Distinct identities of leaf phloem cells revealed by single cell transcriptomics

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Abstract

The leaf vasculature plays a key role in solute translocation. Veins consist of at least seven distinct cell types, with specific roles in transport, metabolism, and signaling. Little is known about leaf vascular cells, in particular the phloem parenchyma (PP). PP effluxes sucrose into the apoplasm as a basis for phloem loading, yet PP has been characterized only microscopically. Here, we enriched vascular cells from Arabidopsis leaves to generate a single-cell transcriptome atlas of leaf vasculature. We identified at least 19 cell clusters, encompassing epidermis, guard cells, hydathodes, mesophyll, and all vascular cell types, and used metabolic pathway analysis to define their roles. Clusters comprising PP cells were enriched for transporters, including SWEET11 and SWEET12 sucrose and UmamiT amino acid efflux carriers. We provide evidence that PP development occurs independently from ALTERED PHLOEM DEVELOPMENT, a transcription factor required for phloem differentiation. PP cells have a unique pattern of amino acid metabolism activity distinct from companion cells (CCs), explaining differential distribution/metabolism of amino acids in veins. The kinship relation of the vascular clusters is strikingly similar to the vein morphology, except for a clear separation of CC from the other vascular cells including PP. In summary, our single-cell RNA-sequencing analysis provides a wide range of information into the leaf vasculature and the role and relationship of the leaf cell types.

Introduction

A key feature of multicellularity is the division of labor. During evolution, when plants moved from aquatic to terrestrial environments, new mechanisms were required for the exchange of nutrients—both photoassimilates from aerial organs to soil-anchored sections, and water and essential nutrients from the soil to the aerial, photosynthetic organs. Terrestrial plants developed complex vascular systems to provide, for example, roots with photoassimilates and to provide a photosynthetic organism with essential materials. For this, the leaf vasculature and its intricate array of cell types are critical. It is estimated that the leaf vasculature consists of at least seven distinct cell types, each with specific roles in transport, metabolism, and signaling. However, little is known about the leaf vascular cells, in particular the phloem parenchyma (PP). PP effluxes sucrose into the apoplasm as a basis for phloem loading, yet PP has been characterized only microscopically. Here, we enriched vascular cells from Arabidopsis leaves to generate a single-cell transcriptome atlas of leaf vasculature.

Clustering of vascular cells

To identify distinct and functionally relevant cell types, we applied single-cell RNA sequencing (scRNA-seq) to enriched leaf vascular cell populations from Arabidopsis. This approach allowed us to generate a single-cell transcriptome atlas of leaf vasculature, which revealed at least 19 cell clusters. The clusters encompassed epidermis, guard cells, hydathodes, mesophyll, and all vascular cell types, including PP cells. We used metabolic pathway analysis to define the roles of these cell clusters, and found that PP cells were enriched for transporters, including SWEET11 and SWEET12 sucrose and UmamiT amino acid efflux carriers. This suggests that PP development occurs independently from ALTERED PHLOEM DEVELOPMENT, a transcription factor required for phloem differentiation. PP cells have a unique pattern of amino acid metabolism activity distinct from companion cells (CCs), explaining differential distribution/metabolism of amino acids in veins. The kinship relation of the vascular clusters is strikingly similar to the vein morphology, except for a clear separation of CC from the other vascular cells including PP.

In summary, our single-cell RNA-sequencing analysis provides a wide range of information into the leaf vasculature and the role and relationship of the leaf cell types.
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Background: Transport processes are critical for all multicellular organisms. In plants, two vascular tissues, phloem and xylem, are mainly responsible for fulfilling this task. The leaf vascular cells are embedded deep inside the leaf and were mostly identified microscopically. Cell type specific markers are rare for most vascular cell types. Especially, little is known about the phloem parenchyma, a cell type of the phloem involved in sucrose transport. To gain more profound knowledge of the transcript profiles of the leaf vascular cells, we optimized protocols to enrich vascular protoplasts (plant cells without cell walls) and performed single-cell RNA sequencing analysis.

Questions: What is the mRNA profile of different vascular cell types? Can we identify the transcriptome of the phloem parenchyma? What types of transporters are represented in different cell types? Which metabolic pathways are enriched in vascular cell types?

Findings: Our single-cell transcriptome atlas contains all cell types in the Arabidopsis leaf vasculature and non-vascular cell types such as bundle sheath, epidermis, guard cells, and mesophyll cells. We were able to identify distinct mRNA signatures which reflect vascular cell type identity. We found that the phloem parenchyma cells are enriched for various transporters, including sugar transporters and amino acid transporters. In a parallel article in this issue, we show that maize homologs of these sugar and amino acid transporters are enriched in the abaxial bundle sheath cells. These observations suggest that maize and Arabidopsis use different routes for phloem loading of sucrose and amino acids. In addition, we found unexpected roles of PP in hormone biosynthesis and defense-related responses.

Next steps: As we now have the resources at hand, our next step will be to explore the vascular cell types at the molecular level to dissect their function. Our goal is to identify specific components, for instance, transcription factors or transport proteins and contribute to the solution of the long-standing riddles: How do plants distribute nutrients across the entire plant body and how are the logistics coordinated?

The vascular parenchyma (VP) is often located at the interface between phloem and xylem. At earlier stages of development, xylem and phloem are separated by meristematic cells (procambium; open-type vasculature), that differentiate toward the tip of the leaf where procambium is absent (closed-type; Kang and Dengler, 2004). Phloem and xylem exchange water and solutes in complex ways, and thus the cells must be equipped with specific sets of transporters. Moreover, it is likely that the different cell types have specialized metabolic activities. In addition, the vasculature plays important roles in communication by translocating hormones, small RNAs, and even proteins; and also appear to be involved in electrical signaling (Nguyen et al., 2018).

PP is one of the poorly defined and least characterized vascular cell types. In Arabidopsis, PP has so-called cell wall ingrowths with a transfer cell appearance which are thought to play a role in amplifying the surface area to allow for higher transport rates (Edwards et al., 2010; Arun Chinnappa et al., 2013). Since in many plant species the interface between CC and SE (SE/CC) contains only few plasmodesmata (PD), photoassimilate translocation requires an apoplastic transport route (Haritatos et al., 2000). SE/CC loading is mediated by the H+-sucrose symporter SUT1 (originally identified in potato; named SUC2 in Arabidopsis), which imports sucrose from the cell wall space (Riesmeier et al., 1994). We recently identified sucrose uniporters of the SWEET family in a specific subset of cells in the phloem that likely represent PP (Chen et al., 2012). SWEET expression in PP was confirmed using translational GFP fusions, using a new confocal microscopy method that enabled unambiguous PP identification (Cayla et al., 2019). A key goal of this work was to characterize the role of PP in more detail by identifying its mRNA outfit. This could serve, for example, as a basis for identifying transporters involved in phloem loading of other nutrients, as well as metabolic pathways active in these cells.

Understanding the complexity of plant cells at a single cell level has long been an active area of interest in plant biology. Comprehension at this level is crucial to determine subtle distinctions between, for example, the multitude of complex vascular cell types. Recently, high-throughput single-cell RNA-sequencing (scRNA-seq) has been used to provide new insights into root cell biology and development (Denyer et al., 2019; Ryu et al., 2019; Shulse et al., 2019; Zhang et al., 2019; Wendrich et al., 2020). These latest technologies have allowed us to build on the resolution of prior atlases derived from the isolation of specific cell types, using fluorescent-activated cell sorting of fluorescently labeled protoplasts, and/or laser-capture microdissection (Birnbaum et al., 2003). To gain insights into the specific
transcript profiles of the leaf vasculature, we optimized protoplast isolation protocols to enrich for leaf vascular cells. Using scRNA-seq, we identified unique and characteristic mRNA signatures, which revealed fundamental differences in amino acid metabolism and transport pathways in PP and CC important for phloem translocation as well as differences in hormone and glucosinolate metabolism.

Results

Enrichment of vascular protoplasts

Standard protoplast isolation protocols are efficient for protoplasting mesophyll cells for downstream applications such as transient expression. However, these procedures do not efficiently release cells from the vasculature. Here, we developed a methodology for isolating vascular-enriched protoplast populations from mature Arabidopsis leaves (Figure 1; Supplemental Figure S1). Initially, a fraction of the nonvascular cell types were removed using the “tapse-sandwich” method which effectively eliminated trichomes, guard cells, and epidermis from the abaxial leaf surface (Supplemental Figure S1B; Wu et al., 2009). Cuts along the midvein further facilitated access of cell wall digesting enzymes to the vasculature (Figure 1A; Supplemental Figure S1B). Elevated concentrations of mannitol lead to an increased release of protoplasts (Figure 1B; Supplemental Figure 1, C and D). We monitored the release by light microscopy, reverse transcription- quantitative polymerase chain reaction (RT-qPCR), and fluorescence microscopy (Figure 1, D and E; Supplemental Figure 1, E and F). The mRNA levels of marker genes were used to evaluate the enrichment of vascular protoplasts during optimization of the protocols (Figure 1, A–C). In addition, the release of intact vascular protoplasts was verified microscopically by monitoring fluorescently labeled cells using stably transformed Arabidopsis lines expressing pAtSWEET11:AtSWEET11-GFP, a tentative marker for PP (Chen et al., 2012), and Q0990, a procambial (PC) marker (Radoeva et al., 2016; Figure 1, D and E; Supplemental Figure S1E). The bulk leaf (not subjected to protoplasting) transcriptome showed high correlation with bulk protoplast transcriptomes (Pearson’s correlation coefficient of 0.9; Supplemental Figure 2A and Supplemental Data Set 1). The high correlation indicates that the protoplasting protocol used here did not impact the relative abundance of cells in the leaf and supported the notion that the scRNA-seq data derived from the protoplasts should cover essentially all cell types present in a mature leaf (the term enriched here thus refers to enrichment over standard protocols).

scRNA-seq of vascular-enriched leaf protoplast population

scRNA-seq libraries were produced from vasculature-enriched leaf protoplast populations, with 5,230 single-cell transcriptomes obtained from two biological replicates. Sequencing to a depth of approximately 96,000 reads per cell was undertaken, and identified a median number of 3,342 genes, and 27,159 unique molecular identifiers (representing unique transcripts), per cell (Supplemental Figure 2B). Unsupervised clustering using Seurat (Satija et al., 2015) identified 19 distinct cell clusters (Figure 2; Supplemental Movie 1 and Supplemental Data Set 2). Plotting the transcriptomes from the two replicates in two dimensions using Uniform Manifold Approximation and Projection (UMAP; McInnes et al., 2018) revealed an overlapping distributions of cells, and a similar proportion of cell identities (Supplemental Figure 2C). mRNA profiles and relative cell numbers in the clusters were highly correlated in the replicates (Supplemental Data Set 1). To assign cell identity to the clusters, we examined the specificity of transcripts of known marker genes (Figure 3 and Supplemental Figure 3, Supplemental Data Sets 3 and 4).

The 19 clusters cover all major cell types of the leaf. The population was dominated by mesophyll cells (~75% of all cells), as indicated by the enrichment of known mesophyll markers (CAB3, LHCB2.1, and CA1; Sawchuk et al., 2008; Endo et al., 2014; Figure 3; Supplemental Figure S3A). The mesophyll group comprised 12 clusters (Clusters, or C1, 2, 3, 5, 6-9, 11, 12, 14; Figures 2A and 3; Supplemental Data Set 4) and formed three major branches on the UMAP plot. Naively, two separate mesophyll clusters can be expected, one for palisade and one for spongy parenchyma. FIL (YAB1) transcripts, markers for spongy parenchyma were enriched in C2 and C7, possibly indicating that these clusters may represent spongy parenchyma (Supplemental Figure S3A; Uemoto et al., 2018). Among the mesophyll clusters, C6 and C9 were enriched for rRNAs (Supplemental Data Set 4). We did not remove rRNA before sample preparation and did not filter cells that showed higher rRNA transcript levels since rRNA levels could vary between cells. Whether C6 and C9 represent unique biologically relevant cell populations in the leaf or are due to artifacts was not further evaluated. While the mesophyll clusters likely contain interesting information, we did not characterize them further, as we decided to focus on the vasculature.

Beyond the mesophyll, epidermal cells were identified in C13, based on enrichment of the epidermis-specific transcription factor AtML1 (Takada et al., 2013) and wax-related genes (Figure 3). Epidermal guard cells were found in C16, indicated by the specific enrichment of FAMA and guard cell-specific MAP kinases (Figure 3; Supplemental Figure 3A).

Enrichment of transcripts of the purine transporter PUP1 and the homolog of carrot chitinase, EP3, indicated that C17 comprises hydathodes cells (Supplemental Figure 3A; Bürkle et al., 2003).

Several clusters showed a vasculature identity, based on the enrichment of known marker genes, indicating that the enrichment for these cell types was successful. The vasculature is composed of phloem and xylem, often separated by a meristematic layer, the procambium, which initiates and maintains vascular organization. We assigned clusters to the seven known cell types present in leaves (Figure 2).
Additionally, we identified cell types with identities that had properties of two cell types, e.g., xylem and procambium. Cluster 4 was found to contain bundle sheath (BS) and xylem cells and was subsequently subclustered into three cell populations, C4.1, C4.2, and C4.3 (Figure 2B; Supplemental Data Set 5). Subclusters C4.2 and C4.3 were enriched for the BS marker SCL23, and the sulfate transporter, SULTR2;2 (Figure 3; Supplemental Figure 3, B and C). C4.2 was enriched for genes involved in photosynthetic processes, possibly indicating a differentiation of BS cells (Supplemental Data Set 5). The existence of two BS clusters is consistent with morphological descriptions (Haritatos et al., 2000). C4.1, C10, C15, and C18, with a total of 478 cells, were assigned as vascular cell types (Figures 2 and 3; Supplemental Figure 3A, Supplemental Data Sets 2 and 3).

Degree of kinship relations among BS, XP, PC and PP
Clusters composed of BS, XP, and PC, and PP cells show a J-shaped relatedness in our UMAP plot (Figure 2). BS is related to two types of xylem cells (XP1 and XP2), that abut the PC. PC subclusters into two subsets closely related to XP1 (PCXP) and PP1 (PPPP), respectively. The PCPP is adjacent to the two PP clusters PP1 and PP2. XP3 and PP2 form
a separate group, which together with XP2, form the hook of the J shape in our UMAP plot (Figure 2B).

Xylem-related clusters
The enrichment of GLR3;6 (Nguyen et al., 2018) and ACL5 markers identified C4.1 as parenchymatic xylem cells (Supplemental Figures 3, A and D–I). Further clustering of C4.1 resulted in two subgroups, XP1 and XP2, distinguished by the relative enrichment of photosynthetic genes (Supplemental Data Set 5). An additional subgroup with xylem identity, XP3, was observed as a subgroup of cells from C18. XP3 was enriched with xylem markers as well as those for VP (Endo et al., 2008; Supplemental Figure 3, D–I; Supplemental Figure 4). Transport proteins were enriched in the xylem clusters (XP1–XP3), as one may expect from their role in supplying tracheal elements with ions and nutrients for xylem translocation. These included amino acid transporters, such as the H⁺/amino acid symporter AAP6 and the UmamiT amino acid exporter family member UmamiT22 (Okumoto et al., 2002; Pilot et al., 2004; Dinkeloo et al., 2018). The transcript of amino acid export protein GDU4 was also enriched in the xylem clusters. Multiple SULTR low-affinity sulfate transporters, which play pivotal roles in sulfate transport in XP cells, were detected in the xylem cells as well as the boron transporter NIP6.1, the glucosinolate importer GTR2/NPF2.11 (Nour-Eldin et al., 2012; Madsen et al., 2014), the heavy metal transporter HMA2, and the plastidic bile acid transporter BATS/BASS5 (Supplemental Data Set 6).

Phloem- and xylem-related subpopulations of the PC subclusters
Arabidopsis procambium constitutes a bifacial stem cell population producing xylem on the adaxial pole and phloem on the abaxial pole (Elo et al., 2009; Sanchez et al., 2012). On our UMAP plot, the XP clusters are in proximity to Cluster 10. Cluster 10 could be further subdivided into three subclusters: the PP1 (C10.2) and two distinct PC populations, C10.1.1 and C10.1.2 (Figures 2 and 4A; Supplemental Data Sets 7 and 8). While cells in PCPP correspond to PC cells involved in the maintenance of meristematic identity and the differentiation of phloem cells, cells of PCXP appear to be more closely related to the formation of xylem cells, and is located closer to the XP clusters (Figure 4; Supplemental Data Sets 7 and 8).

Cells in the PCXP cluster were enriched with transcripts for homeodomain leucine zipper (HD-Zip) transcription factor, HOMEOBOX GENE 8 (HB-8; Baima et al., 2001).
transcripts, known to trigger xylem differentiation (Figure 4B). Adaxial HD-ZIP transcription factors, such as *REVOLUTA/INTERFASCICULAR FIBERLESS1* (*REV/IFL1*), *PHABULOSA* (*PHB*), and *CORONA* (*CNA/AtHB15*), were also enriched in this cluster (Figure 4, C and D; Supplemental Figure S5). Conversely, the PC PP cluster closest to the PC XP was enriched with factors involved in the maintenance of protophloem identity and pluripotency in roots and stem, such as *COTYLEDON VASCULAR PATTERN 2* (*CVP2*), *CVP2 LIKE 1* (*CVL1*; Rodriguez-Villalon et al., 2015), *BARELY ANY MERISTEM 3* (*BAM3*), and *CLAVATA3/EMBRYO SURROUNDING REGION-RELATED 45* (*CLE45*; Rodriguez-Villalon et al., 2014; Gujas et al., 2020; Figure 4, E–G; Supplemental Figure S5). *BRX* and *OCTOPUS* (Truernit et al., 2012), involved in protophloem differentiation at different ends of the developing protophloem cells, were also enriched in the nonoverlapping cells in PC PP. *ALTERED PHLOEM DEVELOPMENT* (*APL*), a major phloem marker, which inhibits xylem differentiation, was also detected in this region (Bonke et al., 2003; Figure 4E). Transcripts related to phloem differentiation, including *CLE41*, *DOF5.1*, and *LBD3* were enriched in the lower region of PC PP (Figure 4, H–J; Supplemental Figure S5). Strikingly, the arrangement in the UMAP plot, of the PC PP cluster facing the PP cluster (C10.2) and the PC XP facing the XP clusters, is analogous to the morphological positioning. However, as the markers used for inferring the clusters are based not only on leaf markers, but also root and stems due to the lack of leaf procambium cell markers, further verification will be required to provide a comprehensive understanding of procambium
Figure 4 Identification of the procambium cell cluster with distinct procambium cell identities. (A) Schematics representing subpopulations of Cluster 10. Genes enriched in the subpopulations are indicated. (B–D) UMAP showing enrichment of transcripts of genes related to xylem differentiation. (E–G) UMAP showing enrichment of transcripts of genes related to maintenance of protophloem pluripotency and differentiation. (H–J) UMAP showing the distribution transcripts related to phloem differentiation.
PP clusters

SWEET11 and SWEET12 efflux transporters had previously been reported to be expressed in the PP (Chen et al., 2012; Cayla et al., 2019). Sucrose released by the SWEETs into the apoplasm is taken up into the neighboring SE/CC via the sucrase/H+ symporter SUC2/SUT1 (Riesmeier et al., 1994; Gottwald et al., 2000). C10.2 (PP1) and C18.1 (PP2) were specifically enriched for SWEET11 and SWEET12 (Figures 3 and 5, A–F). Although the top marker genes were not mutually exclusive in either of the clusters (Supplemental Figure 6, A and B), genes related to callose deposition and cell wall thickening were over-represented in PP1, reflecting transfer cell identities (Supplemental Figure 6, C and D; Maeda et al., 2014). Among them, NAC056, a transcription factor known to play a role in cell wall ingrowth deposition in the PP, was enriched in PP1 in comparison to PP2 (Supplemental Figure 6C; Wu et al., 2018). On the other hand, genes involving photosynthesis were enriched in C18.1 indicating that C18.1 and C10.2 could be cells undergoing differential photosynthetic activity (Supplemental Figure 6E).

Both C10.2 and C18.1 are enriched for transcripts related to transport processes, reflecting the roles of PP as a critical cell type required for loading diverse substrates into the phloem (Supplemental Data Sets 7, 9–12). We identified an additional member of the Clade III SWEET family, SWEET13, in the cells that expressed SWEET11 and SWEET12 (Figure 5, G–I). SWEET13 transcripts show compensatory accumulation in sweet11;sweet12 double knockout mutants, indicating redundant roles with SWEET11 and SWEET12 (Chen et al., 2012).

As it is challenging to unambiguously distinguish phloem cell types by cross-sections, we used a modified protocol from a recently developed confocal imaging method (Cayla et al., 2019), which allows us to distinguish PP cells from the CC, SE, and BS, based on the position of chloroplasts, presence of cell wall ingrowths, and cell size. Using this method, we were able to validate the presence of SWEET11 and SWEET13 in the PP cells in pSWEET11:SWEET11-2A-GFP (Figure 5P) and pSWEET13:SWEET13-YFP (Figures 5Q) expressing lines.

UmamiT family members have been described as cellular exporters for amino acids (Ladwig et al., 2012). We thus speculated that UmamiTs might play analogous roles for the efflux of amino acids from PP as the SWEETs do for sucrose. Consistent with our hypothesis, we found UmamiT18/SIAR1 mRNA in PP (Figure 5, J–L). Vascular expression in leaves is supported by UmamiT18/SIAR1 transcriptional fusion lines (Ladwig et al., 2012). In addition, transcripts for six other members of the UmamiT family (UmamiT12, 17, 20, 21, 28, 30) were enriched in the two PP clusters (Figure 5, M–O; Supplemental Figure 7, A–F; Supplemental Data Sets 7 and 12). Notably, many of the PP-specific UmamiTs were coexpressed with each other as well as SWEET11 and 12 (Supplemental Figure 7G and Supplemental Data Set 13).

As many transcripts related to transport processes were specifically enriched in the PP clusters, and many of them were coregulated (Supplemental Data Set 13), this may indicate that they are subject to control by the same transcriptional networks. We therefore searched for transcription factors that might be responsible for coregulation of the functionally related transporters. We identified multiple leucine zipper genes (bZIP) either specific to the PP clusters (bZIP6, bZIP7, bZIP9), or present in PP and other clusters (TGA7, bZIP11; Supplemental Figure 8). We were able to confirm that bZIP9 promoter activity was specific for the PP cells in the leaf vasculature, using a transcriptional fusion, pbZIP9-GFP-GUS (Figure 6, A–D). Strong GUS activity was also detected in cell types known to be involved in apoplastic transport steps, such as the unloading zones of anthers and seeds, and the veins of petals, receptacles, ovules, transmitting tract, and funiculi (Supplemental Figure 59). The cell specificity of bZIP9 is consistent with a possible role in the activation of target genes involved in transport processes across different tissues. However, as previously reported, bzip9 single mutants did not display phenotypic differences compared with wild type (WT) plants (Supplemental Figure S10, A and B; Silveira et al., 2007; Veerabagu et al., 2014) and transcript level of PP markers were not significantly changed in the bzip9 mutant (Supplemental Figure S10C). As bZIP family members act as dimers or heteromers, hence are functionally connected, it may be necessary to characterize mutants in multiple PP-expressed bZIP family members to elucidate their role.

The CC cluster

CCs, which acquire carbon and nitrogen from the adjacent PP, but also maintain the functionality of the enucleate SEs, cluster separately from all other cell types. C15 was designated as CC based on the enrichment of transcripts for the sucrose/H+ symporter SUC2/SUT1, the H+-ATPase AHA3, and genes such as FTIP and APL (Figure 3; Supplemental Figure S3A). CCs were enriched for transcripts of various transporters, including the amino acid/H+ symporter AAP2 (Zhang et al., 2010), the potassium channel KAT1 (Schachtman et al., 1992), the hexose uniporters SWEET1 and SWEET4 (Chen et al., 2010), and the tonoplast peptide transporter PTR4/NPF8.4 (Weichert et al., 2012, p. 4; Supplemental Data Set 12). All these markers are also enriched in the CC translomate (Mustroph et al., 2009). A subset of cells in CC is specialized for synthesizing FT, a small mobile protein that acts as florigen (Chen et al., 2018). Although we could neither detect FT mRNA nor the presence of specialized CC cells in our data set, many genes involved in the regulation of FT and control of flowering, for instance, FTIP, CDFS, EFM, VOZ1, were enriched in CC (Liu et al., 2012; Yan et al., 2014; Henriques et al., 2017; Kumar et al., 2018; Supplemental Data Set 3).
Figure 5 Three SWEET sucrose transporters and UmamiT amino acid transporters mark the PP cluster. (A–O) UMAP and violin plots of C10 and C18 subclusters showing enrichment of SWEET11 (A–C), SWEET12 (D–F), SWEET13 (G–I), UmamiT18/SIAR1 (J–L), and UmamiT20 (M–O) transcripts in PP clusters. Subcluster 10.1 corresponds to PC, 18.2 to XP3, and 10.2 and 18.1 to PP. Inset show magnification of C10 and C18. (P and Q) Confocal microscopy images of pSWEET11:SWEET11-2A-GFP-GUS (P) and pSWEET13:SWEET13-YFP (Q) leaf showing GFP (P) or YFP (Q) fluorescence specific to PP. Magenta, chlorophyll autofluorescence. Yellow, FM4-64FX. Cyan GFP fluorescence (P) or YFP fluorescence (Q). Scale bars: 10 μm. BS, bundle sheath; CC, companion cell; PP, phloem parenchyma cells are marked. Numbers on the bottom left indicate independent transgenic lines.
scripts were even four-fold higher in the apl mutant. (A) Morphology of apl mutant (right) grown on LD conditions for 2 weeks. WT plant grown under the same condition is shown on the left side. Scale bar: 1 cm. (B) RT-qPCR analysis of CC-marker gene (SUC2) and PP-marker genes (SWEET11 and bZIP9). Segregating seeds from heterozygous parents were plated on MS for 2 weeks. The first and second leaves from plants homozygous for the APL mutation (apl) and heterozygous for the mutation or WT (APL) were collected for RNA extraction and RT-qPCR. Three independent replicates showed similar results and a representative experiment with three technical replicates are shown (mean ± se, n = 3).

Figure 7 Distinct PP and CC marker gene expression in the apl mutant. (A) Morphology of apl mutant (right) grown on LD conditions for 2 weeks. WT plant grown under the same condition is shown on the left side. Scale bar: 1 cm. (B) RT-qPCR analysis of CC-marker gene (SUC2) and PP-marker genes (SWEET11 and bZIP9). Segregating seeds from heterozygous parents were plated on MS for 2 weeks. The first and second leaves from plants homozygous for the APL mutation (apl) and heterozygous for the mutation or WT (APL) were collected for RNA extraction and RT-qPCR. Three independent replicates showed similar results and a representative experiment with three technical replicates are shown (mean ± se, n = 3).

PP and CC cells undergo differentiation through distinct pathways

The clear separation of PP and CC in the UMAP was striking and indicates the presence of distinct differentiation pathways. The MYB coiled-coil-type transcription factor APL plays a key role for the definition of phloem identity (Bonke et al., 2003). In our data set, APL was preferentially expressed in CC, but also detected in the PC-PP cluster (Figure 4E; Supplemental Figure 3A). As APL function is necessary for the development of the SE/CC complex, one may propose two hypotheses: (i) APL serves as a master regulator of all phloem cells or (ii) PP develops independently of the SE/CC differentiation. While apl mutants are characterized by drastic reduction of CC marker SUC2, apl retains expression of multiple PP markers (Figure 7, A and B); PP marker transcripts were even > four-fold higher in the apl mutant (Figure 7B; Bonke et al., 2003). Whether the increase of PP-marker transcripts in apl is due to a compensatory mechanism in which APL represses PP differentiation, as suggested for xylem differentiation (Bonke et al., 2003), remains to be elucidated.

The existence or function of PP has not been described for Arabidopsis roots. One may hypothesize that the root phloem-pole pericycle plays a role in the unloading of small molecules and thus ontogenetically or functionally be related to leaf PP (Ross-Elliott et al., 2017). Cell clusters in the roots with transcript profiles related to leaf PP1 do not appear to correspond to phloem pole pericycle identity. The root single-cell transcriptome atlas from Wendrich et al. (2020), which includes the highest proportion of vascular cells, had identified the phloem and pericycle as distinct clusters. A comparison shows that 19% of the phloem-pole pericycle markers overlap with pericycle markers in root single-cell sequencing data set, indicating that PP and pericycle are distinct (Brady et al., 2007; Wendrich et al., 2020). Furthermore, none of the recent single-cell sequencing analyses of roots provide a detailed analysis of different phloem cells (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). Therefore, we compared the markers from our leaf phloem clusters with the phloem markers identified by single-cell sequencing from roots (Wendrich et al., 2020). Surprisingly, root phloem markers had the highest overlap with C10.2 (Supplemental Figure 11A). Markers enriched in leaf C10.2 and the root phloem include genes related to callose deposition, as found for leaf PP1 (Supplemental Figure 11B). This finding may indicate the existence of a yet unidentified cell type in roots with a leaf PP identity. Whether PP cells are present in the root and whether they play analogous roles as in the leaf remains to be elucidated.

Unique and complementary metabolic landscapes of PP and CC

Amino acid transporters of the AAP and UmamiT families are relatively nonselective and transport many of the proteogenic and other amino acids (Fischer et al., 1995; Dinkeloo et al., 2018). One may therefore assume that the translocation of most amino acids is similar and that relative
a-Gluco-83A1 and markers for glucosinolate biosynthesis such as
cells, as biosynthetic enzymes could not be detected in iso-
analysis revealed that glucosinolates are not produced in S-
markers that are known to be absent in S-cells. Proteomic
cells are presently unavailable, therefore, we searched for
present in the abaxial area of PP in leaves, we questioned
herbivores and pathogens. As S-cells were suggested to be
act as functional shield of the plant vascular system from
10-fold higher sulfur levels, high levels of glucosinolate and
of the leaf (Koroleva et al., 2010). S-cells contain more than
of Arabidopsis but also described as sulfur-rich cells in the PP
Glucosinolate-rich cell types (S-cells) are found in floral stems
Cluster 19 as a candidate for putative S-cell cluster
Glucosinolate-rich cell types (S-cells) are found in floral stems of Arabidopsis but also described as sulfur-rich cells in the PP of the leaf (Koroleva et al., 2010). S-cells contain more than 10-fold higher sulfur levels, high levels of glucosinolate and act as functional shield of the plant vascular system from herbivores and pathogens. As S-cells were suggested to be present in the abaxial area of PP in leaves, we questioned whether S-cells are present in the PP clusters. Markers for S-cells are presently unavailable, therefore, we searched for markers that are known to be absent in S-cells. Proteomic analysis revealed that glucosinolates are not produced in S-cells, as biosynthetic enzymes could not be detected in isolated S-cell extracts. The promoter activities of key enzymes and markers for glucosinolate biosynthesis such as CYTOCHROME P450 83A1 (CYP83A1) and BRANCHED-CHAIN AMINOTRANSFERASE 4 (BCAT4) were not detected in S-cells (Li et al., 2010; Nintemann et al., 2018; Hunziker et al., 2019). We detected transcripts of CYP83A1, BCAT4, as well as other transcripts encoding proteins involved in the synthesis of aliphatic and indolic glucosinolates in the BS, xylem (C4) and PP/PC/XP (C10, C18) clusters (Supplemental Figure 14). The enrichment of HIG1 transcription factor which activate glucosinolate biosynthetic genes (Gigolashvili et al., 2007; Supplemental Data sets 4 and 7) also exclude the cells in the PP clusters (C10.2, C18.1) from containing S-cells and suggest the PP, procambium, xylem, and BS cells as sites for glucosinolate biosynthesis.

A unique characteristic of the S-cells is that these cells undergo programmed cell death at the early stages of differentiation. We searched for cells that could be enriched with the transcripts related to programmed cell death and identified Cluster 19, a cluster distinct from, but closely spaced to the BS cells. This cluster was enriched with transcripts related to programmed cell death, hypersensitive response, defense, and immune response (Supplemental Data set 14). This cluster also showed high activity scores in insect chewing-induced glucosinolate breakdown pathway (Supplemental Figure 12B). This result is in line with the primary role of S-cells in releasing toxic compounds through glucosinolate breakdown upon chewing induced-mechanical disruption. Although we cannot rule out that these subset cells were clustered based on the stress response from the protoplasting process, they could serve as candidates as putative S-cells.

Cell-type-specific expression of plasmodesmatal proteins
PDs are highly complex channels that interconnect plant cells, likely transporting solutes, metabolites, small RNAs,
and proteins. Different cell types have unique types of PD, including that which connects CC to enucleate SEs to supply all components necessary for function. The PDs typically are branched on the CC side and have a single pore on the SE side. We here found that transcripts of PD genes such as the PLAMODESMA-LOCATED PROTEIN (PDLP) and MULTIPLE C2 DOMAINS AND TRANSMEMBRANE REGION PROTEIN (MCTP) showed distinct expression patterns in different cell types (Figure 9, A–D; Supplemental Figure 16A). The cell-type specificity of PDLPs was correlated with their phylogenetic relationships (Figure 9E). Our results are consistent with data from GUS reporter fusions for the PDLP7 (Liu et al., 2012). In comparison, MCTP1, MCTP2, and MCTP3 were mainly enriched in the vasculature (Figure 3; Supplemental Figure 16B). In particular, MCTP1 (FTIP1) transcript was specifically enriched in CC, consistent with a role in trafficking the flowering protein FLOWERING LOCUS T (FT), from CC to SEs (Liu et al., 2012). In comparison, MCTP4, MCTP6, and MCTP5 are ubiquitously expressed in vascular and mesophyll cells (Supplemental Figure 16B). Overall, our data support the concept of PD-type-specific assembly of PDs.

Discussion

Despite the advances in scRNA-seq technologies, application to plant cells still faces challenges since the cell walls of plant cells must be removed for the release of individual cells. Different cell types are likely under different osmotic pressure in the intact plant, which might result in breakage if the osmotic conditions are not adequate. Recent studies using Arabidopsis (Liu et al., 2020), rice (Wang et al., 2020), as well as maize (Bezrutczyk et al., 2021) aerial tissues clearly demonstrate difficulty in capturing vascular cell types reflecting the need for careful strategy development. A key cell type in the vasculature, the PP, was not identified in any of the recently published scRNA-seq data from roots (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).

Here, we systematically optimized protoplast isolation protocols to enrich vascular cell types and produced a single-cell transcriptome and metabolic activity score atlas that covers essentially all known cell types in the Arabidopsis leaf. The scRNA-seq confirmed that except for trichomes, abaxial epidermis, and guard cells, which were intentionally removed, all known nucleus-containing cells of the leaf were represented in the dataset. In particular, all cells from the leaf vasculature were identified.

The vascular cells had identities clearly distinct from those of the epidermis, guard cells, and mesophyll. The BS formed a supercluster together with all vascular cells including the vascular meristem, except the CCs. A major finding within this atlas was that the kinship relation of vascular cells was highly similar to the actual morphology of the vasculature, with the exception of the CC, which formed a unique island separate from BS, xylem, procambium, and the PP. Surprisingly, although we used 6-weeks-old plants for the analysis, in which the developmental processes were expected to be completed, the arrangement of the XP–PC–PP clusters in the UMAP plot reflected a potential developmental trajectory. Future pseudotime trajectory

![Figure 9](image-url)

**Figure 9** Transcript enrichment of PDLP genes in vascular cell types. (A) Violin plot showing transcript enrichment of PDLPs. (B–D) UMAP plot showing the enrichment of PDLP6 (B), PDLP7 (C), and PDLP8 (D) transcripts in the PP clusters (B and C), guard cell cluster (C and D), and CC cluster (D). (E) Phylogenetic analysis of PDLPs in Arabidopsis. The phylogenetic tree was generated with the maximum likelihood method implemented in PhyML (Lemoine et al., 2019). Percent support values from 1,000 bootstrap samples are shown. Protein motifs predictions are based on the SMART database (http://smart.embl-heidelberg.de). DUF26 domains, transmembrane region, and signal peptide are shown in green, blue, and red, respectively.
analyses from the dataset, or comparison to data from younger leaves, will allow us to better understand the transition from meristematic cells to differentiated phloem or xylem cell types and trans-differentiation processes from PP into transfer cells. Some clusters showed gradients, indicating a transition, in which transcriptional modules (e.g. for xylem identity) decreased in favor of phloem modules. One example is the PC, in which cells closer to the xylem shared some xylem properties, while those closer to PP shared PP properties. A similar behavior was seen for the VP. It is conceivable that the well-studied dorsoventral cues, or cues from neighboring cells, are responsible for these gradients.

From their positioning inside the phloem, one may naïvely have expected the PP and CC to cluster together. CCs are unique since they have to fulfill their own tasks, e.g. import sucrose from PP, and at the same time maintain the function of the adjacent enucleate SEs, living cells that act as conduits for assimilate translocation (Bel and Knoblauch, 2000). Based on this dual role, it is possibly not surprising that they form such a distinct island. Whether PP and CC have expected the PP and CC to cluster together. CCs are unique since they have to fulfill their own tasks, e.g. import sucrose from PP, and at the same time maintain the function of the adjacent enucleate SEs, living cells that act as conduits for assimilate translocation (Bel and Knoblauch, 2000). Based on this dual role, it is possibly not surprising that they form such a distinct island. Whether PP and CC share some xylem properties, while those closer to PP shared PP properties. A similar behavior was seen for the VP. It is conceivable that the well-studied dorsoventral cues, or cues from neighboring cells, are responsible for these gradients.

A major discovery was the assignment of Clusters 10.2 and 18.1 as PP, providing insights into the PP transcriptome. Although C10.2 and 18.1 appear to represent different states, C10.2 having transfer cell identity and C18.1 being enriched in photosynthetic activity, it will be interesting to determine if these clusters represent spatially different cell types or developmental trajectories. The PP has essential roles in sucrose transfer to the SE/CC, but also likely many other functions in transport, metabolism and signaling. *SWEET13* is a gene that may act in a compensatory role, as evidenced by higher mRNA levels in *sweet11;12* mutants. Here, we detected SWEET13 in the same cells as *SWEET11* and 12 (Chen et al., 2012). Notably, in a parallel study, we found that while in main veins and rank-1 intermediate veins of maize, the orthologs in maize, named *ZmSWEET13a*, *b* and *c*, are likely also expressed in PP, in the C4-specific rank-2 intermediate veins that are mainly responsible for phloem loading, the three *ZmSWEET13s* are not in PP, but in two abaxial BS cells (Bezrutczyk et al., 2021). These data indicated to us that maize uses a different path for phloem loading of sucrose as compared to, for example, Arabidopsis or potato (Kühn et al., 1997).

In addition to the clade III SWEETs, scRNA-seq provided extensive insights into other genes enriched in PP, in particular, identifying transcripts of seven members of the UmamiT amino acid transporter family. UmamiTs are rather nonselective transporters involved at sites where cellular amino acid efflux is required. For example, *UmamiT18/SIAR1* transcripts, similar to *SWEET11* and 12, were detected in the chalazal region of developing seed (Ladwig et al., 2012; Chen et al., 2015). Knockout mutants showed reduced accumulation of amino acids in seeds, likely due to both a reduction in efflux from leaves and reduced efflux from the chalaza (Ladwig et al., 2012). Of note, the SWEETs and UmamiTs showed high coexpression indices. One may thus hypothesize that the genes share transcriptional control, and thus it will be interesting to analyze whether common transcription factors binding sites are present in the PP-specific genes. SWEETs and SUTs seem to act as a pair, one responsible for sucrose efflux from PP, the other for active import into the SE/CC. UmamiTs also seem to act in pairs, with UmamiTs in PP and AAP amino acid H+/symporters including AAP2, 4 and 5 in the CC (Fischer et al., 1998; Figure 10). These complementary functions are consistent with the clear separation of the clusters in the UMAP plots. A striking result from the comparison of PP and CC was that their metabolism was also highly differentiated. In particular, we found major differences regarding amino acid metabolic pathways. Both isotope labeling studies as well as amino acid localization by 2D-NMR had indicated that amino acids behave very differently regarding entry into metabolism along with the translocation pathway and entry into the phloem (Metzler et al., 1995; Atkins, 2000). This difference could thus likely be explained by the metabolic activities along the path, rather than their nonselective transporters.

Cell type-specific metabolic pathway analysis revealed many other interesting aspects. For example, hormone metabolism varied between cell types. Cell types responsible for hormones biosynthesis are largely unknown. The metabolic analysis provides a map that may guide identification of hormone biosynthesis and transport pathways in leaves. For instance, high activities of ABA, ethylene, JA, and GA biosynthesis pathways were detected in C18 (which includes the PP), consistent with the high number of ABA, JA, and GA transporters in the phloem (Kuromori et al., 2014; Kanno et al., 2016, p. 13; Ye et al., 2016; Nguyen et al., 2017).

As most transporters show functional redundancy and display diverse substrate specificity, this data set serves as a source to identify cell type-specific redundant family members for the generation of multiple mutants, to eliminate or verify interacting partners, and to identify yet-unknown transporters.

Besides the insights into both metabolism and into apoplastic transport pathways, the study also showed that cell types with unique types of PD express particular paralogs of PD-specific proteins, such as PDLPs or MCTPs. The combined analysis of sym- and apoplastic fluxes of ions, metabolites, and signaling molecules at the cellular level will likely enable a much better understanding of the physiology of the leaf.
In summary, scRNA-seq enabled us to identify unique and distinct features of the different vascular cell types present the leaf. Through transcriptomic and metabolic pathway analyses at the single-cell level, we identified potential roles of PP not only in the transport of sugars, but also in amino acid transport. Importantly, we also identified unexpected roles of PP in hormone biosynthesis and defense-related responses. The information in this study provides key resources to develop strategies influencing the flux of ions, metabolites, and signals.

**Materials and methods**

**Enrichment of vascular protoplasts**

Protoplasts were isolated from mature leaves of 6-weeks-old Arabidopsis Col-0 plants grown under short-day (8-h light/16-h dark) conditions at a PAR of 60 μmol m⁻² s⁻¹. For protoplast isolation, the tape-sandwich method (Wu et al., 2009) was modified and applied for removal of the abaxial epidermis, guard cells, and trichomes. The adaxial side of the fully developed leaves was stabilized by placing on the time tape, and the abaxial side was adhered to the 3M Magic tape. Two cuts were made on each side of the major vein of the peeled leaves using a razor blade (Swann-Morton, Sheffield). Seven to nine abaxial epidermis-peeled and cut leaves attached to time tape were immediately immersed in the petri dish containing 15 mL freshly prepared protoplast isolation solution (1% Cellulase Onuzuka R-10 (Duchefa, Haarlem), 0.3% macerozyme R-10, (Duchefa, Haarlem), 0.6 M Mannitol, 20 mM MES (2-(N-
Morpholino)ethanesulfonic acid hydrate) pH 5.7, 20 mM KCl, 1 mM dithiothreitol (DTT), 10 mM CaCl₂, and 0.1% bovine serum albumin, BSA). DTT and BSA were added to the enzyme solution to protect the protoplasts. Leaves were shaken at 30 rpm (IKA Rocker 3D orbital shaker), on a platform shaker for 2 h. The release of the protoplasts was monitored every 30 min by checking released cells under the microscope or by monitoring what was left on the time tape. After confirming the release of the protoplasts into the solution, 10 mL of wash buffer (0.6 M Mannitol, 20 mM MES pH 5.7, 20 mM KCl, 1 mM DTT, 10 mM CaCl₂, and 0.1% BSA) was slowly added to the petri dish containing the protoplasts. The solution was filtered into a round bottom 20 mL tube using a 70-μm pore size filter (Corning, NY, USA). The protoplast solution was centrifuged at 100 x g for 3 min in a swinging rotor. The protoplasts were washed four times with 20 mL washing buffer. After the final wash step, protoplasts were slowly resuspended in 1 mL wash buffer and gently filtered twice using a 40-μm Flowmi cell strainer (Bel-Art SP Scienceware, NJ, USA). The number of protoplasts was counted with the C-Chip Neubauer improved hemocytometer (NanoEnTek, Seoul) under a light microscope (Leica VT1000, Wetzlar). The viability of protoplasts was determined using trypan blue solution (Gibco; Thermo Fisher Scientific, MA, USA).

RT-qPCR
Total RNA was extracted using the RNeasy Kit (Qiagen, Hilden). cDNA synthesis and gDNA removal steps were performed using QuantiTect Reverse Transcription Kit (Qiagen, Hilden), qPCR was performed on Stratagene Mx3000P (Agilent Technologies, CA, USA) using the Lightcycler 480 SYBR Green I Master Mix (Roche, Penzberg). Transcript levels were quantified using the relative standard curve method. Values were normalized to the level of the internal control, UBO10. Primers used for RT-qPCR are listed in Supplemental Table 1. For RT-qPCR of *apl* mutants, segregating seeds from heterozygous parents were sown on MS media. RNA was extracted using 2-weeks-old plants grown under LD conditions.

Single cell RNA-seq library preparation and sequencing
Two biological replicates were performed with the aim of capturing approximately 7,000 leaf protoplasts for each replicate. Two batches of freshly isolated protoplasts were obtained from eight leaves from different plants (per batch) adjusted to 700–900 cells/μL and loaded into the 10× Genomics Chromium single cell microfluidics device according to the Single Cell 3’ Reagent Kit v2 protocol (10× Genomics, CA, USA). Eleven cycles were used for cDNA amplification and 12 cycles were used for final PCR amplification of the adapter-ligated libraries. The quality and size of the final library was verified on a DNA High Sensitivity Bioanalyzer Chip (Agilent Technologies, CA, USA), and libraries were quantified using the NEBNext Library Quantification Kit for Illumina (New England Biolabs, MA, USA). scRNA-seq library sequencing was performed on a NextSeq platform (Illumina Inc, CA USA), using the sequencing parameters 26,8,0,98 (c.ATG, Tübingen).

Generation of single-cell expression matrices
Reads were aligned to the *Arabidopsis thaliana* reference genome (Araport 11) using Cell Ranger 3.0.2 (10× Genomics, CA, USA) with default parameters. The output files for the two replicates were aggregated into one gene–cell expression matrix using Cell Ranger aggregate with the mapped read depth normalization option.

Dimensionality reduction, UMAP visualization, cell clustering analysis, and correlation analysis
The Seurat R package (version 3.1.0; Satija et al., 2015; Butler et al., 2018) was used for dimensionality reduction analysis. The SCTransform option was used for normalization, scaling the data and finding variable genes using default parameters (Hafemeister and Satija, 2019). During normalization, potential variation due to mitochondrial mapping percentage was removed. We did not remove rRNA before sample preparation and did not filter cells with higher rRNA transcript levels, since rRNA levels can vary among cells. Fifty principal components were selected as input for a graph-based approach to cluster cells by cell type using a resolution value of 0.8 in all clustering analyses. UMAP dimensional reduction (McInnes et al., 2018) was used for two-dimensional visualization using 10 principal components, 30 neighboring points and a minimum distance of 0.1. Subclustering was performed using the same parameters. For the correlation analysis between single-cell replicates across individual clusters, the average expression of cells within a cluster was calculated and the Pearson correlation coefficient was determined.

Identification of differentially expressed genes and cluster-specific marker genes
Genes differentially expressed across clusters or subclusters were identified by comparing average transcript levels in cells of a given cluster to that of cells in all other clusters using the Seurat package likelihood ratio test (Bimod). The following cutoffs were applied: average expression difference ≥ 0.25 natural Log and q < 0.01. Cluster-specific marker genes were selected from among the differentially expressed genes based on the criteria that marker genes must be expressed in >10% of cells within the cluster (PCT1, percentage of cells where the feature is detected in the first group), and <10% of cells across all other clusters (PCT2, percentage of cells where the feature is detected in the second group).

Bulk RNA-seq library preparation and sequencing
Total RNA was extracted from leaves (not protoplasted) and leaf protoplasts isolated using the same method for the single-cell sequencing using the RNeasy Kit (Qiagen, Hilden). On-column DNase treatment was performed to remove residual gDNA using the RNase-free DNase kit (Qiagen,
Bulk RNA-seq analysis

Paired-end reads (150 bp) were aligned to the A. thaliana reference genome (Araport11) using STAR (Dobin et al., 2012; maximum intron length of 2 kbp). Differential expression analysis was carried out in R (v3.6.1) using Bioconductor (v3.9) and DESeq2 (v1.24; absolute Log2FC ≥ 1 and q-value < 0.05). For correlation analysis of gene expression between protoplasted and nonprotoplasted bulk tissues, the Log2 (mean FPKM + 1) expression values were calculated for each gene. Pearson correlation coefficient was determined in R.

Pathway activities in different cell types

We followed the formulation introduced in Xiao et al. (2019) to calculate a PAS, which depends on the mRNA level of its constituent genes. The mRNA count of gene \( i \) in cell \( j \) was denoted as \( g_{i,k} \). We first normalized the mRNA count of a gene in a cell by the average for all genes in the cell, and denoted this normalized level as \( g'_{i,k} \). The mean transcript level \( E_{i,j} \) of gene \( i \) in cell type \( j \) is then defined as the mean \( g'_{i,k} \) over all cells of that type, \( E_{i,j} = \frac{1}{n_j} \sum_{k=1}^{n_j} g'_{i,k} \), where \( n_j \) is the number of cells classified as cell type \( j \), and \( k \) is the index for individual cells. \( E_{i,j} \) is normalized by the average transcript level of gene \( i \) across all cell types to become the relative transcript level \( r_{i,j} \) of gene \( i \) in cell type \( j \): \( r_{i,j} = E_{i,j}/\left( \frac{1}{N} \sum_{a=1}^{N} E_{i,a} \right) \), where \( N \) is the total number of cell types. The PAS of pathway \( t \) in cell type \( j \), denoted as \( p_{t,j} \), is then a weighted average of the relative transcript levels across the pathway genes: \( p_{t,j} = \sum_{i \in \text{pathway}} w_i r_{i,j} \); here, \( m_i \) is the number of genes in pathway \( t \), and the weight \( w_i \) of gene \( i \) is defined as the reciprocal of the number of pathways that include gene \( i \), to ensure a stronger influence of pathway-specific genes. As the relative mRNA levels \( r_{i,j} \) are centered around 1, the same is true for the pathway activity \( p_{t,j} \) with \( p_{t,j} < 1 \) corresponding to underrepresentation of pathway \( t \) in cell type \( j \) relative to the pathway’s activity across all cell types; conversely, \( p_{t,j} > 1 \) indicates a higher than average activity in cell type \( j \). To assess the statistical significance of a \( p_{t,j} \) value, we performed a permutation test, shuffling the cell type labels of the genes a thousand times to simulate the null distribution of \( p_{t,j} \) under the assumption of no systematic cell type-specific pathway activity; we defined an empirical \( P \)-value by comparing \( p_{t,j} \) to this null distribution. AraCyc pathway lists (Mueller et al., 2003) used for the analysis can be found in Supplemental Data Set 15.

Transgenic plants and T-DNA insertion lines

For the generation of pbZIP9:GFP-GUS lines, a 2,289-bp fragment of the \( bZIP9 \) gene promoter was amplified using \( bZIP9 \) pro attrB1 and \( bZIP9 \) pro attrB2 primers using Col-0 genomic DNA as template. The corresponding PCR fragment was purified and used for GATEWAY BP reaction into pDONR221 and cloned into the destination vector pBGWFS7.0 through LR reaction. For generating pSWEET11:SWEET12-GFP-GUS lines, a 4,784-bp fragment consisting of promoter and SWEET11 genomic region including all exons and introns was amplified using SWEET11-2A-attrB1 and SWEET11-2A-attrB2 primers. The SWEET11-2A-attrB2 reverse primer contained the 2A cleavage sequence. The genomic SWEET11 with the 2A cleavage site was cloned into pDONR221 and subcloned into the pKGWFS7.0 vector through LR reaction. To generate pSWEET13:SWEET13-YFP, a 3,941-bp fragment including SWEET13 promoter and its genomic region was amplified using SWEET13attB1 and SWEET13attB2 primers and cloned into the donor vector pDONR221-f1 by BP reaction. Subsequently, the fragment was sub-cloned into a gateway-compatible vector, pEG-TW1 (Kim et al., 2019) to generate pSWEET13:SWEET13-YFP construct by LR reaction. Transformation of plants was performed using the floral dip method (Zhang et al., 2006). Primers used for amplification are listed in Supplemental Table 1. The \( bzip9 \) mutant (salk_093416c) was obtained from the Nottingham Arabidopsis Stock Centre. Homozygous mutants were identified by PCR as described (http://signal.salk.edu/tdnaprimers2.html) using the primer pairs listed in Supplemental Table 1.

GUS histochemistry

GUS staining was performed as described with minor modifications (Martin et al., 1992). Tissues were fixed with 90% ice-cold acetone. After applying vacuum for 10 min, acetone was removed and replaced by pre-staining solution (1 mM EDTA, 5 mM potassium ferricyanide, 5 mM potassium ferrocyanide, 100 mM sodium phosphate (pH 7.0), 1% Triton-X-100). After 10 min vacuum, the pre-staining solution was replaced with staining solution containing 1 mM EDTA, 5 mM potassium ferricyanide, 5 mM potassium ferrocyanide, 100 mM sodium phosphate (pH 7.0), 1% Triton-X-100, and 2 mM X-Gluc and further vacuum infiltrated for 15 min in the dark. The tissues in staining solutions containing...
X-Gluc were incubated at 37°C in the dark for 4 h. Ethanol series were performed from 25% to 70% ethanol in 30 min steps.

Sample preparation for imaging leaf vasculature
Rosette leaves were excised by cutting the leaf petiole from 5-weeks-old plants grown under SD conditions. Doublesided tape was used to attach the petiole and the upper lamina of the leaf on the slide glass (abaxial side facing upward, as in the tape-sanwich method). The abaxial epidermis was peeled using the 3M Magic tape and immediately covered with water. The region of interest was excised with a razor blade and moved to a new slide glass. Samples were applied with FM4-64FX (Sigma-Aldrich, Missouri; 5 μg/mL) for 5 min for staining the cell membrane and imaged immediately. PP cells were identified based on several criteria only clearly detectable in longitudinal sections, i.e. plasma membrane invaginations, and cell diameter (about 8–10 μm). CCs were distinguished from the PP cells based on the regularly aligned chloroplast strings. BS cells were identified by their large size and chloroplast alignment on one side of the cell (Cayla et al., 2019).

Confocal imaging
Fluorescence images were captured using a Leica TCS SP8 confocal microscope with a 20× or 40× objective with water immersion. GFP, YFP, FM4-64FX, and chlorophyll autofluorescence signals were acquired using the following settings: GFP, excitation 488 nm (white-light laser) and emission 492–552 nm; YFP, excitation 514 nm and emission 492–552 nm; FM4-64FX, excitation 561 nm and emission 510–565 nm; chlorophyll autofluorescence, excitation 638 nm, and emission 465–738 nm.

Phylogenetic analysis
Reported amino acid sequences from A. thaliana PDLPs were aligned using the tool MAFFT with 0.123 and 1.53 gap extend penalty and gap opening penalty values, respectively. Maximum likelihood tree was constructed with full-length amino acid sequences with the software NGPhylogeny.fr. The sequence alignment file and the tree file are provided in Supplemental Files 1 and 2. Bootstrap percentages at the branch points were estimated from 1,000 bootstrap replications. The evolutionary distances were computed using the amino acid replacement LG matrix and the units are the number of amino acid substiutions per site.

Data availability
All sequencing data, both raw and processed, have been deposited in Gene Expression Omnibus GEO (www.ncbi.nlm.nih.gov/geo/) under the accession number GSE161482. The Arabidopsis leaf scRNA-seq data set will be deposited in the PscB (https://www.zmbp-resources.uni-tuebingen.de/timmermans/plant-single-cell-browser/; Ma et al., 2020) for visualization.

Supplemental data
Supplemental Figure 1. Optimization of vascular protoplast enrichment.
Supplemental Figure 2. Reliability of Arabidopsis leaf scRNA-seq data set.
Supplemental Figure 3. Enrichment of known marker genes for assigning cell types to clusters.
Supplemental Figure 4. Vascular parenchyma-specific marker genes enriched in Cluster 18 (XP3).
Supplemental Figure 5. Distinct procambium cell identities in C10 subclusters.
Supplemental Figure 6. Distinct identities of PP1 and PP2.
Supplemental Figure 7. Transcripts of UmmiT amino acid transport family members are enriched in the PP.
Supplemental Figure 8. Transcript enrichment of multiple bZIP transcription factors in the PP clusters.
Supplemental Figure 9. GUS staining of pbZIP9:GFP-GUS plants showed promoter activity in multiple tissues and organs.
Supplemental Figure 10. Phenotypic analysis of bzip9.
Supplemental Figure 11. Comparison of root and leaf phloem markers.
Supplemental Figure 12. Hormone and glucosinolate pathway activity across clusters.
Supplemental Figure 13. Metabolic pathway activity across all cell types.
Supplemental Figure 14. Enrichment of transcripts related to glucosinolate biosynthesis and transport.
Supplemental Figure 15. Enrichment of transcripts related to glucosinolate transport.
Supplemental Figure 16. Cell type-specific transcript enrichment of plasmodesmal proteins.
Supplemental Table 1. Primers used in this study.
Supplemental Movie 1. Three-dimensional UMAP plot of Arabidopsis leaf single cells.
Supplemental File 1. Arabidopsis thaliana PDLPs amino acid sequence alignment file (fasta).
Supplemental File 2. Arabidopsis thaliana PDLPs tree file (nhx).
Supplemental Data Set 1. Overview of the RNA-seq data generated in this study and list of differentially expressed genes (DEGs) in bulk RNA-seq of protoplasted and nonprotoplasted leaves.
Supplemental Data Set 2. Correlation of two scRNA-seq biological replicates and the number of cells in the clusters and subclusters.
Supplemental Data Set 3. Known marker genes used for cluster annotation.
Supplemental Data Set 4. Cluster specific marker genes and DEGs across all clusters.
Supplemental Data Set 5. Cluster 4 subcluster markers.
Supplemental Data Set 6. List of genes enriched in the xylem clusters (XP1, XP2, XP3).
Supplemental Data Set 7. Cluster 10 subcluster markers.
Supplemental Data Set 8. Cluster 10.1 subcluster markers.
Supplemental Data Set 9. Cluster 18 subcluster markers.
Supplemental Data Set 10. GO enrichment analysis of C18.1 vs. C10.2 markers.
Supplemental Data Set 11. GO enrichment analysis of Cluster 10 marker genes.
Supplemental Data Set 12. List of genes related with transport processes enriched in the CGs (C15) and phloem parenchyma (C10.2, C18.1).
Supplemental Data Set 13. List of genes coexpressed with UmamiT20 and SWEET11.
Supplemental Data Set 14. GO enrichment analysis of Cluster 19 marker genes.
Supplemental Data Set 15. Gene-to pathway table.

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