Evidence for phloem loading via the abaxial bundle sheath cells in maize leaves

Margaret Bezrutczyk 1, Nora R. Zöllner 1, Colin P. S. Kruse 2, Thomas Hartwig 1, Tobias Lautwein 3, Karl Köhrer 3, Wolf B. Frommer 1,4,* and Ji-Yun Kim 1

1 Institute for Molecular Physiology, Heinrich-Heine-University Düsseldorf, Düsseldorf 40225, Germany
2 Los Alamos National Laboratory, Los Alamos, New Mexico 87545, USA
3 Biological and Medical Research Center (BMFZ), Genomics and Transcriptomics Laboratory (GTL), Medical Faculty, Heinrich Heine University Düsseldorf, Düsseldorf 40225, Germany
4 Institute of Transformative Bio-Molecules (WPI-ITbM), Nagoya University, Chikusa, Nagoya 464-8601, Japan

*Author for correspondence: frommew@hhu.de
†Senior author.
Conceptualization, MB, J.K., W.B.F.; methodology, MB, C.P.S.K, T.H., T.L.; investigation, MB, N.Z., T.L., C.P.S.K.; resources, K.K., W.B.F.; writing, MB, W.B.F.; and supervision, J.K., W.B.F.

The author(s) responsible for distribution of materials integral to the findings presented in this article in accordance with the policy described in the Instructions for Authors (https://academic.oup.com/plcell/pages/General-Instructions) is: Wolf B. Frommer (frommew@hhu.de).

Abstract

Leaves are asymmetric, with different functions for adaxial and abaxial tissue. The bundle sheath (BS) of C₃ barley (Hordeum vulgare) is dorsoventrally differentiated into three types of cells: adaxial structural, lateral S-type, and abaxial L-type BS cells. Based on plasmodesmatal connections between S-type cells and mestome sheath (parenchymatous cell layer below bundle sheath), S-type cells likely transfer assimilates toward the phloem. Here, we used single-cell RNA sequencing to investigate BS differentiation in C₄ maize (Zea mays L.) plants. Abaxial BS (abBS) cells of rank-2 intermediate veins specifically expressed three SWEET sucrose uniporters (SWEET13a, b, and c) and UmamiT amino acid efflux transporters. SWEET13a, b, c mRNAs were also detected in the phloem parenchyma (PP). We show that maize has acquired a mechanism for phloem loading in which abBS cells provide the main route for apoplasmic sucrose transfer toward the phloem. This putative route predominates in veins responsible for phloem loading (rank-2 intermediate), whereas rank-1 intermediate and major veins export sucrose from the PP adjacent to the sieve element companion cell complex, as in Arabidopsis thaliana. We surmise that abBS identity is subject to dorsoventral patterning and has components of PP identity. These observations provide insights into the unique transport-specific properties of abBS cells and support a modification to the canonical phloem loading pathway in maize.

Introduction

Leaves are typically asymmetric; there are often differences in the relative stomatal and trichome densities and cuticle properties between the abaxial (lower) and adaxial (upper) leaf surfaces (Nelson et al., 2002; Juarez et al., 2004b).

Although maize leaves are amphistomatic (stomata on both surfaces of the leaf), asymmetry remains apparent, with higher density of stomata on the abaxial side, bulliform cells only on the adaxial side and a conjoined, collateral and closed-type vasculature having adaxial xylem and abaxial phloem. Dorsoventral patterning in maize leaf is initiated in
Background: Maize is among the most productive crop plants. Maize leaf mesophyll cells synthesize sucrose, the transport form of carbohydrates. Sucrose is then translocated from the mesophyll cells to the sieve-element companion cells for export to sink tissues in organs such as roots and developing seeds. Although three maize sucrose uniporters (SWEET13a, b, and c) were shown to be critical for leaf sucrose export, the complete source-to-sink route in maize is not fully understood. In barley, it appears that a subset of bundle sheath cells are specialized for translocating nutrients into the vein for export. Here, we used a combination of single-cell RNA sequencing and in situ hybridization to try to reconstruct the route in maize.

Questions: Can we identify key steps in the translocation route that transports sucrose from the mesophyll to the sieve element companion cell complex? Are all maize bundle sheath cells the same, or is there a subset of bundle sheath cells that is specialized in sugar translocation in maize plants?

Findings: Using single cell sequencing combined with in situ hybridization and analysis of translational report fusions, we identified abaxial bundle sheath cells as being specialized in nutrient transport. Transcripts of SWEET13a, b, and c, as well as other sugar and amino acid transporters were enriched in the abaxial bundle sheath cells. This indicates that maize has developed a new route for transferring sucrose from the large surface area of the two ventral bundle sheath cells to the phloem.

Next steps: Can we observe sucrose gradients in a maize leaf? Do these SWEET uniporters show polar localization in the plasma membrane facing the phloem? Is the pattern similar in the maize ancestor teosinte and in related species? How did this feature evolve, and could it be used to increase productivity of other crops?

IN A NUTSHELL

Maize has developed an apoplastic route for transporting sucrose from the large surface area of the two ventral bundle sheath cells to the phloem, a new route that is distinct from the typical sink route in maize. Whether this route also exists in other C4 plants such as teosinte is unknown. We used single cell sequencing combined with in situ hybridization and analysis of translational report fusions to identify abaxial bundle sheath cells that are specialized in nutrient transport. Our results indicate that maize has developed a new route for transferring sucrose from the large surface area of the two ventral bundle sheath cells to the phloem, a feature that may be useful for increasing productivity of other crops.
We compared published protocols for releasing protoplasts from maize and varied parameters such as incubation time, enzyme concentration, enzyme blend, and preincubation in L-Cys (Ortiz-Ramírez et al., 2018). Efficiency of the release of putative vascular cells was monitored using qRT-PCR with cell type-specific markers (PEPC, NADP-mDH for MS, NADP-ME for BS and SWEET13α, β, and γ; 18S rRNA, actin as ubiquitously expressed genes), under the assumption that the SWEET13 paralogs, analogous to their Arabidopsis homologs, would be specific to PP. Although none of the protocols were able to yield efficient release of putative vascular cells, we obtained the apparent release of both BSCs and vascular cells with Protocol 4, though the majority of cells released had MS characteristics (Supplemental Figure S1).

An estimated 7,000 protoplasts were pipetted together with reagents for cDNA synthesis into a 10 × chromium microfluidic chip, where protoplasts were partitioned into bar coded gel beads (GEMs), together reagents for single-cell cDNA synthesis. Single-cell cDNA libraries were sequenced using next-generation sequencing (NGS). Gel bead-specific barcodes were used to identify mRNAs present in specific cells. After filtering this dataset (dataset A) to select for healthy cells, 3,763 cells with an average of 4,874 mRNA molecules per cell were analyzed. Unsupervised clustering was performed using Seurat (Butler et al., 2018) to determine the relationship between mRNA expression profiles, and this was ultimately represented in a two-dimensional Uniform Manifold Approximation and Projection plot (UMAP, a dimension reduction technique for visualization; Figure 1B). Cell identities were assigned to the clusters based on established marker genes for different cell types (Supplemental Table S1).
A second independent scRNA-seq experiment yielded 3,242 cells. Data set A (from the above experiment) was analyzed in detail, the sparser data set B (from the second experiment), was used only for validation and comparison (Supplemental Figure S2). Analyzing data set A, we obtained six clusters, five of which formed a large supercluster that all had MS identities plus one separate cluster corresponding to BS identity (Figure 1C). The distribution of marker genes was consistent with the roles of MS and BS cells in C₄ photosynthesis (Supplemental Figure S3 and Table S1). The ratio of MS cells to BSCs was approximately 75:1, indicating a low efficiency of BSC retrieval. To our surprise, no vascular cells were recovered. The BS cluster was further divided into two subclusters, the upper and lower subclusters in the UMAP plot, which later were assigned as abaxial (abBS) and adaxial (adBS) BS cells (Figures 1, D and 2; see below). Importantly, the separation of the BS cluster into subclusters which contained either high or low (or undetectable) levels of SWEET13a, SWEET13b, and SWEET13c mRNA, respectively, was robust when tested with multiple clustering parameters (Supplemental Figure S4).

MS and BS clusters show canonical expression of C₄ photosynthesis-related genes

Unsupervised clustering resulted in five clusters of MS cells based on the presence of genes involved in photosynthesis that are known to be differentially expressed between MS cells and BSCs (Friso et al., 2010; Li et al., 2010; Denton et al., 2017; Figure 1C; Supplemental Data Set 1 and Table S1). At first sight, the presence of five clusters was surprising. Marker genes and enriched genes are those which either are differentially expressed in a single cluster compared to all other clusters, or are differentially expressed between two specific clusters (as noted), and meet statistical significance criteria described in Supplemental Data Set 1. Cluster MS1 included most MS cells and likely represents the core MS. Cluster MS1 was enriched in photosynthetic processes. Cluster MS2 was enriched for the GO terms triose phosphate transport (GO:0035436; GO:0015717), nucleic acid metabolic process, immune system process (GO:0035436; GO:0015717), nucleic acid transport (GO:0016070), and RNA metabolic process (GO:0016070) (Supplemental Data Sets 1 and 2). Clusters MS3 and MS4 contained high levels of ribosomal protein-related transcripts. Whether these ribosomal protein-enriched clusters represent biologically relevant cell populations in the leaf or are due to artifacts was not further evaluated due to our focus on BSCs. An additional subcluster, MS5, had an apparent MS identity, but was clearly separated from other MS clusters. The main determinants for this separate clustering were iron/metal-related processes. However, MS5 was not detected in data set B (Supplemental Figure S2). Because the MS cluster was not reproducible, it was not analyzed further (Supplemental text). Our clustering data are supported by the presence of transcripts for the glutamine synthetase GLN4 (corresponding to gln1–3 and GS1–3 protein) in the MS1–4 cells (Supplemental Figure S5), consistent with previously published in situ and immunolocalization data that detected glutamine synthetase specifically in the MS (Martin et al., 2006).

NADP-ME C₄ photosynthesis-related transcripts are present in both BSC subsets

In maize, photosynthetic activity is partitioned between the MS and BSC (Li et al., 2010; Chang et al., 2012; Friso et al., 2010), allowing us to differentiate between these cell populations based on their mRNA profiles. Maize leaves utilize a combination of the NADP-ME- and PCK-dependent C₄ pathways (Pick et al., 2011; Wingler et al., 1999), and MS cells and BSC must exchange metabolites via plasmodesmata, with specific enzymes highly upregulated in one cell type or the other. To identify the clusters, we selected several key marker genes that are known to be differentially expressed in either MS or bundle sheath.

Most of the genes involved in primary carbon metabolism that showed significant differential expression in the 2010 proteomics survey of MS and BS chloroplasts (Friso et al., 2010) were expressed in the expected cell types in dataset A (Supplemental Figure S3, Supplemental Data Set 1, and Supplemental Table S1). For example, transcripts of NADP-dependent malic enzyme (ME1) were almost exclusively found in BSCs, while carbonic anhydrase (CAH1) transcript levels were high in MS cells (Supplemental Figure S3, Supplemental Data Set 1, and Supplemental Table S1), in agreement with both proteomics data (Friso et al., 2010) and their corresponding contributions to C₄ photosynthesis (Schlüter and Weber, 2020). All cells in the BSC subcluster showed high levels of mRNA for photosynthesis-related genes that function in the BSCs in NADP-ME/PCK C₄ plants, including RBCS, ME1, and PCK1 (Supplemental Figure S3, Supplemental Data Set 1, and Supplemental Table S1). The Rubisco small subunit mRNAs (RBCS1, RBCS2) were equally distributed in both abaxial and adaxial BS subclusters (Figure 1D; Supplemental Data Set 1). The bundle sheath-enriched glutamine synthetase GLN3 (corresponding to gene gln1–4 and protein GS1–4) was found in both BS subclusters, consistent with in situ and immunolocalization data that showed no evidence for a specific pattern for GS1–4 in the BS (Supplemental Figure S5; Martin et al., 2006).
Figure 2  The abaxial BS cluster contains high levels of mRNAs encoding transport proteins. A, Feature plots show normalized levels of mRNAs for maize genes differentially expressed between the two clusters of bundle sheath cells from data set A plotted in UMAP space. B–F, In situ hybridizations of SWEET13a, SWEET13b, and SWEET13c to localize their mRNAs. Rank-2 intermediate veins from sections hybridized with antisense probes for SWEET13a, SWEET13b, and SWEET13c showed that the mRNA localization of these SWEET13 genes was largely limited to abaxial bundle sheath cells. B and C, SWEET13a mRNA localization was predominantly in abBScs in the majority of veins (77.5%) and to PP in a subset of veins (22.5%; \( n = 824 \)). D, No staining was visible after hybridization with the SWEET13a sense probe (negative control). E and F, SWEET13b and SWEET13c probes showed staining predominantly in abBScs. G, RBCS1 antisense probe hybridized sections showed staining in all BS cells. H, No staining was detected in sections hybridized with an RBCS1 sense probe. I, SUT1 mRNA was localized to a vascular cell which is likely a companion cell in rank-2 intermediate veins (arrow). J, No staining was detected after hybridization with a SUT1 sense probe (negative control). See Supplemental Figure S7 for intermediate rank-1 and major veins. Bars = 100 \( \mu m \).
Abaxial BS cluster is enriched for genes encoding transport proteins

The BS cell subclusters (Figure 1D) could represent either developmental trajectories along the leaf axis (Li et al., 2010), BS cells from the three different vein classes (major vein or rank-1 or rank-2 intermediate veins; Esau, 1943; Russell and Evert, 1985; Sharman, 1942; Hughes et al., 2019), different physiological states, or dorsoventral patterning. Whereas the majority of mRNAs corresponded to BS identity, only 5 genes were enriched in abBS (the lower subcluster) and 39 to abBSCs (the upper subcluster; Table 1). Surprisingly, among the genes with the largest difference in mRNA levels between the two BS subclusters were the three SWEET13a, b, and c paralogs. SWEETs are uniporters (Chen et al., 2010), and maize SWEET13a, b, and c function as sucrose transporters.

Arabidopsis SWEET11 and SWEET12 are phylogenetically related to maize SWEET13a, SWEET13b, and SWEET13c, and in both species these SWEETs are critical for phloem loading of sucrose (Bezrutczyk et al., 2018). However, we did not know either if the maize SWEET13 proteins were co-expressed in the same cells or which cell types they function in. All three SWEET13 paralogs were present in BS cells in maize, whereas their homologs in Arabidopsis, At-SWEET11, At-SWEET12, and At-SWEET13, are enriched in PP (Kim et al., 2021). This observation is consistent with the qRT-PCR results performed during the optimization of the protoplast protocol, which detected SWEET13 transcripts, but no vascular cells, recovered by scRNA-seq. The upper BS subcluster, abBS, showed a striking enrichment for transport proteins, with 9 of 39 abBS-specific genes involved in transport (Table 1). Importantly, this included not only SWEET13a, b, and c but also an STP hexose transporter (STP3), the amino acid efflux transporter UmamiT21a (Supplemental Figure S6, Supplemental Data Set 3), two members of the H+/amino acid symporter family, AAP56 and AAP45 (Deng, 2014), a member of the nitrate peptide transport family, and the H+/ATPase AHA3 (Table 1). Notably, in Arabidopsis, transcripts for the closest Arabidopsis homolog of Zm-UmamiT21a, At-UmamiT21, were enriched in PP (Figure 3) and co-expressed with At-SWEET11 and At-SWEET12 in Arabidopsis. On the basis of the presence of SWEETs, UmamiT21, and other abBS-enriched candidates, we speculate that the Arabidopsis transcription factors that are involved in PP identity have been recruited in maize to abBS to drive the unique set of genes expressed in the abBS. The transcription factor DNA binding with one finger3 is implicated in the regulation of SWEET gene expression in rice (Wu et al., 2018) and shows abBS-preferential expression in this dataset. Two other transcription factors, bZIP4 (ABA-insensitive S-like protein) and MYB25 (just below p-value cutoff), were enriched in the abBS. In contrast, abBS-enriched, bZIP and MYB25 were not BS-specific, but were only sparsely expressed in MS cells (Supplemental Figure S5). One of the more highly expressed genes in the adaxial BS cluster was cystatin3 (CC3), the function of which is not known.

Other BSC-expressed genes such as RBCS1 showed equal transcript distribution across all BS cells, excluding the possibility of an artifact, for example, a gradient of cells differing in unique molecular identifier (UMI) counts. This included UmamiT20a, which was BS-enriched but equally expressed across both BS subclusters. The lack of specificity of many genes for subsets of BSCs is consistent with published data from in situ hybridization and immunolocalization of RBCS1 and glutamine synthetase, neither of which showed dorsoventral patterning (Langdale et al., 1988a; Martin et al., 2006; Langdale et al., 1988b; Figure 2, G and H).

Because the cells partitioned were all collected from plants grown under the same conditions on the same day, we cannot exclude the possibility that transcriptional states could be influenced by particular conditions such as undetected pathogen infection or differences in growth conditions. Therefore, a second scRNA-seq experiment was generated (data set B). Data sets A and B were combined so that the transcriptional profiles of cells from both replicates could be directly compared. In both datasets, SWEET13a, SWEET13b, and SWEET13c were expressed in a subset of BS cells in both replicates along with genes encoding other transport proteins such as UmamiT21a and STP3, as well as the abaxial-localized transcription factor KANADI1. In both scRNA-seq datasets, RBCS1 and other BS-specific C4 genes were highly expressed in all BS cells (Supplemental Figure S2). A major difference was the absence of cluster M5, indicating that M5 likely represented a subset of cells in a different physiological state (Supplemental text).

To test the hypothesis that the two BS subclusters represent spatially discrete BSC populations, in situ hybridization was used to localize the SWEET13a, b, and c mRNAs (Figure 2, B–F; Supplemental Figure S7). Notably, all three SWEET13 transcripts were specifically detected in the two abaxial BSCs (abBSCs) adjacent to the phloem. Additionally, SWEET13a, b, and c transcripts localized to the PP (Figure 2C; Supplemental Figure S7) similar to those of At-SWEET11, At-SWEET12, and 13 (Kim et al., 2021; Chen et al., 2012). Using three independent sets of probes, we discovered that transcripts of all three SWEET13s were almost exclusively found in the abBS of rank-2 intermediate veins, which are a special adaptation of C4 monocots (Langdale et al., 1988a) and serve as the main sites of phloem loading (Fritz et al., 1989, 1983). SWEET13a was detected in both abBS and PP cells in about 23% of the rank-2 intermediate veins. Thus, in the veins that are the main loading sites, sucrose efflux toward the SE/CC for phloem loading must occur predominantly from the abBS into the apoplasm toward the phloem, and only to a smaller extent by direct release at the SE/CC from PP. Rank-1 intermediate veins seemed to have a more balanced distribution of SWEET13a between the abBS and PP. In major veins, SWEET13 transcripts were also present in the medial vascular parenchyma, and the main path appeared to be through release from PP.

Because protein abundance does not always correlate with mRNA levels (Walley et al., 2016), we evaluated the cell
specificity of the SWEET13a protein. Maize lines were generated that stably expressed translational GUS reporter fusions (ProSWEET13a:SWEET13a-GUS) comprising 6-kb upstream of the ATG and all exons and introns. Six transgenic lines from three independent transformation events showed consistent localization of the SWEET13a–GUS fusion protein in the abBS and PP of rank-1 and -2 intermediate veins, and in the PP of the major veins (Figure 4; Supplemental Figure S7).

In summary, both in situ hybridization and immunolocalization showed that SWEET13a, b, and c transcripts and SWEET13a protein were found not only in the PP of maize but also in a subset of BSCs, specifically the abBS. This is different from the cellular expression of their homologs in the dicot Arabidopsis, suggesting that an additional sucrose phloem loading route has evolved in C₄ monocots (Langdale et al., 1988a).

| **abBS** cluster | logFC | FDR   | Description                          |
|------------------|-------|-------|--------------------------------------|
| Zm00001d023677   | 4.990 | 2.3E-08 | SWEET13a                             |
| Zm00001d041067   | 3.959 | 3.2E-08 | SWEET13c                             |
| Zm00001d016625   | 3.675 | 3.3E-07 | Os02g0519800 protein                 |
| Zm00001d023673   | 2.156 | 2.2E-06 | SWEET13b                             |
| Zm00001d035717   | 2.140 | 3.3E-07 | UmatiT21a                            |
| Zm00001d033551   | 1.412 | 3.3E-07 | Phosphoglycerate mutase-like family protein |
| Zm00001d033980   | 1.252 | 4.5E-03 | Ustilago maydis induced12            |
| Zm00001d019062   | 1.180 | 2.9E-04 | Membrane H⁺ ATPase3                  |
| Zm00001d035243   | 1.123 | 1.4E-03 | AAP46                                |
| Zm00001d038753   | 1.114 | 7.7E-05 | Ubiquitin domain-containing protein   |
| Zm00001d017966   | 1.098 | 5.0E-04 | Dihydropyrimidine dehydrogenase (NADP( + )) chloroplastic |
| Zm00001d012231   | 1.055 | 9.2E-04 | AAP56                                |
| Zm00001d018867   | 0.992 | 1.1E-05 | Syntaxin 132                         |
| Zm00001d000299   | 0.920 | 1.1E-05 | Endosomal targeting BRO1-like domain-containing protein |
| Zm00001d002489   | 0.809 | 4.5E-03 | PLATZ transcription factor family protein |
| Zm00001d035651   | 0.806 | 5.0E-04 | DNA binding with one finger3         |
| Zm00001d027268   | 0.753 | 6.5E-04 | STP3 (sugar transport protein 3)     |
| Zm00001d005344   | 0.737 | 1.4E-03 | Histidine-containing phosphotransfer protein 2 |
| Zm00001d013296   | 0.737 | 4.5E-03 | ATP sulfurylase 1                    |
| Zm00001d030103   | 0.735 | 8.1E-03 | Probable xylanucan endotransglycosylase/hydrolase protein |
| Zm00001d052038   | 0.723 | 3.8E-03 | Putative LH DNA-binding domain superfamily protein |
| Zm00001d012559   | 0.692 | 2.1E-03 | Stomatal closure-related actin-binding protein 1 |
| Zm00001d015025   | 0.684 | 4.5E-03 | AMP binding protein                  |
| Zm00001d038768   | 0.636 | 5.0E-03 | Reticulon-like protein B4            |
| Zm00001d044768   | 0.633 | 3.8E-03 | Protein NRT1/ PRT FAMILY 5.8         |
| Zm00001d033981   | 0.633 | 4.8E-03 | ATP sulfurylase 1                    |
| Zm00001d015618   | 0.619 | 8.1E-03 | Probable cinnamyl alcohol dehydrogenase |
| Zm00001d041710   | 0.597 | 5.0E-03 | Glutathione synthetase chloroplastic |
| Zm00001d036401   | 0.596 | 5.0E-03 | Endoplasmic homolog                  |
| Zm00001d018758   | 0.584 | 8.1E-03 | Succinate dehydrogenase1             |
| Zm00001d019670   | 0.578 | 8.5E-03 | Kinesin-like protein KIN-4A          |
| Zm00001d022042   | 0.574 | 9.4E-04 | Eukaryotic translation initiation factor 5A |
| Zm00001d049597   | 0.567 | 4.9E-03 | External alternative NAD(P)H-ubiquinone oxidoreductase B4 |
| Zm00001d016662   | 0.536 | 3.8E-03 |                                    |
| Zm00001d018178   | 0.535 | 4.7E-03 | bZIP4 (ABSCISIC ACID-INSITIVE 5-like protein 5) |
| Zm00001d032249   | 0.532 | 2.1E-03 | KANADI 1                             |
| Zm00001d002625   | 0.531 | 3.4E-03 | Probable methyltransferase PMT15     |
| Zm00001d021773   | 0.517 | 9.9E-03 |                                    |
| Zm00001d039270   | 0.512 | 3.5E-03 | Glutaredoxin family protein          |

The table shows the genes of interest that were differentially expressed between clusters abBS and adBS of data set A. Bold type = 9 out of 39 genes enriched in the abBS cluster that are specific to transmembrane transport. Criteria for inclusion were average log-fold change > 0.5 for all cells in the subcluster and an FDR-adjusted P < 0.01. The abBS specificity was validated for three genes, SWEET13a, b, and c. Whether genes with lower FDR-adjusted P-values also show high specificity will require experimental validation. BS, bundle sheath; FC, fold-change.
Abaxial BS transcripts are co-regulated during the sink–source transition

In Arabidopsis, many PP-expressed genes were found to also be co-regulated (Kim et al., 2021). We therefore tested whether several of the transporter genes identified in the abBS might also be co-regulated. SUT1 H+\/sucrose symporter genes are typically expressed to a low level in young net-importing leaves and are induced during the sink-to-source transition (Bürkle et al., 1998; Riesmeier et al., 1993). RNA was extracted from different segments of leaf 3 of V2 plants, in which the tip had transitioned to a sucrose source while the base was still in the sink state (Tausta et al., 2014; Figure 5A) for qRT-PCR. SWEET13a, transcripts of UmamiT21, AAP45, and SUT1 were 115-, 34-, 23-, and 10-fold higher, respectively, in the tip of leaf 3 (source) compared with the base (sink; Figure 5, A and B). SWEET13a protein levels were also higher in source regions of the leaf (Figure 5C). SWEET13a was detected neither in stem sections near the base of the plant, which contains whorls of developing leaves, nor in root tip (Figure 5A and B). SWEET13a protein levels were also higher in source regions of the leaf (Figure 5C). SWEET13a was detected neither in stem sections near the base of the plant, which contains whorls of developing leaves, nor in root tip (Figure 5A and B).
Discussion

Although single-cell RNA sequencing was successfully used to identify the transcriptomes of vascular cells in Arabidopsis (Kim et al., 2021), the suberin–lignin barrier surrounding the BS of maize leaves prevented access to vascular cell transcriptomes in maize. With our optimized protocol, BSCs were released and could be identified based on a broad range of known marker genes. BSCs separated into two subclusters. mRNA for BSC markers such as RBCS and GS1–4 were present at equal levels in both subclusters, whereas others were specifically enriched in one of the two subclusters. Because only moderately and highly expressed mRNAs are captured with droplet-based scRNA-seq protocols such as 10× chromium, we cannot exclude the possibility that transcripts that appear to be specific are also present at lower levels in the other cell types.

A major surprise was the finding that mRNAs for all three SWEET13 paralogs were present in BSCs, in clear contrast to the distribution of their homologs in Arabidopsis (Chen et al., 2012). Other groups had shown that SWEET13a, SWEET13b, and SWEET13c transcripts are abundant in mechanically isolated BS strands, which likely also include vascular cells within the strands (see Supplementary discussion in Chang et al., 2012). Therefore, the data shown here provide multiple lines of evidence that SWEET13a, SWEET13b, and SWEET13c transcripts are found in specialized BSCs. Because barley BSCs seem clearly differentiated, with S-type cells suspected to represent a preferential site of sucrose transfer into the phloem (Williams et al., 1989), we tested whether SWEET13a, b, and c mRNA would be present in adaxial and lateral BSCs.

To our surprise, in situ hybridization and the analysis of translational GUS fusions showed that all three SWEET13s were preferentially expressed in the abBSCs of rank-2 intermediate veins, which are considered the main sites of phloem loading in maize (Fritz et al., 1983, 1989). In maize, these two abaxial BSCs are smaller compared with the medial BSCs (Bosabalidis et al., 1994). In rank-2 intermediate veins of maize, it appears that the abBSCs may have recruited sucrose-transporting SWEETs in order to export sucrose toward the abaxially localized phloem (Figure 6). Based on the amplification of SWEET13 genes just prior to the evolution of C₄ photosynthesis in both Andropogonae and Paniceae lineages, Emms et al. (2016) hypothesized that SWEET13 transporters may export sucrose directly from the BS to the apoplasm in C₄ plants. This conclusion was based on expression RNAseq data from BS strands (Chang et al., 2012) and laser capture dissection (Li et al., 2010), techniques that both cause contamination by adjacent tissues such as vascular parenchyma (see Supplementary materials and methods in Chang et al., 2012).

The presence of SWEET13s in the abBS possibly identifies a novel route for phloem loading in which BSCs likely export photosynthetically derived sucrose to the apoplasm of the phloem on the abaxial side of the leaf. Rank-2 veins are thought to be an emergent phenomenon of C₄ grasses (Sedelnikova et al., 2018). Rank-2 veins increase the ratio of BS to MS cells, the vein density, and the capacity for nutrient transport, and they appear to be the main path for sucrose phloem loading. It is thus conceivable that this phloem loading route coevolved with the evolution of the rank-2 intermediate veins. A minority of rank-2 intermediate veins showed transcripts of SWEET13a in vascular cells, presumably PP, in addition to abBSCs, suggesting that, in a subset of rank-2 intermediate veins, sucrose may be exported from PP as well.

Given the findings of Williams et al. (1989), which indicate that barley uses adaxial and medial BSCs for phloem loading, our results suggest that the two species use distinct sets of
BSCs for transferring sucrose from the BS to the phloem. This may occur in other C₃ and C₄ species, and it will be interesting to explore whether SWEETs are also present in medial BSCs of barley. abBS and adBS transcript profiles are highly similar, possibly explaining why this differentiation of BSCs had previously not been identified.

In rank-1 intermediate and major veins, SWEET13a, b, and c transcripts were detected in cells in the vasculature that most likely correspond to PP, thus similar to the canonical route in Arabidopsis (Chen et al., 2012; Cayla et al., 2019). Phylogenetic and functional analyses had shown that the PP transporters SWEET11 and SWEET12 from Arabidopsis are phylogenetically related to the three maize SWEET13 isoforms and fulfill overall a related function, that is, cellular sucrose efflux as a key step for phloem loading (Bezrutczyk et al., 2018). Subsequent to SWEET-mediated efflux, sucrose is taken up actively by SUT1/SUC2 
H⁺/sucrose symporters in both maize and Arabidopsis (Gottwald et al., 2000; Slewinski et al., 2009). This is supported by in situ hybridization of SUT1 in maize leaves (Supplemental Figure S7). The CC localization of SUT1 is consistent with previously published results (Baker et al., 2016). We could not confirm previous data that indicated that SUT1 may also be expressed substantially in BSCs (Supplemental Figure S7). Localization of SUT1 in vascular parenchyma is consistent with a role in sucrose retrieval on the side of the phloem that faces the xylem.

Our data are compatible with the presence of two distinct sites for phloem efflux in maize leaves, one from the abBS and a more standard path from PP in a minority of rank-2 veins (Figure 6). This route may be a specific adaptation of maize leaves in the context of C₄ photosynthesis (Emms et al., 2016) to provide higher rates of sucrose flux toward the phloem. No doubt, SWEET13a, b, and c are key transporters for phloem loading, though at present we cannot assess the relative contribution of this new efflux step. This model could be tested by inhibiting SWEET activity specifically either in BSC or PP. However, because transcription factors driving expression of genes specifically in maize abBS or PP are not currently known, this hypothesis could be tested by generation of lines in which SWEET13 mRNA levels have been repressed through BSC-specific RNAi.

Notably, transcripts for other transporter genes were also enriched in the abBS. This includes UmamiT21a, a member of the SWEET family, which is known to function in sucrose transport in Arabidopsis. Further studies are needed to determine the functional relevance of these transcripts in maize phloem transport.

Figure 6 Phloem loading occurs via the abBS in maize. A, Arrangement and relative numbers of major veins, rank-1 intermediate veins, and rank-2 intermediate veins in a mature maize leaf. Note that rank-1 intermediate veins are distinguished from rank-2 by the presence of hypodermal sclerenchyma. B, A rank-2 intermediate vein surrounded by bundle sheath (blue outline) and mesophyll (gray) cells. Sucrose movement down its concentration gradient is indicated by blue arrows. C, The inset shows details of sucrose movement either from bundle sheath cells into the apoplasm via SWEET13 transporters or to PP (teal) via plasmodesmata, where sucrose is then exported to the apoplastic space by SWEETs. Sucrose in the apoplas is taken up by SUT1 into the sieve element (orange, pink) complex of the companion cells for long distance transport.
of the UmamiT amino acid transporter family. One of the key findings from the analysis of PP in Arabidopsis by scRNA-seq was that multiple members of this family were enriched in PP (Figure 3 and Table 1; Kim et al., 2021). Because they appear to play roles analogous to that of SWEETs in cellular efflux of amino acids in Arabidopsis, it appears that the abBS, besides having clear BSC identity, has acquired components or subnetworks of the PP identity. In both maize and Arabidopsis, BS and PP cell lineages are thought to diverge relatively early during vein development (Bosabalidis et al., 1994; Dengler et al., 1985; Esau, 1943), making it unlikely that abBS expression of SWEET13s is determined by cell lineage. Instead, positional cues may determine which cells adjacent to the SE/CC express the genes involved in nutrient transport. Mobile signals are known to be critical for patterning of root vasculature in Arabidopsis (De Rybel et al., 2016), and it is possible that positional cues, possibly originating in companion cells, contribute to the induction of transporter genes in adjacent cells.

The co-regulation of at least some of the abBS-enriched genes further strengthens this hypothesis. Interestingly, we also observed weak enrichment of the abaxial KANADI transcription factor (Figure 2A; Supplemental Data Set 1). KANADI plays a key role in determining abaxial identity in leaves (Candelé et al., 2008). We therefore hypothesize that one or several transcriptional regulators that are involved in the regulation of the efflux of sucrose and amino acids from PP have been brought under control of both a polarity cue and the BS identity cues in order to increase nutrient flux toward the maize phloem. It will be fascinating to identify the transcription factors that are involved in controlling the PP and BSC identities and to dissect the SWEET promoters to determine which cis-elements are involved in the acquisition of abBSC fate. Several transcriptional regulators have been identified as candidates for the induction of abBSC fate; however, the limitations of sequencing depth in 10x genomics chromium droplet-based scRNA-seq preclude a comprehensive profiling of all transcriptional regulators. Methods that provide higher sensitivity may help to address this aspect. Importantly, the abBS genes identified provide unique insight into the specialized nature of this cell type.

Comparison of this phenomenon in other grasses, those that use both C₃ and C₄ photosynthesis, as well as a careful analysis of the evolution of rank-2 intermediate veins may provide insights into how widely distributed this mechanism is and may provide hints regarding the evolution of this regulatory rearrangement. Finally, new methods will be required to gain access to the vascular cells of maize, which are not accessible through the current methods. In summary, scRNA-seq enabled the identification of cells with a unique combination of properties on the adaxial side of the bundle sheath. The identification of this new property may be relevant to bioengineering of staple crops, for example, C₄ rice.

### Material availability
Plasmids generated in this study have been deposited to Addgene under the code Plasmid #159535.

### Data and code availability
The raw data that support the findings of this study are available from the corresponding author upon request. All sequencing data have been deposited in the Gene Expression Omnibus (www.ncbi.nlm.nih.gov/geo/) under the accession number GSE157759.

### Materials and methods

#### Plant growth
Maize (Zea mays L.) B73 seeds (obtained from Maize COOP) were germinated on filter paper with deionized, distilled water in darkness at 22–25°C and transferred to soil (Einheitserde ED73, Meyer KG, Frankfurt a.M., Germany) mixed with 20% perlite upon coleoptile emergence. Plants were subsequently grown at 28–30°C in a greenhouse supplemented by sodium lamps (400 μmol m⁻² s⁻¹) for 12 h from 8:00 to 20:00. Protoplasts for scRNA-seq were generated from the last 6 cm of the distal portion of V2 leaf 2. For each pool of protoplasts tested, leaf segments from six concurrently grown plants were used. In situ hybridization was performed on sections taken from the distal portion of leaf 2 of V2 plants, the distal portion of leaf 5 of V4 plants, and the distal portion of the flag leaf from VT plants, with similar results. All images shown are from V4 leaf 5. For GUS staining, tissue segments were taken 10 cm from the tip of the third leaf below the flag leaf of T0 plants at growth stage VT (mature leaf tip). 4 cm from the tip of leaf 3 of T1 V2 plants (seeding leaf tip), 12 cm from the tip of T1 V2 plants (seedling leaf base), a stem section 1 cm above soil surface of T1 V2 plant (seedling stem), or the seminal root tip from T1 V2 plant (seedling root).

#### Genes analyzed
Gene IDs are provided as Supplemental Table S2.

#### Probe preparation for in situ hybridization
RNA was extracted from leaves of V2 B73 seedlings by a phenol–chloroform extraction method as previously described (Bezruczyk et al., 2018). cDNA synthesis was performed using QuantiTect Reverse Transcription Kit (Qiagen, Hilden, Germany). cDNA was amplified for each gene (primers are listed in Supplemental Data Set 4) using Takara PrimeSTAR GXL polymerase then subcloned into pJET1.2 using CloneJet PCR cloning kit (ThermoFisher, Meerbusch, Germany). For SWEET13a, b, and c, three unique regions in the 5’- and 3’-UTRs and in the coding region with lengths of ~100 bp were selected as probe templates. The three probes specific for one of the genes were combined for detection of the respective target gene. For SUT1, two regions in the 5’- and 3’-UTRs, unique to SUT1 but common to all six isoforms, were selected as probe templates. For RBCS1, two regions in the coding region of the gene were selected as probe templates. All cDNA sequence alignments were performed using Geneious R11 (https://www.geneious.com; Supplemental Figure S8), Probe template regions were amplified with SP6 sequences flanking the forward primers for the sense probe, and reverse primers for antisense probes.
(Supplemental Data Set 4). Digoxigenin (DIG) labeled probes were generated using the MEGAscript SP6 Transcription kit (ThermoFisher) with a 1:2 ratio of DIG-labeled UTP:UTP. Probes were precipitated after the DNase reaction by addition of 2 mg/mL glycogen, 0.1 volume 10% acetic acid, 0.1 volume NaOAc, and 2.5 volumes ethanol and were centrifuged at 4°C at 20,000g for 30 min. Pellets were washed with 70% ethanol in diethyl pyrocarbonate (DEPC)-treated water, allowed to dry, and resuspended in 25-μL RNase-free 10 mM Tris–EDTA pH 8 and 25-μL formamide.

In situ hybridization
In situ hybridization was adapted from Jackson and Simon lab protocols (Jackson, 1992; Stahl and Simon, 2010). Leaf-tip sections 1 cm in length were dissected from V2 or V5 plants and placed into 4% paraformaldehyde, vacuum infiltrated sections 1 cm in length were dissected from V2 or V5 plants with BSA block solution for 15 min each, then in Buffer C (30 mM Na2HPO4, 10 mM Tris–Cl pH 6.8, 5 mM EDTA, 50% formamide) three independent times.

Gene/probe combination were performed as a minimum of EP50 camera. In situ hybridization experiments for each Olypmus CKX53 cell culture microscope equipped with an Olympus CKX53 cell culture microscope, and veins corresponding to cell-type-specific expression in the different vein classes were counted.

Generation of Zm-SWEET13-GUS constructs
Zm-SWEET13a (Supplemental Table S2), including 5,751-bp upstream of the start codon and 684-bp downstream of the stop codon, was isolated from B73 gDNA (Supplemental Data Set 4) and inserted into pJET using the CloneJET PCR cloning kit. The final construct consists of GusPlus inserted directly upstream of the Zm-SWEET13a stop codon, preceded by a 9-Ala linker, in the Golden Gate vector pGGGb-Ag, the in silico cloning of which was performed using Geneious R11 (Supplemental Figure S8). The assembly of all fragments with the vector pGGGb-Ag was performed using the Takara InFusion HD cloning kit, and validated by Sanger sequencing.

Maize transformation
Agrobacterium tumefaciens strain EHA105 was transformed with the Zm-SWEET13a-GUS vector at the Crop Genome Engineering Facility at VIB Ghent (Vlaams Instituut voor Biotechnologie; https://www.psb.ugent.be/cgef). Transformed EHA105 cells carrying the respective plasmids were used to transform the inbred maize line B104 via Agrobacterium-mediated transformation of 600 immature embryos according to previously described methods (Coussens et al., 2012). In brief, callus formation was induced using auxins and transgenic cells selected over several weeks using phosphino- tricin selection. Plantlets were then regenerated on hormone-free medium, and presence of the transgene confirmed using TraitCheck (Romer Labs; Butzbach, Germany) and PCR analysis. Three independent transformation events were derived from different starting immature embryos, yielding six plants in total: three plants from event A (ProSWEET13a:SWEET13a-GUSplus®), two from event B (ProSWEET13a:SWEET13a-GUSplus®), and one from event C (ProSWEET13a:SWEET13a-GUSplus®).

GUS histochemistry
Tissue segments were taken 10 cm from the tip of the third leaf below the flag leaf of T0 plants at growth stage VT (maturing leaf tip), 4 cm from the tip of leaf 3 of T1 V2 plants (seedling leaf tip), 12 cm from the tip of T1 V2 plants (seedling leaf base), on the stem 1 cm above the soil surface of T1 V2 plant (seedling stem), or from the seminal root tip of T1 V2 plant (seedling root) at hour 13:00. Tissue segments were cut with a Leica RM 2155 microtome and mounted on ProbeOn Plus slides (Fisher). After deparaffinization (Malcomber and Kellogg, 2004), sections (10 μm) were cut and placed into 4% paraformaldehyde, vacuum infiltrated sections 1 cm in length were dissected from V2 or V5 plants with BSA block solution for 15 min each, then in Buffer C (30 mM Na2HPO4, 10 mM Tris–Cl pH 6.8, 5 mM EDTA, 50% formamide) three independent times.

Hybridization buffer (300 mM NaCl, 10 mM Na2HPO4, 10 mM Tris–Cl pH 6.8, 5 mM EDTA, 50% formamide, 12.5% dextran sulfate, 1.25 mg/mL tRNA) Probe incubation in slide pairs was performed at 55°C overnight. Slides were rinsed three times with 0.2× SSC pH 7 (600 mM NaCl, 60 mM sodium citrate) at 55°C for 1 h, washed with block reagent solution (Roche), washed with PBS (1.3 M NaCl, 0.07 M Na2HPO4, 0.03 M NaH2PO4 pH), and fixed with 4% paraformaldehyde for 10 min. Slides were washed with PBS and acetylated with 0.1 M triethanolamine and acetic anhydride for 10 min, then washed and dehydrated with an increasing ethanol series from 15% to 100%. Probes for each construct were mixed (e.g. all three antisense probes for SWEET13a), diluted 1:50 with formamide, denatured at 95°C for 3 min, and further diluted 1:4 with hybridization buffer (300 mM NaCl, 10 mM Na2HPO4, 10 mM Tris–Cl pH 6.8, 5 mM EDTA, 50% formamide, 12.5% dextran sulfate, 1.25 mg/mL tRNA). Probe incubation in slide pairs was performed at 55°C overnight. Slides were rinsed three times with 0.2× SSC pH 7 (600 mM NaCl, 60 mM sodium citrate) at 55°C for 1 h, washed with block reagent solution (Roche), washed with BSA blocking solution (10 mg/mL bovine serum albumen, 0.1 M Tris–Cl, 150 mM NaCl, 0.3% Triton X-100) for 45 min, and incubated with anti-DIG AP-conjugated antibody (Roche, Lot # 16646822) for 2 h at 22°C. Slides were rinsed four times with BSA block solution for 15 min each, then in Buffer C (100 mM Tris pH 9.5, 50 mM MgCl2, 100 mM NaCl) for 15 min, and incubated with 50 μL nitro blue tetrazolium chloride (NBT) and 37.5 μL 5-bromo-4-chloro-3-indolyl-phosphate (BCIP) in 5 mL buffer C for 24–48 h. Slides were washed with water, dehydrated with an increasing ethanol series as noted above, and mounted with Eukitt Quickhardening mounting medium. Images were taken with an Olympus CKX53 cell culture microscope equipped with an EP50 camera. In situ hybridization experiments for each gene/probe combination were performed as a minimum of three independent times.

Quantification of mRNA in situ hybridization staining patterns in veins
Observations of hundreds of veins led us to determine that veins could be classified by the pattern of cell types stained as intermediate rank-2 type I (abBS cells only) or type II (BS cells + vascular cells, presumably PP); intermediate rank-1 type I (abBS cells only) or type II (vascular cells only); major type I (abaxial PP only) or type II (abaxial and medial PP). Sections hybridized with antisense probes were harvested from two different leaves and observed under an Olympus CKX53 cell culture microscope, and veins corresponding to cell-type-specific expression in the different vein classes were counted.
were dissected into cold acetone and vacuum infiltrated for 2 min, then vacuum infiltrated with GUS wash buffer (20 mM EDTA, 40 mM C\textsubscript{6}N\textsubscript{6}FeK\textsubscript{3}, 40 mM C\textsubscript{6}FeK\textsubscript{4}, 20% methanol, 57.6 mM Na\textsubscript{2}HPO\textsubscript{4}, 42 mM NaH\textsubscript{2}PO\textsubscript{4}, 0.1% Triton X-100), and incubated with GUS wash buffer including 0.2% X-Gluc at 37°C for 1–48 h. GUS activity was detected by production of a dark blue chlorobromomoidigo precipitate by cleavage and subsequent oxidation from X-Gluc. Sections were dehydrated in 20%, 30%, and 50% ethanol for 30 min, fixed in formalin-alcohol-acetic acid (FAA: 50% ethanol, 3.7% formaldehyde, 5% acetic acid) for 30 min, and further dehydrated in 75% and 100% ethanol. Embedding was performed by incubating sections at 60°C in tert-butyl ethanol:para- plast dilutions at 3:1, 1:1, and 1:3 ratios. Melted Paraplast generations with similar results. The embedding procedure was repeated three times in both T0 and T1 generations with similar results.

Single-cell sequencing: protoplast preparation

Tissue was sampled from the distal portion of leaf 2 (from 1 to 7 cm, as measured from the tip) from V2 plants. This region was selected because it is thought to be nonexpanding, nondifferentiating source tissue based on results from the RNAseq-defined developmental transcriptome of the maize leaf (Li et al., 2010). For each pool of protoplasts, leaf segments from six concurrently grown plants were used. Leaf segments were harvested at hour 9:00, and tape was applied to the adaxial epidermis to stabilize the tissue, which was scored every 5 mm from the midvein to leaf edge with a razor manifold consisting of scalpel blades taped together to ensure minimum distance between scores. Taped sections were placed abaxial-side down in pretreatment solution (2 mM l-Cys, 164 mM sorbitol) and vacuum infiltrated for 10 min with 2 min of active pumping. Taped sections were incubated with gentle agitation (30 r.p.m., IKA Rocker 3D orbital shaker) for an additional 20 min in pretreatment solution, then transferred to enzyme solution (cellulase Onozuka RS 1.25%, cellulase Onozuka R-10 1.25%, pectolysin Y-23 0.4%, macerozyme R-10 0.42%, sorbitol 0.4 M, MES 20 mM, KCI 20 mM, CaCl\textsubscript{2} 10 mM, BSA 0.1%, β-mercaptoethanol 0.06%) for 3.5 h with gentle agitation (30 r.p.m. on an orbital shaker). Protoplasts were carefully pipetted onto a prewetted Corning 70-μm nylon mesh strainer placed in a tilted 50 mL Falcon tube to remove large debris, and were then slowly pipetted into a round bottom tube using a wide-bore 1 mL pipette. Protoplasts in the round-bottomed tube were centrifuged for 1.5 min at 100 g in a Hettich Rotina 38R centrifuge. The enzyme solution was gently removed and replaced with cold wash solution (sorbitol 0.4 M, MES 20 mM, KCI 20 mM, CaCl\textsubscript{2} 10 mM, BSA 0.1%). Protoplasts were carefully resuspended in wash solution and centrifuged for 30 s at 100 g, then strained through a 70-μm Scienceware Flowmi Cell Strainer to remove large debris. Washing solution steps were repeated four additional times to remove chloroplasts and small debris. Cell viability and concentration were quantified under an Olympus CKX53 cell culture microscope: 1 μL 0.4% Trypan blue was added to 9 μL of resuspended protoplasts in wash solution and pipetted into the chamber of a C-Chip Neubauer Improved Disposable Haemocytometer (NanoEntek; Seoul, South Korea); healthy (unstained) cells were counted. Protoplasts were resuspended to a concentration of 1,200 cells/μL. A variety of approaches to degrade the suberin-lignin-containing BSC walls with the addition of other enzymes failed to produce healthy cells (data not shown): Laccase (Sigma) and manganese peroxidase (Sigma), as well as enzymes provided by Novozymes (Copenhagen, Denmark), namely a cutinase, a fungal carbohydrate blend produced in Aspergillus aculeatus, a fungal β-glucanase blend produced in Humicola insolens, a pectinase preparation produced in Aspergillus, a xylanase blend, and a multienzyme complex containing carbohydrateases, including arabinase, cellulase, β-glucanase, hemicellulose, and xylanase, were each added to the existing protocol to a final concentration of 1%–2% active enzyme weight/volume. Visual inspection of protoplasts during isolation revealed that addition of these enzymes caused protoplasts to rupture.

Generation of scRNA-seq data set B

To generate data set B, tissues from six plants at the same developmental stage as in the first experiment (B73, V2, leaf 2 tip) were grown at a different time under the same growth conditions and were harvested for protoplast release. 10× Genomics chromium cell partitioning and sequencing were performed as described for the first replicate: percent.pt < 4 and percent.mt < 0.75 and nFeature_RNA > 1,800 and nFeature_RNA < 7,000; 3,242 cells passed this threshold. Normalization, scaling, and variable feature detection were performed using SCTransform (Hafemeister and Satija, 2019a). See Quantification and statistical analysis of Materials and methods section for further analysis.

Protoplast protocol optimization

Several variations of the above protocol were tested prior to the final protoplast preparation, and the presence of diverse cell types was verified by qRT-PCR using primers specific to MDH6, ME1, SWEET13a, SWEET13b, and SWEET13c (Supplemental Data Set 4). Briefly, RNA was extracted from protoplasts using the RNEasy Mini Kit, and first-strand cDNA was synthesized using a Quantitect reverse transcription kit (Qiagen, Hilden, Germany). Quantitative reverse transcription PCR (qRT-PCR) was performed using Roche LightCycler 480 SYBR Green I Master polymerase on a Stratagene Mx3000P, and relative expression of transporter genes was calculated relative to 18S rRNA and Actin using the 2− △ACT method. Modifications to the standard protoplast isolation protocol of 2 h in enzyme solution (Protocol 1, see previous section) included doubling the concentration of enzymes in solution (Protocol 2), isolating BS strands
released after 2 h followed by continued incubation of filtered BS strands in fresh enzyme solution to deplete MS cells (Protocol 3), and incubating the leaf tissue in pretreatment solution (Protocol 4, 2 mM L-Cys and 164 mM sorbitol). The protocol that yielded the highest ratio of BS:MS marker genes (MET:MDH6) included a pretreatment incubation step (164 mM sorbitol, 2 mM L-cys in MilliQ water) and 1 × enzyme solution (Supplemental Figure S1).

Cell partitioning, library preparation, and NGS
To aim for partitioning of 7,000 cells with the expectation that 3,500 cells would be sequenced, 6 µL of the protoplast suspension with an estimated 1,200 cells/µL was applied to the 10 × genomics chromium microfluidic chip (Chemistry V3.0). Thereafter the standard manufacturer’s protocol was followed. Twelve cycles were used for cDNA amplification, and the completed cDNA library was quantified using an Agilent AATI Fragment Analyzer. Sequencing was performed at Novogene (Sacramento, CA, USA) on a single lane with the Hi-Seq platform and the standard PE150 sequencing parameters.

Generation of single-cell expression matrices
Cellranger count (10 × Genomics) was used to process fastq files provided by Novogene, with read 1 trimmed to 26 bp (r1-length = 26), as the first 26 bp of a 10 × library R1 comprise the cell barcode and UMi index, and the remaining part contains the poly-A tail with no further information. A formatted reference genome was generated using Cellranger mkref using the maize B73 RefGen 4 (Jiao et al., 2017) whole genome sequence and annotation (fasta and gff3 downloaded from Ensembl B73 RefGen V4), to which reads were aligned using STAR (Dobin et al., 2013). For analysis of single-cell sequencing data, see Quantification and statistical analysis section.

Phylogeny of UmamiT transporters
BLAST results from the Arabidopsis seed sequence At-UmamiT12 to maize (AGP v4 of the MaizeSequence database (Jiao et al., 2017) and barley (IBSC v2 from the International Barley Genome Sequencing Consortium; Mascher et al., 2017) were combined with BioMart (Smedley et al., 2009) results and filtered for the WAT1-related protein domain (panther ID PTHR31218). Genes passing this filter were selected as UmamiT family candidate genes. Two trees were generated: one using an alignment of all known splice variants, and one with only the representative transcript, with similar results. Alignments were performed in MEGA7 (Kumar et al., 2016) using MUSCLE with the following parameters: gap open penalty −2.9, gap extend penalty 0, hydrophobicity multiplier 1.2, max iterations 8, clustering method UPGMA for iteration 1, 2; UPGMB for all subsequent iterations, and lambda 24. The maximum likelihood tree was created from these alignments using IQTREE webserver (Trifinopoulos et al., 2016) using the BLOSUM62 substitution model and 1,000 bootstraps (Supplemental Data Set 3).

Quantification and statistical analysis
Sample selection for scRNA-seq, qRT-PCR, and RNAseq
Plants chosen for protoplast release, qPCR, and RNAseq were randomly selected from among a larger number of individuals that had been grown concurrently and were at the same growth stage. True biological replicates (i.e. independently grown plants) were used as replicates for statistical analyses. The number of plants per sample and number of replicates is given in the respective figure legends or in specific sections of Methods. To ensure reproducibility, the plants used in successive experiments were grown in the same greenhouse under controlled conditions. Samples for a repeat of a given experiment were taken at the same developmental stage at the same time of day.

Dimensionality reduction and cell clustering
The Seurat R package (v3.1; Butler et al., 2018) was used for dimensionality reduction analysis and dataset filtering. To remove cells with low mRNA count (nFeature_RNA) and doublets, as well as damaged cells with high chloroplast (pt) or mitochondria (mt) genome-derived transcripts, cells were filtered (percent.pt < 4 and percent.mt < 0.75 and nFeature_RNA > 1,800 and nFeature_RNA < 7,000). Normalization, scaling, and variable feature detection were performed using SCTransform (Hafemeister and Satija, 2019a). Cells were clustered using FindNeighbors to create a K-nearest neighbors graph using the first 50 principle components. FindClusters was used to iteratively group cells using a resolution of 0.2 or 23. These clusters were used as input for nonlinear dimensional reduction using UMAP (McInnes et al., 2018). This allowed us to visualize the relatedness of cells in the single-cell dataset based on their transcriptomic profiles, as represented by their positions in 2D space.

Differential gene expression analysis across clusters
Genes differentially expressed across clusters or subclusters were identified by comparing the average normalized mRNA counts in cells of a given cluster to that of cells in all other clusters using the Seurat function FindMarkers. Genes with a false discovery rate (FDR) corrected $P < 0.05$ and an average log-fold change (logFC) $> 0.5$ were considered marker genes.

Identification of cluster identities
Photosynthesis-related genes known to be differentially expressed between MSCs and BSCs in NADP-ME/PCK $C_4$ plants were used as markers to define MSC and BSC clusters (Denton et al., 2017; Li et al., 2010; Schlueter and Weber, 2020; Friso et al., 2010; Supplemental Table S1). The cluster identified as BSCs was subdivided into two subclusters when FindClusters was applied with a resolution of 23, and differential gene expression analysis was performed on these two subclusters with FindMarkers (for subclusters: logFC $> 0.5$, FDR $\leq 0.01$; Supplemental Data Set 1)
Integration of scRNA-seq data sets A and B
The two scRNA-seq replicates were combined using the integration function in Seurat. First, the same parameters were used for filtering for both datasets: to remove cells with low mRNA count (nFeature_RNA) and doublets, as well as damaged cells with high chloroplast (pt) or mt genome-derived transcripts, cells were filtered (percent.pt < 4 and percent mt < 0.75 and nFeature_RNA > 1,800 and nFeature_RNA < 7,000). Each dataset was separately scaled and normalized using SCTransform (Butler et al., 2018; Hafemeister and Satija, 2019b). Integration was performed using nfeatures = 1,000, and all features (genes) were used to find anchors. UMAP plots and feature plots were generated as previously described. After cell selection, the integrated dataset consisted of 7,005 cells (3,763 from data set A and 3,242 from data set B), with four MS clusters and one BS cluster at low resolution (FindClusters resolution = 0.2) and two BS clusters at high resolution (FindClusters resolution = 26). The integrated dataset was used to produce Supplemental Figure S2 only. All other analyses were performed using data set A.

qRT-PCR of transporter genes in seedling leaves
Leaf segments were harvested from the distal and proximal end (tip and base) of leaf 3 of early V2 plants at hour 13:00. Tissue was ground in liquid nitrogen and RNA was extracted as previously described (Bezrutczyk et al., 2018). First-strand cDNA was synthesized using Quantitect reverse transcription kit (Qiagen). qRT-PCR to determine relative mRNA levels was performed using a Stratagene Mx3000P with primers for 18S, Actin, SWEET13a, 13b, 13c, SUT1, UmamiT21a, and STP3 (Supplemental Data Set 4). Relative expression of transporter genes was calculated relative to 18S and Actin using the 2^-ΔΔCT method for quantification, with similar results. Values shown in Figure 5B are the average of three technical (qRT-PCR) replicates of three pools of two plants each; error bars represent SEM. Students two-tailed paired t test values are shown. Two independent repeats confirm the data.

Gene ontology term analysis for MS clusters
Marker genes for each of the five MS clusters (LogFC > 0.5; FDR-adjusted p-value < 0.05) were used as input for Gene Ontology (GO; Ashburner et al., 2000; The Gene Ontology Consortium, 2018) analysis via the online portal GO Gene Ontology database (doi: 10.5281/zenodo.3727280; released 2020-03-23). GO terms with false discovery rate (FDR)-corrected P < 0.05 can be found in Supplemental Data Set 2.

Protoplast and bulk leaf tissue RNAseq
Protoplasts were generated according to the previously described method. Whole leaf tissue from sibling plants was ground in liquid nitrogen at the time leaf tissue was harvested for protoplast isolation. RNA from two pools of protoplasts made from four leaves each (P1 and P2), and two pools of four whole leaf segments each (L1 and L2) was extracted using the RNEasy Mini Kit (Qiagen), and four cDNA libraries were generated using the NEBNext Ultra DNA Library Prep Kit for Illumina (New England Biolabs, MA, USA) with modifications to select for 250–500 bp fragments. Sequencing of the four libraries was performed at Novogene (Sacramento, CA, USA) on a single lane with the Hi-Seq platform and the standard PE150 sequencing parameters. Reads were analyzed using a custom implementation of the Wtyeq RNAsseq analysis pipeline (https://github.com/astauff/WySeq). Briefly, reads were trimmed using TrimGalore (v 0.6.5) and aligned to the AGPv4 B73 reference genome using STAR (v 2.5.1b). Counts were generated using Subread featureCounts (v 2.0.1), and differential expression was calculated using the R-packages EdgeR (v 3.30.3) and limma (v 3.44.3) using trimmed mean M-value normalization factors. Reads corresponding to genes enriched in either BS or MS cells were normalized separately to compensate for the expected difference in cell populations represented in protoplasts and whole leaf tissue. Genes were filtered to remove those with a coefficient of variation > 75th percentile within replicate groups prior to correlation analysis. Pearson correlation (Supplemental Figure S9) and differentially expressed genes enriched in BS and MS cells (logFC > 1 or < -1) are presented (Supplemental Data Set 5). None of the genes in the abBS subclusters were induced by protoplast isolation. Rather, several showed reduced mRNA levels in the protoplast sample.

Accession numbers
Accession numbers for maize genes are provided in Table 1; accession numbers Arabidopsis are provided in Supplemental Table 2. Additional genes used in this study include: actin—Zm00001d010159 and 18s rRNA—ENSRNA049479027.

Supplemental data

Supplemental Figure S1. qRT-PCR of putative BSC and vascular-expressed genes as an indication of protoplast cell type diversity prior to sequencing.

Supplemental Figure S2. UMAP and feature plots of integrated datasets.

Supplemental Figure S3. Schematic of C4 photosynthesis-related genes and relative expression in BS and MS clusters.

Supplemental Figure S4. BS subclusters are robust to different clustering parameters.

Supplemental Figure S5. UMAP plots of glutamine synthetase, transport-related proteins, and transcription factors in bundle sheath cells.

Supplemental Figure S6. Neighbor joining tree of family of UmamiT amino acid transporters in Arabidopsis, maize, and barley.

Supplemental Figure S7. SWEET and SUT mRNA localization and SWEET13a protein localization in Rank-2 intermediate and major veins.

Supplemental Figure S8. Probe design for in situ hybridization and ProSWEET13a:SWEET13a-GUS construct schematic.
Supplemental Figure S9. Correlation of mRNA counts between protoplasted cells and whole leaf.

Supplemental Table S1. mRNA enrichment of C_{4} photosynthesis-related genes in MS and BS clusters.

Supplemental Table S2. Genes referenced in this study.

Supplemental Text.

Supplemental References.

Supplemental Data Set 1. Cluster-specific marker genes and degs across all clusters.

Supplemental Data Set 2. GO term analysis for MS clusters.

Supplemental Data Set 3. Raw data for phylogenetic trees.

Supplemental Data Set 4. Primers used in this study.

Supplemental Data Set 5. Protoplast-enriched and protoplast-depleted mRNAs

Acknowledgments

We would like to thank Reinout Laureyns (VIB-Ugent) for advice regarding in situ hybridization, Kajetan Linkert (HHU) regarding GUS histochemistry, Max Blank and Tatjana Buchmann (WF lab) for cDNA library preparation, and Thomas Kleist (HHU) for advice on phylogenetic analyses.

Funding

This research was supported by the National Science Foundation (SECRETome Project: Systematic Evaluation of CelluR ExporT from plant cells, IOS-1546879) and Deutsche Forschungsgemeinschaft (DFG, German Research Foundation) for abaxial/adaxial patterning in maize leaves. Plant Cell 20: 2073

Candela H, Johnston R, Gerhold A, Foster T, Make S (2008) The milkweed pod1 gene encodes a KANADI protein that is required for abaxial/adaxial patterning in maize leaves. Plant Cell 20: 2073

Cayla T, Le Hir R, Dinant S (2019) Live-cell imaging of fluorescently tagged phloem proteins with confocal microscopy. Methods Mol Biol (Clifton NJ) 2014: 95–108

Chang Y-M, Liu W-Y, Shih AC-C, Shen M-N, Lu C-H, Lu M-YJ, Yang H-W, Wang T-Y, Chen SC-C, Chen SM, et al. (2012) Characterizing regulatory and functional differentiation between maize mesophyll and bundle sheath cells by transcriptomic analysis. Plant Physiol 165: 165

Chen LQ, Hou BH, Lalonde S, Sosso D, Osorio S, Ferme AR, Frommer WB (2012) Sucrose efflux mediated by SWEET proteins as a key step for phloem transport. Science 335: 207–11

Cossens G, Aesaert S, Verelst W, Demeulenaere M, De Buck S, Njuguna I, Inzé D, Van Lijssebets M (2012) Brachypodium distachyon promoters as efficient building blocks for transgenic research in maize. J Exp Bot 63: 4263–4273

De Rybel B, Mähönen AP, Helaerhu Y, Weijers D (2016) Plant vascular development: from early specification to differentiation. Nat Rev Mol Cell Biol 17: 33–43

Deng L (2014) A genome-wide analysis of the AAAP gene family in maize. J Proteomics Bioinform 7: 1–11

Dengler NG, Dengler RE, Hattersley PW (1985) Differing ontogenetic origins of PCR (“Kranz”) sheaths in leaf blades of C_{4} grasses (Poaceae). Am J Bot 72: 284–302

Denton AK, Maß J, Kühaglu C, Lercher MJ, Bräutigam A, Weber AP (2017) Freeze-quenched maize mesophyll and bundle sheath separation uncovers bias in previous tissue-specific RNA-Seq data. J Exp Bot 68: 147–160

Dobin A, Davis CA, Schlesinger F, Drenkow J, Zaleski C, Jha S, Batut P, Chaisson M, Gingeras TR (2013) STAR: ultrafast universal RNA-Seq aligner. Bioinformatics 29: 15–21

Emms DM, Cowshoff S, Hibberd JM, Kelly S (2016) Independent and parallel evolution of new genes by gene duplication in two origins of C_{4} photosynthesis provides new insight into the mechanism of phloem loading in C_{4} species. Mol Biol Evol 33: 1796–1806

Esau K (1943) Ontogeny of the vascular bundle in Zea mays. Hilgardia 15: 325–368

Friso G, Majeran W, Huang M, Sun Q, van Wijk KJ (2010) Reconstruction of metabolic pathways, protein expression, and homoeostatic machineries across maize bundle sheath and mesophyll chloroplasts: large-scale quantitative proteomics using the first maize genome assembly. Plant Physiol 152: 1219

Fritz E, Evert RF, Heyser W (1983) Microautoradiographic studies of phloem loading and transport in the leaf of Zea mays. Planta 159: 193–206

Fritz E, Evert RF, Nasse H (1989) Loading and transport of assimilates in different maize leaf bundles. Planta 178: 1–9

The Gene Ontology Consortium (2018) The Gene Ontology Resource: 20 years and still GOing strong. Nucleic Acids Res 46: D330–D338

Gottwald JR, Krysan PJ, Young JC, Evert RF, Sussman MR (2000) Genetic evidence for the in planta role of phloem-specific plasma membrane sucrose transporters. Proc Natl Acad Sci 97: 13979

Hafemeister C, Satija R (2019) Normalization and variance stabilization of single-cell RNA-seq data using regularized negative binomial regression. Genome Biol 20: 296

Henderson DC, Zhang X, Brooks III L, Scanlon MJ (2006) RAGGED SEEDLING2 is required for expression of KANADI2 and REVOLUTA homologues in the maize shoot apex. Genesis 44: 372–382

Heyser W, Heyser R, Eschrich W, Fritz E (1977) The influence of externally supplied sucrose on phloem transport in the maize leaf strip. Planta 137: 145–151
Hughes TE, Sedelnikova OV, Wu H, Becraft PW, Langdale JA (2019) Redundant SCARECROW genes pattern distinct cell layers in roots and leaves of maize. Development 146: dev17543

Jackson D (1992). In situ hybridization in plants. In SJ Gurr, MJ McPherson, DJ Bowles, eds, Molecular Plant Pathology: A Practical Approach. Oxford University Press, Oxford, pp 163–174.

Jiao Y, Peluso P, Shi J, Liang T, Stitzer MC, Wang B, Campbell MS, Stein JC, Wei X, Chin CS, et al. (2017) Improved maize reference genome with single-molecule technologies. Nature 546: 524–527 10.1038/nature22971 28605751

Juarez MT, Kui JS, Thomas J, Heller BA, Timmermans MCP (2004b) Specification of cell position and bundle sheath cells from maize leaves for photosynthetic studies. Plant Physiol 133: 128–139

Langdale JA, Lane B, Freeing M, Nelson T (1989) Cell lineage analysis of maize bundle sheath and mesophyll cells. Dev Biol 133: 128–139

Langdale JA, Rothermel BA, Nelson T (1988a) Cellular pattern of photosynthetic gene expression in developing maize leaves. Genes Dev 2: 106–115

Langdale JA, Zelitch I, Miller E, Nelson T (1988b) Cell position and light influence C₄ versus C₃ patterns of photosynthetic gene expression. EMBO J 7: 3643–3651

Li P, Ponnala L, Gadotra N, Wang L, Si Y, Tausta SL, Kebrom TH, Provart N, Patel R, Myers CR, Reidel EJ, et al. (2010) The developmental dynamics of the maize leaf transcriptome. Nat Genet 42: 1060–1067

Malcomber ST, Kellogg EA (2004) Heterogeneous expression patterns and separate roles of the SEPALATA gene LEAFY HULL STERILE1 in grasses. Plant Cell 16: 1692

Martin A, Lee J, Kichey T, Gerentes D, Zivy M, Tatout C, Dubois F, Ballieu T, Valot B, Davanture M, et al. (2006) Two cytosolic glutamine synthetase isoforms of maize are specifically involved in the control of grain production. Plant Cell 18: 3252–3274 10.1105/tpc.106.042689

Mascher M, Gundlach H, Himmelbach A, Beier S, Twardziok SO, Wicker T, Radchuk V, Dockter C, Hedley PE, Russell J, et al. (2017) A chromosome conformation capture ordered sequence of the barley genome. Nature 544: 427–433 10.1038/nature22043

Mcllnnes L, Healy J, Melville J (2018) UMAP: uniform manifold approximation and projection for dimension reduction. arXiv eprint arXiv:1802.03426

Nelson JM, Lane B, Freeing M (2002) Expression of a mutant maize gene in the ventral leaf epidermis is sufficient to signal a switch of the leaf’s dorsoventral axis. Development 129: 4581

Ortiz-Ramirez C, Arevalo ED, Xu X, Jackson DP, Birnbaum KD (2018) An efficient cell sorting protocol for maize protoplasts. Curr Protoc Plant Biol 3: e20072–e20072

Pick TR, Bräutigam A, Schlüter U, Denton AK, Colmsee C, Scholz U, Fahrensticht H, Pierschka R, Rascher U, Sonnewald U, et al. (2011) Systems analysis of a maize leaf developmental gradient redefines the current C₄ model and provides candidates for regulation. Plant Cell 23: 4208

Riesmeier JW, Hirner B, Frommer WB (1993) Potato sucrose transporter expression in minor veins indicates a role in phloem loading. Plant Cell 5: 1591–1598

Riesmeier JW, Willmitzer L, Frommer WB (1994) Evidence for an essential role of the sucrose transporter in phloem loading and assimilate partitioning. EMBO J 13: 1–7

Russell SH, Evert RF (1985) Leaf vasculature in Zea mays L. Planta 164: 448–458

Schlüter U, Weber APM (2020) Regulation and evolution of C₄ photosynthesis. Annu Rev Plant Biol 71: 183–215

Sedelnikova OV, Hughes TE, Langdale JA (2018) Understanding the genetic basis of C₄ Kranz anatomy with a view to engineering C₃ crops. Annu Rev Genet 52: 249–270

Sharman BC (1942) Developmental anatomy of the shoot of Zea mays L. Ann Bot 26: 245–282

Slewinski TL, Meeley R, Braun DM (2009) Sucrose transporter1 functions in phloem loading in maize leaves. J Exp Bot 60: 881–892

Smedley D, Haider S, Ballester B, Holland R, London D, Thorisson G, Kasprzyk A (2009) BioMart—biological queries made easy. BMC Genomics 10: 22

Stahl Y, Simon R (2010) mRNA detection by whole mount in situ hybridization (WISH) or sectioned tissue analysis of maize bundle sheath and mesophyll cells. BMC Genomics 11: 53

Trifinopoulos J, Nguyen L-T, von Haeseler A, Minh BQ (2016) W-IQ-TREE: a fast online phylogenetic tool for maximum likelihood analysis. Nucleic Acids Res 44: W232–W235

Wangler A, Walker R, Chen Z, Leegood R (1999) Phosphoenolpyruvate carboxykinase is involved in the decarboxylation of aspartate in the bundle sheath of maize. Plant Physiol 120: 539–546

Wu Y, Lee S-K, Yoo Y, Wei J, Kwon S-Y, Lee S-W, Jeon J-S, An G (2018) Rice transcription factor OsDOF11 modulates sugar transport by promoting expression of sucrose transporter and SWEET genes. Mol Plant 11: 833–845