlapping, expression within vasculature, mesophyll, and epidermal cell states (Figure 3A, Figure S5). G1/S-phase transition genes *CYCLIN-DEPENDENT KINASE A;1* (*CDKA;1*), *CYCLIN D3;2* (*CYCD3;2*), and *RETINOBLASTOMA-RELATED PROTEIN 1* (*RBR1*) were broadly expressed (Figure 3B), while members of the G2/M-phase *CDKB* and *CYCB* gene families were expressed in distinct clusters (Figure 3C). Of note, we found that *CYCD* gene family members exhibited both broad and tissue-specific expression profiles (Figure 3D-E). While the gene *CYCD3;2* was broadly expressed, *CYCD3;3* and *CYCD2;1* were selectively enriched in distinct mature phloem clusters that expressed the transcriptional regulator *APL*. Conversely, *CYCD6;1* was co-expressed with the procambium/xylem bHLH transcriptional regulator *TMO5*. Onset of *CYCD7;1* in the epidermis exclusively overlapped with the transcriptional regulator *MUTE* and extended into a domain shared with the emergence of *FAMA* expression. *CYCD5;1* was broadly detected in a few cells, and may overlap with *CYCD6;1* and *CYCD7;1* in the vasculature and epidermis, respectively. Interestingly, the kinase *WEE1* is expressed in the epidermal and vascular clusters where restricted *CYCDs* converge (Figure 3E). *WEE1* is a negative regulator of the G2/M transition and ensures size control in yeast (Nurse and Thuriaux, 1980). In *Arabidopsis*, *WEE1*'s role has been described only in response to DNA damage (Cools et al., 2011; De Schutter et al., 2007; Sorrell et al., 2002). *CYCA* gene family members also displayed a range of expression profiles within the atlas (Figure 3B). For instance, while *CYCA2;3* was broadly expressed, the G1/S-phase *CYCA3;1* (Takahashi et al., 2010) was selectively co-expressed with *MUTE*. A closer examination of cell cycle genes revealed that *MUTE* may coordinate with G1/S-phase regulators, while *FAMA* coordinates with G2/M-phase regulators. (Figure 3C, 3E). This is consistent with proposed roles of these transcription factors in driving cell

cycle progression (Han et al., 2018; Ohashi-Ito and Bergmann, 2006), though our findings also raise the possibility that *MUTE* and *FAMA* gene expression is reciprocally responsive to cell cycle state.

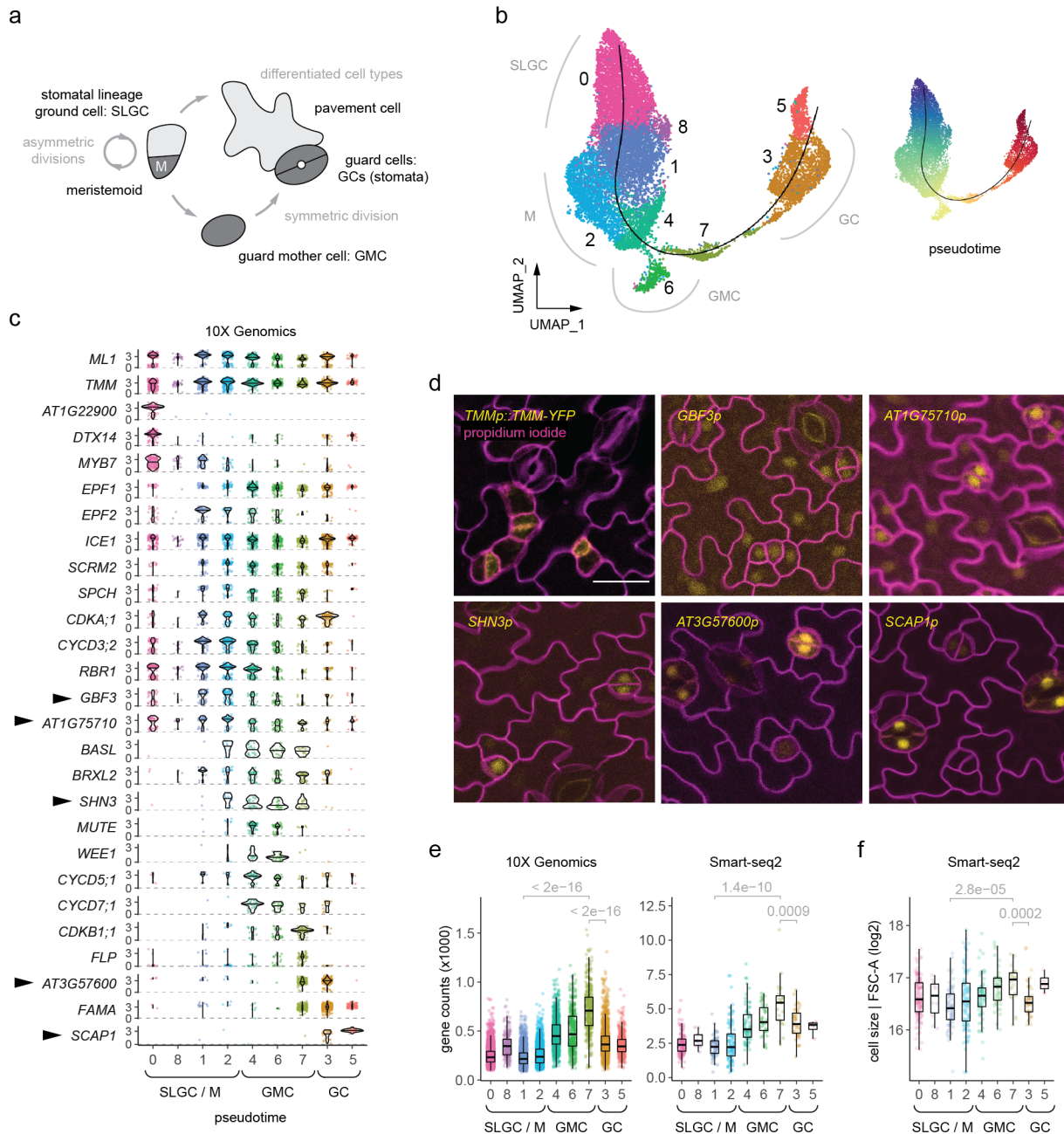
### Integrated analysis elucidates flexible stomatal lineage cell states along a continuum

As distinct models of cell state differentiation were observed in our scRNA-seq leaf atlas (Figure 2), we next aimed to further elucidate genetic programs within the well-characterized and tractable stomatal lineage. We performed scRNA-seq on FACS-isolated cells expressing the leucine-rich repeat (LRR) receptor-like gene *TOO MANY MOUTHS* (*TMM*; *TMMp::TMM-YFP*) (Nadeau and Sack, 2002). *TMM* is highly expressed in stem cells of the stomatal lineage, including both meristemoids and stomatal lineage ground cells (SLGCs). Such cells are defined as flexible given that they are able to either asymmetrically divide to yield another meristemoid/SLGC pair or differentiate into distinct epidermal cell types (Nadeau and Sack, 2002) (Figure 4A). SLGCs eventually undergo endoreduplication and differentiate into pavement cells, though this mature cell type is not captured in our datasets here due to cell size selection. Conversely, meristemoids differentiate into guard mother cells (GMCs), which divide symmetrically to yield a pair of stomatal guard cells (GCs). While decisions to divide or differentiate are known to be regulated by cell-cell signaling cascades, cell polarity, and transcriptional control (Lee and Bergmann, 2019), previous studies focused on discrete major fate transitions. The extent of cell state heterogeneity across flexible cell identities of meristemoids and SLGCs remains unclear.

We leveraged an integrated scRNA-seq approach focused solely on stomatal lineage cells to address characteristic and generalizable developmental features displayed by the stomatal lineage. These included a potential continuum of early lineage cell states and possible relationships among cell size, transcriptional diversity, and cell commitment during GMC fate commitment. To take advantage of a variety of computational analyses, some that require many cells and others that require deep sequence coverage, we used two different technologies to profile *TMMp::TMM-YFP* expressing young stomatal lineage cells. Cells captured for droplet-based 10X Genomics (Zheng et al., 2017) yielded transcriptomes for 12,933 cells, with a median of 353 detected UMIs and 271 genes. This dataset was integrated with another we generated by plate-based Smart-seq2 (Picelli et al., 2014), which yielded transcriptomes for 478 cells, with a median of 2,786 detected genes. After integration with Seurat v3 (Butler et al., 2018), potential cell states were identified through graph-based clustering and differential gene expression analysis (Figure S6-S7, Figure 4B, Table S7). To identify a putative lineage trajectory model that reflects transitions between cell states, we

reconstructed pseudotime with simultaneous principle curves using slingshot (Street et al., 2018). Compared with other methods, slingshot offers relatively more stringent predictions of putative transitions and has been previously identified as a high-performing

program (Saelens et al., 2019). Depicted as a smooth curve overlaid on the UMAP, the pseudotime axis revealed potential developmental relationships between cell states.



**Figure 4. Stomatal lineage flexibility defined as a continuum of cell states.** (a) Model of stomatal lineage highlights that early stem cells, meristemoids (Ms) and stomatal lineage ground cells (SLGCs), either divide asymmetrically (left) or commit to differentiation (right). Differentiation programs ultimately yield guard cells (GCs) and pavement cells. (b) UMAPs of integrated scRNA-seq (10X Genomics v3, 12,933 cells; and Smart-seq2, 478 cells) derived from *TMMp::TMM-YFP*-expressing shoot tissue from 10 dpg seedlings. Colors depict graph-based clustering with resolution 0.4 (left) and the relative pseudotime scale (right). Overlaid on each UMAP is the pseudotime axis (black line). (c) Violin plots depict a range of gene expression profiles within pseudotime, aligned with cluster resolution 0.4 (colors). Included are known regulators of the lineage, as well as genes for which reporters were generated (arrowheads). Pseudotime is predicted to initiate with *SPCH* expression, and it represents cell state transitions from Ms to SLGCs in one direction (towards left), as well as transitions from Ms to GMCs and GCs (towards right). (d) Confocal images of promoter-driven YFP-NLS reporters corroborate predicted expression profiles. The abaxial epidermis was imaged from first true leaves of 10 dpg seedlings. All images were taken at the same magnification (scale bar: 20  $\mu$ m). (e) Boxplots depict transcriptional diversity within pseudotime, based on total gene counts from

respective datasets. (f) Boxplots depict relative cell size (log<sub>2</sub> values of the median forward scatter area, FSC-A) within pseudotime. FSC-A was measured and indexed by FACS, corresponding to the Smart-seq2 dataset. Colors in (e, f) represent cluster resolution 0.4, and clusters are aligned with pseudotime, as in (c). Adjusted p-values in (e, f) are from pairwise comparisons using the Wilcoxon rank sum test.

Cell states were annotated using a similar approach as with the leaf atlas, assessing expression profiles of genes that encode known players in the lineage, along with genes that display dynamic profiles across the clusters (Figure 4C). Flexible cell states early in the lineage (meristemoids and SLGCs) indeed appeared to exist along a continuum within pseudotime, defined by early expression of *SPCH* and *INDUCER OF CBF EXPRESSION1 (ICE1)/SCREAM (SCRM)* bHLH transcriptional regulators (Kanaoka et al., 2008). Separate and distinct clusters represented transitions through deterministic guard cell fate commitment, with the expression of *MUTE* followed by *FAMA*. Surprisingly, *SPCH* expression extended into the expression domains of *MUTE* and *FAMA*, which was unexpected given functional genetic analyses placing *SPCH* early in stomatal lineage progression (Davies and Bergmann, 2014; MacAlister et al., 2007; Pillitteri et al., 2007). We further validated our lineage model of cell state dynamics *in vivo*. We created reporters of genes with no known function in the lineage, that displayed both broader and specific expression within pseudotime (Figure 4D, Figure S8). Imaging these reporters in 10 dpg leaves corroborated expression profiles from the scRNA-seq lineage trajectory, which indicates that our pseudotime model of cell states may reflect *in vivo* dynamics.

#### Transcript diversity correlates with cell size measured by FACS

We next asked whether cell states within our lineage pseudotime could be leveraged to uncover mechanisms for GMC fate commitment, a deterministic directed process that promotes one symmetric division event to yield a pair of stomatal guard cells. We found that the continuum of early lineage cell states included the onset of GMC commitment (Figure 4B-C). This indicates that there may be a gradual, stochastic process by which meristemoids differentiate. We were thus especially intrigued to discover increased transcriptional diversity within differentiating GMCs (Figure 4E), as this feature of lineage progression has been recently described in a range of animal and human systems (Gulati et al., 2020). Furthermore, stomatal lineage cells range in size, and it remains unclear how this is regulated. We therefore examined whether increased transcriptional diversity was correlated with cell size. Using FACS forward scatter data with indexed cells captured in our Smart-seq2 scRNA-seq analysis, cell size was positively correlated with transcriptional diversity; it gradually increased with the onset of GMC fate commitment in pseudotime, with a drop-off in size after the inferred symmetric division (Figure 4F). This is in contrast to the early lineage clusters where a range of cell sizes were detected without a clear correlation with transcriptional diversity, as expected for flexible cell states among asymmetrically dividing cells.

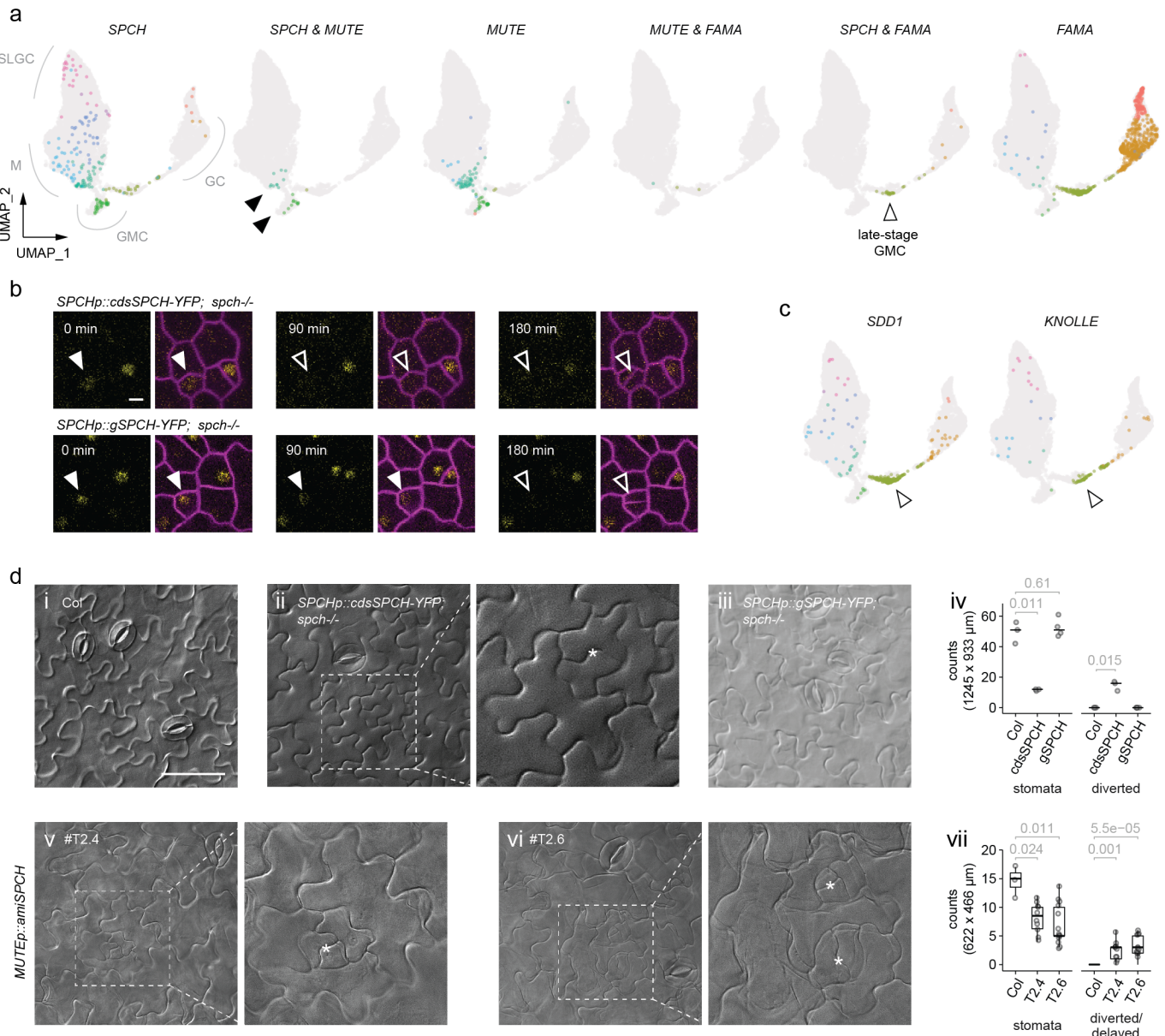
Given the observation of increased transcriptional diversity, we asked whether specific transcription factors and chromatin remodelers were expressed during the onset of GMC fate commitment to modulate the transcriptional landscape. As seen in

here (Figure 4C, 4F), *MUTE* expression is detected in GMCs of all sizes, but *FAMA* is limited to only the largest GMCs. In addition to activators iteratively expressed to gradually activate downstream programs, we might expect a requirement for negative regulators of the lineage progression repressors. We therefore identified putative transcriptional regulators that were differentially expressed within pseudotime (Table S7). Our stomatal lineage scRNA-seq dataset thereby reveals potential candidates that will serve as a framework for future studies to examine underlying mechanisms that lock flexible cell states into a given deterministic fate.

#### Broadened functional role of an early-lineage core regulator

A paradigm of *Arabidopsis* stomatal development is that sequential expression of *SPCH*, *MUTE*, and *FAMA* defines discrete transitions of cell fates (MacAlister et al., 2007; Ohashi-Ito and Bergmann, 2006; Pillitteri et al., 2007). However, there are indications from our lineage pseudotime described here (Figure 4B-C) that developmental progression is more complicated. In our gene expression overlays of *SPCH*, *MUTE* and *FAMA*, expression of *SPCH* persisted beyond early stages of the lineage (Figure 4C). Distinct cell states even appear to co-express *SPCH* with either *MUTE* or *FAMA* (Figure 5A). Given the potential conflict of these findings with the sequential model of the lineage, we reexamined *SPCH* expression *in vivo* within dividing GMCs by time-lapse imaging. We imaged a previously published reporter (coding sequence *SPCHp:cdsSPCH-YFP*, *cdsSPCH*) (Davies and Bergmann, 2014; Vatén et al., 2018), as well as a new reporter that contained more of the *SPCH* genomic context (genomic sequence *SPCHp:gSPCH-YFP*, *gSPCH*), which may more accurately recapitulate regulatory control at the endogenous locus. Each reporter was characterized in a *spch-3* (null) background. Expression of the *SPCH* reporters indeed extended beyond early stage, asymmetrically dividing cells (in 4 dpg cotyledons), with stronger signal potentially detected using the *gSPCH* genomic reporter (Figure 5B). This ultimately corroborates our new model based on scRNA-seq and suggests that *SPCH* serves an unappreciated role within dynamic cell states.

To examine the diversity of potential modules regulated by *SPCH*, we identified putative target genes from ChIP-seq (Lau et al., 2014) and enriched gene ontologies across cell states in the stomatal lineage scRNA-seq dataset. We found that many differentially expressed genes may be directly regulated by *SPCH* (i.e. bound by *SPCH*), and we thereby detected similar enriched modules from gene ontology, spanning from stimulus and immune responses to regulation of growth and division processes (Table S7). We also identified genes that correlate with induced gene expression of *SPCH*, by leveraging a previous RNA-seq dataset (Lee et al., 2019). Differentially expressed genes in the clusters that are bound by *SPCH* and positively correlate with induced *SPCH* expression were generally enriched in early lineage state clusters (Table S7). Of these, within the late-stage GMC cluster,



**Figure 5. SPCH is expressed and required beyond the early stages of the lineage.** (a) Extended SPCH gene expression is depicted. UMAPs indicate cells and clusters (resolution 0.4) that express combinations of the genes SPCH, MUTE, and FAMA. Clusters that express both SPCH and MUTE (black arrowheads) are distinct from the cluster that expresses SPCH and FAMA (white arrowhead). (b) Time-lapse confocal images of *SPCH<sub>p</sub>::cdsSPCH-YFP* (protein-coding sequence) and *SPCH<sub>p</sub>::gSPCH-YFP* (genomic sequence), each complementing *spch-3*. Images are from the abaxial epidermis of 4 dpg seedlings, and arrowheads follow the GMC division that yields young stomatal guard cells (at 180min). Outside the frame of the time-lapse, young guard cells are presumed to enlarge and form a pore. Filled white arrowhead: reporter expression detected, outlined arrowhead: no expression detected. All images were taken at the same magnification (scale bar: 5  $\mu$ m). (c) Examples of genes, and their respective UMAP expression profiles (colors: cluster resolution 0.4), that may be regulated by SPCH beyond early stages of the lineage. Putative state-specific direct targets were found to be bound by SPCH and differentially expressed within pseudotime. (d) Diverted cell fates (asterisks: borders of diverted/delayed cell identities) in representative differential interference contrast (DIC) images of abaxial cotyledon epidermis from Col wildtype control (i), *SPCH<sub>p</sub>::cdsSPCH-YFP* *spch-/-* (ii), *SPCH<sub>p</sub>::gSPCH-YFP* *spch-/-* (iii), and two independent lines of *MUTE<sub>p</sub>::amiSPCH* T2.4 (v) and T2.6 (vi). Images were all taken at 15 dpg with the same magnification (scale bars: 100  $\mu$ m), except for insets demarcated by white dashed boxes. Diverted/delayed fate events are quantified in (iv, vii). Diverted fates are defined by presumed exit of the stomatal lineage, wherein sister cells undergo lobing – rather than give rise to a meristemoid and SLGC pair or generate a pair of guard cells. Early lobing of an SLGC is depicted in (vi, top asterisk). *MUTE<sub>p</sub>::amiSPCH* epidermis also displayed delayed lineage commitment, defined by putative additional rounds of asymmetric divisions before the commitment to yield guard cells. Plots in (iv, vii) include grey points that each represent counts from an individual seedling (at least n=3 seedlings were measured per genotype). median: black line. p-values: two-sample t-test for respective comparisons except for diverted events in (iv), wherein the p-value represents one-sample t-test.

we found that *SPCH* appears to control genes implicated in cell fate decisions and cell cycle control (Figure 5C), which include the protease gene *STOMATAL DENSITY AND DISTRIBUTION 1* (*SDD1*) (von Groll et al., 2002) and the cytokinesis-specific syntaxin gene *KNOLLE* (Lauber et al., 1997; Lukowitz et al., 1996). Conversely, genes that negatively correlate with *SPCH* expression were detected in the guard cell clusters. As *SPCH* function likely shifts within different cellular states, it is interesting to consider the potential new role of *SPCH* in GMCs, where it appears to sequentially overlap with both *MUTE* and *FAMA* expression.

Given that we corroborated extended expression of *SPCH* and identified potential modules putatively controlled by the regulator across cell states, we tested whether extended expression of *SPCH* was required for proper cell identity and fate decisions. The first hint that *SPCH* might contribute to later differentiation steps came from examination of mature tissue (15 dpg cotyledons) in lines where the *spch-3* null mutant was rescued by *cdsSPCH* or *gSPCH*. While *gSPCH* leaves exhibited similar stomatal counts and broad cellular features compared with wildtype, *cdsSPCH* leaves had fewer stomata, and displayed a novel phenotype, which we call “diverted” stomatal lineage cells based on morphology and position relative to other epidermal cells (Figure 5D, Figure S9). Diverted cells undergo lobing, rather than yield either meristemoid and SLGC pairs or pairs of guard cells. This finding could be due to disrupted levels and/or timing of *SPCH* expression, particularly since the *cdsSPCH* transgene was more lowly expressed at later stages relative to *gSPCH* (Figure 5B). We therefore examined whether *SPCH* is specifically required at this later developmental transition by selectively depleting late-stage expression with the *MUTE* gene promoter driving artificial microRNAs against *SPCH* (*MUTEP::amiSPCH*). With these lines, we identified two phenotypes: a delay in commitment (as evidenced by additional rounds of early lineage divisions before producing stomata) and diverted cell identities, similar to what we found in the *cdsSPCH* rescue lines (Figure 5D, Figure S9). Diverted and delayed cellular developmental outcomes were not observed in control seedlings. Overall, this functional approach indicates that fine-tuned expression of *SPCH* is required to control and promote late cell fate commitment in GMCs, and thereby, along with proposed modules from functional genomics, reveals a new role for the critical transcriptional regulator.

## DISCUSSION

Plants display extreme biological flexibility that offers insight into exceptional developmental strategies. Here, we uncovered distinct models of cell state differentiation within *Arabidopsis* leaf tissue by leveraging single-cell transcriptomics and functional molecular genetics. A comparative analysis of cell states in the mesophyll, vasculature, and epidermal clusters revealed tissue-specific modules and enriched cellular programs, along with adaxial/abaxial tissue polarity signatures. Our scRNA-seq atlas of the developing leaf thereby serves as a useful resource to pursue questions and generate hypotheses that will unravel how these tissues arise. We also identified underlying heterogeneity within cell states of the flexible epidermal stomatal lineage, which appear to exist along a continuum, with progressive cell specification. Our findings indicated that the core early transcriptional regulator

*SPCH* potentially controls a range of cellular programs within the lineage continuum. Early roles may be foundational to adjusting stomatal lineage expansion in line with environmental fluctuations (Kumari et al., 2014; Lau et al., 2018). scRNA-seq analysis showed, unexpectedly, that *SPCH* was expressed beyond the early stages of the lineage, and we demonstrated that it was required for cell fate commitment to yield stomatal guard cells. Overall, our work has refined the stomatal lineage paradigm.

*SPCH* serves an unappreciated role in promoting stomatal guard cell fate, which is in conflict with the previous model of sequential and discrete control by *SPCH*, *MUTE*, and *FAMA* (MacAlister et al., 2007; Ohashi-Ito and Bergmann, 2006; Pillitteri et al., 2007). When *SPCH* was specifically depleted by *amiRNAs* driven by the *MUTE* promoter, we detected extended asymmetric amplifying divisions that may indicate a failure to commit and a delay in lineage progression. We also observed that some divisions, rather than resulting in a guard cell pair, yield a pair of lobed cells that are presumed to have diverted from the guard cell fate genetic program. Extended amplifying divisions have also been detected in the *mute* null, though they represent a distinct phenotype, as such divisions are defined by rounded cells within *mute* leaves (MacAlister et al., 2007; Pillitteri et al., 2007). Diverted cell fates as observed in the *MUTEP::amiSPCH* leaves have not been described for other loss-of-function mutations, but ectopic overexpression of the secretory peptide gene *EPIDERMAL PATTERNING FACTOR 1 (EPF1)* leads to a similar phenotype (Hara et al., 2009, 2007; Triviño et al., 2013). Consistent with this, EPF1 represses *SPCH* function through kinase signaling cascades (Lampard et al., 2008). Thus, in our new model of the stomatal lineage, we postulate that *SPCH* serves dynamic roles during early flexible cell states, and also complements *MUTE* and *FAMA* at later stages to drive cell fate commitment and differentiation of guard cells. Signaling and regulatory modalities that selectively implicate *SPCH* may be recruited into these later stages. For example, when *Arabidopsis* is subjected to environmental stresses (e.g. heat, starvation, or drought), the stomatal lineage response has centered on transcriptional and post-translational control of *SPCH* (Han et al., 2020; Kumari et al., 2014; Lau et al., 2014). The preference for *SPCH* was often ascribed to its role in early, flexible development. However, *SPCH* protein also has unique domains that render it a partner or target of regulatory proteins that *MUTE* and *FAMA* cannot engage (Lampard et al., 2008; Putarjunan et al., 2019), and thus it may be that *SPCH* is necessary to transmit such environmental information to later decisions.

Our work also highlighted distinct and overlapping features of common regulatory modules within a developing leaf. We found that cell cycle regulators integrated with tissue-specific regulatory contexts and may coordinate cell fate. For example, we see parallels to the stomatal lineage in the vasculature cluster, two places where cell states are dependent on complexes containing RBR1 and a lineage-specific transcription factor (*FAMA* and *SCARECROW (SCR)*, respectively) (Cruz-Ramírez et al., 2012; Matos et al., 2014). We found that there may be shared regulatory logic between CYCD5;1, WEE1, and CYCD6;1/CYCD7;1 in these decisions as well. CYCD6;1 promotes asymmetric divisions of the root cortex-endodermal initials and is a direct target of a complex

containing SCR and SHORTROOT (SHR) (Sozzani et al., 2010). The rapid upregulation of the *CYCD6;1* gene in response to SHR induction in roots is well characterized, though less appreciated may be the concomitant upregulation of *WEE1* and *CYCD5;1* (Sozzani et al., 2010). Moreover, *SHR* is required for leaf growth in part through upregulation of genes that promote the G1/S transition, including *CYCD5;1* (Dhondt et al., 2010). In the stomatal lineage, MUTE targets *CYCD5;1*, and *WEE1* is also upregulated upon induced *MUTE* expression (Han et al., 2018). However, the stomatal lineage-restricted *CYCD7;1* is not a target of MUTE (Han et al., 2018; Weimer et al., 2018). *CYCD5* and *WEE1* may thus be induced by core transcriptional regulators to regulate formative divisions, but there are also independent routes of tissue-specific expression, as evidenced by *CYCD7;1*.

It is not clear why certain cell states employ additional core cell cycle regulators. In the stomatal lineage, loss of either *CYCD5;1* or *CYCD7;1* results in the accumulation of larger GMCs and a delay transiting into guard cell fate (Han et al., 2018; Weimer et al., 2018). Loss-of-function *WEE1* phenotypes have only been measured in response to DNA damage in roots, but include a prolonged cell cycle and premature differentiation into vascular cells (Cools et al., 2011). If the role of *WEE1* is consistent among cell types, then a plausible outcome for cells expressing both specialized CYCDs and *WEE1* would be the production of larger and more transcriptionally active cells, a situation we observed in the profiled stomatal lineage cells, and that may be required during other formative divisions. The cell cycle represents just one regulatory module, and it will be interesting to leverage our developing leaf atlas to examine signaling cascades and complementary modules. Moreover, a comparison of vasculature cell states in the leaf with that of the stem and root may resolve robust programs that control such tissue-specific differentiation both above and below ground, as vasculature threads throughout the plant body. Finally, detecting *WEE1* expression in scRNA-seq data, when it has often been undetectable (Cools et al., 2011; Dhondt et al., 2010), bodes well for being able to identify other conserved modules that evaded previous detection due to low or highly restricted expression patterns.

Single-cell transcriptomics enabled cell states within the developing leaf to be defined by putatively dynamic instances of a given cell transition or identity, corresponding to cell function and lineage relationships (Morris, 2019; Sagar and Grün, 2020; Wagner et al., 2016). This ultimately provides a framework to compare molecular mechanisms of specific cell fate transitions in plant development between species, as well as with animal development. It will be exciting to consider evolutionarily divergent and convergent deployment of specific molecular regulators and developmental strategies. Furthermore, complementary genomics and proteomics approaches will help shape our understanding of underlying cellular modules and programs, such as by providing insight into gene regulatory networks and protein-protein interactions.

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## AUTHOR CONTRIBUTIONS

C.L.A. conceived, designed, performed all experiments (except as noted below), analyzed results, performed statistical analysis, and wrote the manuscript. A.V. created the *MUTEp::amiSPCH* and *SPCHp::gSPCH-YFP* genomic *SPCH* rescue lines. M.X.A.G. contributed to imaging *SPCH* rescue lines. N.K.S., N.S., and V.G. contributed to creation and imaging of reporter lines. Y.G. performed time-lapse imaging. A.K.W. provided cell cycle insight. D.C.B. analyzed results and wrote the manuscript.

## DECLARATION OF INTERESTS

The authors declare no competing interests.

## MATERIALS AND METHODS

**Plant material and growth conditions.** All transgenic lines were examined in the *Arabidopsis thaliana* Columbia-0 (Col) ecotype, which served as the wildtype control in all experiments. The following previously described transgenics were used in this study: *ML1p::YFP-RCI2A* (Roeder et al., 2010), *TMMp::TMM-YFP* (Nadeau and Sack, 2002), *SPCHp::cdsSPCH-YFP* and *SPCHp::gSPCH-YFP* (Vatén et al., 2018). Seedlings were grown on half-strength ( $\frac{1}{2}$ ) Murashige and Skoog (MS) medium (Caisson Labs MSP01) at 22°C under long-day conditions (16 hr light and 8 hr dark cycles) and were used at the indicated day post germination (dpg) for respective experiments.

**Vector construction and plant transformation.** Transgenic promoter reporters fused to a nuclear localized version of YFP (YFP-NLS) were generated using the Gateway system (Invitrogen). In brief, promoters were PCR-amplified from genomic DNA and cloned into pENTR then recombined with vectors bearing YFP-NLS, into the binary R4pGWB destination vector system (Nakagawa et al., 2008; Tanaka et al., 2011). To generate the *MUTEp::amiSPCH* transgene, the *artificial microRNA* (*amiRNA*) sequence was designed with the Web MicroRNA Designer platform (<http://wmd3.weigelworld.org>), engineered using the pRS300 plasmid as template, and together with the *MUTE* promoter, was subcloned into the destination vector pGWB. Primer

sequences used to generate transgenics are provided in [Table S2](#), and all referenced genes are included in [Table S3](#). Transgenic plants were generated by Agrobacterium-mediated transformation (Clough, 2005; Clough and Bent, 1998), and seedlings were selected on ½ MS plates supplemented with respective antibiotics (e.g. 12 mg/L Basta).

#### Analysis of transcriptional and translational reporters.

Seedlings from at least two independent lines were collected and analyzed, as indicated for each respective experiment. Fluorescence images were taken with a Leica TCS SP5 microscope (Leica Microsystems) and processed with Fiji (Schindelin et al., 2012). Cell outlines were visualized by either the plasma membrane marker *ML1p:mCherry-RCI2A* or with propidium iodide (Molecular Probes, P3566).

**Quantification of developmental phenotypes.** Seedlings from at least two independent and homozygous lines were collected, except for the two independent, segregating lines of *MUTEp::amiSPCH*. Seedlings were cleared in 7:1 ethanol:acetic acid, treated for 30 min with 1 N potassium hydroxide, rinsed in water, and mounted in Hoyer's medium. Differential contrast interference (DIC) images of the middle region of adaxial epidermis of cotyledons were obtained with a Leica DM2500 microscope (Leica Microsystems).

#### Single-cell (sc)RNA-seq and data analysis

**Single-cell isolation.** Single-cells (i.e. protoplasts, plant cells without cell walls) from respective transgenic reporter lines were isolated as described previously (Adrian et al., 2015; Bargmann and Birnbaum, 2010). In brief, whole aerial tissue or first true leaves harvested from ~180-260 total 10 dpg seedlings were placed into 15 mL protoplasting solution: pH 5.7 with 1 M Tris-Cl pH 7.5, contains 1.25% [w/v] cellulase (Yakult R-10), 0.3% [w/v] macerozyme (Yakult R-10), 0.4 M mannitol (Sigma M1902), 20 mM MES (Caisson Labs M009), 20 mM KCl, 0.1% BSA (Fisher BP1605), and 10 mM CaCl<sub>2</sub>. After 2 hours of gentle shaking, the protoplast solution was filtered through a 40 µm cell strainer (BD Falcon 352340) and centrifuged for 5 min at 500 x g. Protoplast pellets were resuspended in ~1-3 mL of protoplasting solution and sorted using a FACSAria II (BD Biosciences) instrument fitted with a 100 µm nozzle.

**Droplet-based 10X Genomics.** 300,000 single-cell positive events were sorted into ~1-2mL of protoplasting solution. Kept at room temperature, cells were lightly centrifuged and resuspended in protoplasting solution to yield a sample with ~2,000 cells/µL in a total volume of ~20-30 µL. Cell concentration was determined with a hemocytometer, and limited cell lysis was detected by staining with 1:1 trypan blue (Thermo Scientific 12250061). Cells were loaded into a microfluidic device (10X Genomics) with 3' v2 or v3 chemistry to capture ~5,000-10,000 cells per sample (Zheng et al., 2017). mRNA was reverse transcribed and cDNA quality was assessed using High Sensitivity NGS Fragment Analyzer (Agilent). Illumina libraries were constructed with Gene Expression v2 and v3 kits (10X Genomics), and sequencing of paired-end 75 bp reads was performed on a HiSeq4000 (Illumina), according to the manufacturer's instructions. Illumina BCL files were processed via

Cell Ranger v2.1.1 and v3.1.0 pipelines, with reads aligned to the Arabidopsis TAIR10 genome assembly (Lamesch et al., 2012). Subsequently, spliced and unspliced transcript matrices were generated using samtools (Li et al., 2009) and velocity (La Manno et al., 2018).

**Plate-based Smart-seq2.** Single-cell positive events were indexed and sorted directly into single wells of a 96-well plate, which contained 4 µL of lysis buffer with 1 U/µL of Recombinant RNase Inhibitor (RRI, Clontech 2313B), 0.1% [w/v] Triton X-100 (Thermo Scientific 85111), 2.5 mM dNTP (Thermo Scientific 10297018), and 2.5 µM oligoDT30VN (5'AAGCAGTGGTATCAACGCAGAGTACT30VN-3', IDT). Once sorted, cells were immediately spun down and frozen at -80°C for temporary storage. cDNA synthesis was performed as previously described using the Smart-seq2 protocol (Picelli et al., 2014), with SMARTScribe (Clontech 639538) reverse transcriptase and KAPA HiFi DNA polymerase (Kapa Biosystems KK2602). cDNA quality was assessed using High Sensitivity NGS Fragment Analyzer Kit (Advanced Analytical DNF-474), and barcoded Illumina libraries were made using the miniaturized Nextera XT protocol (Mora-Castilla et al., 2016). Sequencing of single-end 75 bp reads was performed on a NextSeq500 (Illumina) high output flow cell, according to the manufacturer's instructions. FASTQ files were generated from Illumina BCL files and low-quality sequences were trimmed by cutadapt (Marcel, 2011). Reads were aligned to the TAIR10 genome assembly (Lamesch et al., 2012) with STAR (Dobin et al., 2012), and a gene-by-cell raw count matrix was generated by HTSeq (Anders et al., 2015).

**scRNA-seq dataset analyses.** Transcripts Per kilobase Million (TPM) values (Smart-seq2) or gene-barcode matrices (10X Genomics) from single cells were analyzed with Seurat v3 (Butler et al., 2018). Data processing and plotting was done with R/RStudio (R Core Team, 2019; RStudio Team, 2019), except for as noted otherwise. Cells were pre-processed to filter out poor-quality cells with relatively few detected transcripts (i.e. less than 100-500 UMIs/cell), as well as potential cell doublets (i.e. two cells with a shared cell barcode identifier) with relatively high numbers of detected genes. Cells with substantial detected mitochondria and chloroplast transcripts (i.e. more than 25% of UMIs attributed to respective transcripts) were also filtered out, under the assumption that such profiles may represent plastids released from cells.

After pre-processing, the data were log-normalized by NormalizeData and variable genes were identified with FindVariableGenes, with parameters selection.method = "vst" and nfeatures = 2000. For the *TMMp::TMM-YFP* datasets, the n=3 samples from 10X Genomics were integrated before subsequent integration with the Smart-seq2 dataset, using FindIntegrationAnchors and IntegrateData. Unwanted sources of variation derived from cell cycle stage across different tissues were regressed out of the *ML1p::YFP-RCI2A* dataset to avoid clustering based on cell cycle (Tirosh et al., 2016). Cell cycle stage genes were defined by known homology and previous time-course experiments (Menges et al., 2005, 2003; Vandepoele et al., 2002). Representative genes induced during S and G2/M phases ([Table S4](#)) were used to assign 'cell cycle scores' with CellCycleScoring,

which were used in downstream cell cycle regression. The data were then scaled with ScaleData. Principle component analysis (PCA) dimensionality reduction was performed with RunPCA to calculate 50 principal components. A Shared Nearest Neighbor (SNN) graph was generated by FindNeighbors, and graph-based cell clustering based on the SNN was performed by FindClusters. The clustree package (Zappia and Oshlack, 2018) was implemented to visualize how clusters breakdown from one resolution to another. Uniform Manifold Approximation and Projection (UMAP) embedding was performed by RunUMAP, using all 50 principal components with parameters n\_neighbors = 30, min\_dist = 0.3, umap.method = "uwot", and metric = "cosine". Data visualization was performed with ggplot2 (Wickham, 2016).

Differentially expressed genes in each cluster were identified by the non-parametric Wilcoxon rank sum test (default) using Seurat v3 (Butler et al., 2018) FindAllMarkers, with parameters only.pos = "TRUE", min.pct = 0.25 and logfc.threshold = 0.223 (1.25-fold change). The min.pct argument requires that genes are detected within a minimum percentage (i.e. 25%) of cells of a given cluster, and the logfc.threshold argument requires a minimum fold change (i.e. 1.25-fold) for a gene to be identified as differentially expressed. Enriched GO terms associated with the top 100 differentially

expressed genes were identified by clusterProfiler (Yu et al., 2012) compareCluster and gofilter (level = 3) with the following parameters fun = "enrichGO", OrgDb = "org.At.tair.db", ont = "BP", pAdjustMethod = "BH", pvalueCutoff=0.05, qvalueCutoff=0.05.

Transcriptional dynamics (i.e. predicting future cell states by leveraging unspliced and spliced transcripts) in the *ML1p::YFP-RCI2A* 10X Genomics dataset was assessed by a steady-state deterministic model using scVelo (Bergen et al., 2019), with scv.pp.filter\_and\_normalize parameters min\_shared\_counts = 100 and min\_counts\_u = 500. scVelo was implemented with python using Jupyter/Python Notebook (Perez and Granger, 2007; Van Rossum and Drake, 2009). Pseudotime lineage inference was performed with the integrated *TMMp::TMM-YFP* datasets in R/RStudio using Slingshot (Street et al., 2018), which constructs a minimum spanning tree and fits simultaneous principle curves through the tree, with the parameter reduceDim = "UMAP".

**Accession numbers.** The scRNA-seq datasets generated in this study are deposited in GEO under accession number GSXXXXX.

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