Characterizing Regulatory and Functional Differentiation between Maize Mesophyll and Bundle Sheath Cells by Transcriptomic Analysis


To study the regulatory and functional differentiation between the mesophyll (M) and bundle sheath (BS) cells of maize (Zea mays), we isolated large quantities of highly homogeneous M and BS cells from newly matured second leaves for transcriptome profiling by RNA sequencing. A total of 52,421 annotated genes with at least one read were found in the two transcriptomes. Defining a gene with more than one read per kilobase per million mapped reads as expressed, we identified 18,482 expressed genes; 14,972 were expressed in M cells, including 53 M-enriched transcription factor (TF) genes, whereas 17,269 were expressed in BS cells, including 214 BS-enriched TF genes. Interestingly, many TF gene families show a conspicuous BS preference in expression. Pathway analyses reveal differentiation between the two cell types in various functional categories, with the M cells playing more important roles in light reaction, protein synthesis and folding, tetrapyrrole synthesis, and RNA binding, while the BS cells specialize in transport, signaling, protein degradation and posttranslational modification, major carbon, hydrogen, and oxygen metabolism, cell division and organization, and development. Genes coding for several transporters involved in the shuttle of C4 metabolites and BS cell wall development have been identified, to our knowledge, for the first time. This comprehensive data set will be useful for studying M/BS differentiation in regulation and function.

C4 plants, with few exceptions, require the coordination of the mesophyll (M) and bundle sheath (BS) cells, arranged in a wreath structure called Kranz leaf anatomy (Hatch and Agostino, 1992), to confer high rates of photosynthesis. The initial carboxylation phase of the C4 pathway takes place in the M cells, while the decarboxylation phase is restricted to the BS cells. The high photosynthetic capacity of C4 plants implies a massive efflux of C4-related metabolites between M and BS cells and between the cytosol and organelles in each cell type (Weber and von Caemmerer, 2010).

Although research in the past few decades has greatly increased our understanding of the biochemical reactions and the enzymes involved in the C4 pathway of photosynthesis, little is known about the specific genes involved in the development of the Kranz leaf anatomy, the C4 biochemical pathway, and the underlying regulatory mechanisms for the high-level expression of C4-specific genes in a cell-, organ-, or development-specific manner. With the advancement in genomics, the genomic sequences of several C3 and C4 model plants have become available. These advances have allowed in-depth comparative proteomic and transcriptomic analyses of the whole leaves of typical C3 and C4 plants and their closely related C3-C4 intermediate species of Cleome and Flaveria ( Gowik et al., 2011). These comparative studies allow deduction on how many genes are required to make a C4 plant and possibly on how they may have been regulated at the genetic level. In addition, attempts have been recently made to characterize the transcriptomic profile of maize (Zea mays) leaf in a development-dependent manner ( Li et al., 2010). Also, the laser-capture microdissection (LCM) technique was used to isolate both M and BS cells from mature maize leaves for transcriptome analysis. However, LCM has a limitation in the amount of cells that can be isolated and has the potential of cross-contamination.

To gain a comprehensive understanding of the C4 photosynthesis pathway and related processes, we have modified current methods to isolate large quantities of...
RESULTS AND DISCUSSION

Isolation of Highly Pure M Protoplasts and BS Strands

In this study, highly purified mesophyll protoplasts (MP) and bundle sheath strands (BSS) were obtained in large quantities from newly matured second leaves (Supplemental Fig. S1) for transcriptome analyses. Examination by light microscopy showed that the mechanically purified BSS was attached with very little M tissue, except M cell wall and vascular and epidermal tissues. To further assess the purity of the isolated cells, the activities of two key C₄ photosynthesis marker enzymes for M and BS cells were assayed. On a chlorophyll basis, the high activity of NADP-ME, a key C₄ acid decarboxylase located in the BS cells of maize, was almost exclusively associated with the BSS isolated (Supplemental Table S1). On the other hand, the high activity of phosphoenolpyruvate carboxylase (PEPC) was mainly associated with the MP isolated by enzymatic digestion, although a low activity of PEPC was also detected in the BSS isolated, which might be due to the presence of the C₃-type PEPC in the BS cells. Immunoblot analysis using specific antibodies raised against the maize proteins also demonstrated the high homogeneity of the two cell types isolated (Supplemental Fig. S2).

Raw Read Processing and Definition of Expressed Genes

To clean the sequencing raw reads, we deleted reads that contain any N (unknown nucleotide) or have a poor quality score (Table I). Using Tophat version 1.3.3 (Trapnell et al., 2009; http://tophat.cbcb.umd.edu/), we found 78% and 79% of the cleaned reads can be mapped to the maize B73 genome (Maize Genome AGP version 2, release 5a.59). We used Cufflinks version 1.2.1 (Trapnell et al., 2010; http://cufflinks.cbcb.umd.edu/) to calculate the expression level of each gene in terms of reads per kilobase per million mapped reads (RPKM; Mortazavi et al., 2008). RPKM is normalized by the length of the exon and the total mappable reads and can be used to quantify the expression level of a gene from the mapped read count. A total of 52,421 annotated genes with at least one read were found in the two transcriptomes (Table II). (All genes with at least one read are listed in Supplemental Data S1 with their RPKM values.) The maximum RPKM were 35,421 and 31,561, and the median RPKM were 0.55 and 0.73 in the M and BS cell transcriptomes, respectively. Only approximately 2% of the genes had a very high expression level (RPKM ≥ 100) in either M or BS cells.

The quality of the transcriptomes, the potential contamination of BS cells by epidermal and vascular bundle tissues, and the potential effects of long incubation time during the isolation of M protoplasts are discussed in Supplemental Materials and Methods S1 and Supplemental Fig. S3.

Following the criterion of Li et al. (2010), we defined an “expressed gene” as a gene with RPKM > 1 in the transcriptome. In all analyses below, only expressed genes were considered. If the Filtered Gene Set (FGS) was chosen, we identified 14,972 and 17,269 expressed genes in the M and BS transcriptomes of maize, respectively (Fig. 1A). In total, we identified 18,482 genes expressed in either the M or BS transcriptome or both, which is 1,844 genes more than were identified by Li et al. (2010; Fig. 1A). If the Working Gene Set (WGS) was chosen, then we gained an additional 2,224 expressed genes (Fig. 1A).

Low Degrees of Cross-Contamination between the Two RNA Samples

It is important to know the degree of cross-contamination between the two transcriptomes, because cross-contamination can affect the estimation of the number of expressed genes in a transcriptome (one cell type) and the number of differentially expressed genes between the two transcriptomes (two cell types). To address this issue, we used the expression levels of 10 selected C₄ marker genes to estimate the degrees of cross-contamination in the two RNA samples from the isolated M cells and BS cells (Table III). The proteins encoded by these genes play key roles in C₄ photosynthesis and are assumed to be exclusively or almost exclusively expressed in only one cell type (Supplemental Data S1). Among the 10 selected genes, NADP-ME (GRMZM2G085019), DCT2 (for dicarboxylate transporter; GRMZM2G086258), PCK

Table I. Statistics of the transcriptome data

<table>
<thead>
<tr>
<th>Cell Type</th>
<th>Total Reads</th>
<th>Trimmed Reads</th>
<th>Mapped Reads (Mappable Rate) a</th>
<th>Unique Hits</th>
</tr>
</thead>
<tbody>
<tr>
<td>M</td>
<td>177,363,768</td>
<td>125,841,884</td>
<td>98,048,470 (78%)</td>
<td>89,522,392</td>
</tr>
<tr>
<td>BS</td>
<td>193,418,692</td>
<td>135,111,940</td>
<td>107,090,765 (79%)</td>
<td>100,493,851</td>
</tr>
</tbody>
</table>

aThe mappable rate is defined as the ratio of the mapped reads to the trimmed reads.
(GRMZM2G001696), and two rbcSs (GRMZM2G098520 and GRMZM2G13033) are supposed to be BS specific, while PEPC (GRMZM2G083841), CA (GRMZM2G121878), NADP-MDH (GRMZM2G129513), OMT (for 2-oxoglutarate/malate transporter; GRMZM2G383088), and PPDK-RP (GRMZM2G131286) are supposed to be M specific. Using Equations 2 and 4 as shown in “Materials and Methods,” we obtained an upper estimate of \( c_1 = 1.2\% \) for the contamination rate of M cells by BS cells and an upper estimate of \( c_2 = 4.5\% \) for the contamination rate of BS cells by M cells (Table III). These are considered the upper estimates because the assumption of exclusive expression of a key C\(_4\) gene in a cell-specific manner (e.g. PEPC in M cells) may not be absolutely true. In any case, these calculations suggest that the degrees of cross-contamination in our two RNA samples were reasonably low, which is crucial for us to accurately determine the expression profile of a given gene in a cell-specific manner.

We checked the above conclusion on the purity of our isolated M and BS cells by quantitative reverse transcription-PCR (Supplemental Table S2 and S3). For the three BS-specific genes studied (rbcS-2, rbcS-4, and NADP-ME), the respective M/BS expression ratios were 0.001, 0.049, and 0.004, which are indeed similar to the ratios obtained from the transcriptome data (0.022, 0.022, and 0.007). These data support the conclusion of very low contamination of our M cell sample by BS cells. For the three M-specific genes studied (PEPC, NADP-MDH, and OMT), the respective BS/M ratios were 0.055, 0.050, and 0.037, which are also very close to the BS/M ratios of 0.048, 0.039, and 0.045 from our transcriptome data (Supplemental Table S2), supporting an estimate of approximately 4.5\% contamination of our BS sample by M cells.

Li et al. (2010) had reported two sets of deep-sequencing transcriptomic data from the maize M and BS cells isolated by LCM. In their data, nine out of the above 10 C\(_4\) marker genes, except PPDK-RP (GRMZM2G131286), were listed as cell type-enriched genes. Applying the same equations to analyze their data, we estimated the upper contamination rates of \( c_1 = 9.2\% \) and \( c_2 = 21.6\% \) for their M and BS cell preparations. These results indicate that our isolation methods caused a much lower level of cross-contamination than the LCM method they used. Consequently, our data would give a more accurate estimation of genes differentially expressed between the M and BS cells.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>M Cells</th>
<th>BS Cells</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Detected genes (at least one read)(^a)</td>
<td>43,934</td>
<td>48,888</td>
<td>52,421</td>
</tr>
<tr>
<td>Expressed genes in the FGS</td>
<td>14,972</td>
<td>17,269</td>
<td>18,482</td>
</tr>
<tr>
<td>Expressed TF genes in the FGS</td>
<td>653</td>
<td>819</td>
<td>880</td>
</tr>
<tr>
<td>Enriched expressed genes in the FGS</td>
<td>1,545</td>
<td>3,422</td>
<td>4,967</td>
</tr>
<tr>
<td>Enriched expressed TF genes in the FGS</td>
<td>53</td>
<td>214</td>
<td>267</td>
</tr>
</tbody>
</table>

\(^a\)The total numbers of annotated genes in the WGS and the FGS of AGPv2 (release 5a.59) were 110,028 and 39,656, respectively.

Genes Differentially Expressed between the Two Cell Types

To estimate the differential expression of genes between the two cell types, we defined the degree of cell specificity of gene \( i \) as \( R_i = |m_i - b_i|/\max(m_i, b_i) \), where \( m_i \) and \( b_i \) represent the RPKM of gene \( i \) in the M and BS RNA samples, respectively. By this definition, \( R_i = 1 \) if gene \( i \) is exclusively expressed in only one cell type, and \( R_i = 0 \) if gene \( i \) is equally expressed in the two cell types. To define “differentially expressed genes,” we used the criterion of \( R_i = 0.8 \), which corresponds to a 5-fold difference in RPKM value between the two cell types. This criterion is more stringent than the criterion used by Li et al. (2010); 72\% of their differentially expressed genes had only 4-fold or smaller differences. In the differentially expressed gene set, there were 1,545 and 3,422 genes enriched in the M and BS transcriptomes, respectively (Fig. 1B). That is, 4,697 (1,545 + 3,422) genes, or approximately 25\% of all expressed genes in the FGS, were differentially expressed between the two cell types. In contrast, Li et al. (2010) identified only 3,427 (2,028 + 1,399) differentially expressed genes, or approximately 21\% of all expressed genes (16,638), even though they used a looser criterion (Fig. 1B). The lower percentage could be due to the high cross-contamination rates in the study of Li et al. (2010; see above). In our data, there were many more BS-enriched genes than M-enriched genes. Only 30\% and 16\% of M- and BS-enriched genes in this study overlapped with the Li et al. (2010) data (Fig. 1B). Thus, many more cell-specific genes have been identified from our study. For genes with \( R_i > 0.99 \), which can be taken as truly cell-specific genes, there were only 65 genes in M cells but 688 genes in BS cells (Supplemental Data S2). This result suggests that maize BS cells have assumed a more important metabolic role in C\(_4\) photosynthesis and other metabolic processes.

The major difference in photosynthetic biochemistry between C\(_3\) and C\(_4\) plants is that in C\(_4\) plants the CO\(_2\)-concentrating mechanism and the Calvin cycle are divided between M and BS cells. To achieve an effective CO\(_2\)-concentrating mechanism, genes that are involved in carbon fixation, such as PEPC (GRMZM2G083841), CA (GRMZM2G121878), NADP-MDH (GRMZM2G129513), and PPDK-RP (GRMZM2G131286), and plastid membrane
transporters such as OMT (GRMZM2G383088), are all enriched or specifically expressed only in the M cells. In our M transcriptome, the RPKM values for PEPC, CA, NADP-MDH, PPDK-RP, and OMT were 9,485, 6,692, 2,621, 476, and 988, respectively, much higher than the corresponding values 431, 281, 95, 22, and 74 in the BS transcriptome (Table III). Therefore, the cell-specificity $R_i$ values for these five genes were 0.95, 0.96, 0.96, 0.95, and 0.93, respectively. (These $R_i$ values were probably underestimated because of the potential contamination of our BS cell sample by M cells.) On the other hand, the genes involved in releasing CO$_2$ from C$_4$ acid decarboxylation, such as NADP-ME (GRMZM2G085019) and PCK (GRMZM2G001696), plastid membrane transporter, such as DCT2 (GRMZM2G086258), and the Calvin cycle, such as rbcSs (GRMZM2G098520 and GRMZM2G113033), were all enriched or specifically expressed in BS cells, each with $R_i > 0.98$ (Table III). Taken together, these results indicate that most of the known C$_4$ genes are very tightly regulated and specifically expressed in M or BS cells in mature maize leaves, allowing an effective CO$_2$-concentrating mechanism through effective carboxylation and decarboxylation in the two cell types. An exception to this rule is PPDK (GRMZM2G306345): it is preferentially expressed in M cells with a $R_i$ of only 0.78. This is consistent with the

![Figure 1](image_url)

**Figure 1.** M- and BS-expressed genes and TF genes. A, Numbers of expressed genes in M and BS cells found in the FGS and the additional numbers of expressed genes found in the WGS, excluding transposable elements, gene fragments, and pseudogenes. In Li et al. (2010), no data were available for expressed genes found in the WGS. B, Venn diagrams of M- and BS-enriched genes and TFs compared between this study and Li et al. (2010). C, Functional distribution of all cell type-enriched genes. Genes of unknown function were not included. D, Distribution of functional preferences of cell type-enriched genes. In each row, the red and blue bars represent, respectively, the percentages of BS- and M-enriched genes within the functional category. The numbers in the red and blue bars represent the numbers of M- and BS-enriched genes. E, Distribution of cell type-enriched TF families. The red and blue bars represent the percentages of BS- and M-enriched TFs within each TF family, respectively. The numbers in the red and blue bars represent the numbers of M- and BS-enriched TFs.
biochemical study by Aoyagi and Nakamoto (1985), which found substantial activity and protein of PPDK present in the maize BS cells. The conversion of phosphoenolpyruvate (PEP) to pyruvate and ATP catalyzed by the reversible PPDK is another metabolic way for cells to produce ATP.

There is strong biochemical evidence that the PCK enzyme plays a substantial role in some NADP-ME subtype C₄ plants, such as maize (Wingler et al., 1999; Leegood and Walker, 2003). Consistent with this notion, PCK was found to be expressed at an extremely high level and exclusively in BS cells (i.e. \( R_i = 1.00 \); Table III). Two other enzymes that are also required to support the operation of the PCK decarboxylation system in both cell types, namely aspartate aminotransferase (AspAT) and alanine aminotransferase (AlaAT). There are two major paralogs of AspAT found in maize with a high level of expression: one (GRMZM5G836910) is M specific (\( R_i = 0.97 \)), which encodes a chloroplast enzyme, confirming the biochemical results of Gutierrez et al. (1974), while the other (GRMZM2G094712) is BS enriched (\( R_i = 0.79 \)), which encodes a mitochondrial enzyme (Supplemental Data S3). In contrast, the two AlaAT paralogs shown to moderate expression levels, both with an M cell preference. The presence of a large amount of AspAT transcript in M cells is necessary for the operation of the PCK decarboxylation system in maize leaves, as OAA is very labile and once produced should be taken up quickly by the M chloroplast for reduction to malate by NADP-MDH or transaminated to Asp by AspAT.

### Functional Differentiation between M and BS Cells and Identification of Genes Involved in Kranz Structure Formation

For the 1,545 M-enriched and 3,422 BS-enriched genes identified, their biological functions were further elucidated using MapMan (http://mapman.gabipd.org/): 71% of the enriched genes (3,358 genes: 1,015 M enriched and 2,343 BS enriched) have been assigned to known biological processes or metabolic pathways. Except for those that have been classified as “others” (14%), the three most abundant functional groups are protein metabolism (13%), RNA metabolism (11%), and transport (6%; Fig. 1C). The other abundant functional groups are miscellaneous (5%), signaling (4%), cell (3%), stress (3%), hormone metabolism (3%), photosynthesis (2%), and development (2%; Fig. 1C). When considering the functions of these genes in M or BS cells, we identified 15 cell type-enriched functional groups using the Fisher exact test \( (P < 0.05 \) for each group; Fig. 1D). The difference in the number of cell type-enriched genes between M and BS cells within each function is quite large, except for the protein-targeting category. Overall, the number of BS-enriched genes (3,422) is much higher than that of M-enriched genes (1,545). As expected, the transcripts of genes coding for photosynthesis light reaction components were more abundant in M than in BS cells. Interestingly, over 90% of the genes related to protein synthesis (for ribosomal proteins) and protein folding were preferentially expressed in M cells, while those involved in protein degradation were predominantly expressed in BS cells. These observations indicate that many proteins were synthesized and folded in M cells but eventually degraded in BS cells, which surround the vasculature. This arrangement facilitates the transport of amino acids released from protein degradation to other plant parts. Genes involved in RNA regulation of transcription, transport, signaling, cell division, organization and vesicle transport, development, and major carbon, hydrogen, and oxygen metabolism were predominantly expressed in BS cells, but genes involved in secondary metabolism, tetrapyrrrole synthesis, and RNA binding showed an M cell expression preference. Most of the secondary metabolites accumulate in the epidermis cells, adjacent to M cells.

For the 85 BS-enriched genes involved in development, we found that 22 genes encode the nodulin MtN3 and MtN21 family proteins, which were originally identified as root nodule proteins of the legume Medicago truncatula (Gam et al., 1996). MtN3 and MtN21 also have recently been shown to be crucial for exine pattern formation and cell integrity of

### Table III. Differential expression of C₄ photosynthesis-specific genes between M and BS cells, calculated from the transcriptome data

<table>
<thead>
<tr>
<th>Gene Identifier</th>
<th>Symbol</th>
<th>Our Data</th>
<th>Li et al. (2010)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>M Cells</td>
<td>BS Cells</td>
</tr>
<tr>
<td>GRMZM2G083841</td>
<td>PEPC</td>
<td>9,485</td>
<td>431</td>
</tr>
<tr>
<td>GRMZM2G121878</td>
<td>CA</td>
<td>6,692</td>
<td>281</td>
</tr>
<tr>
<td>GRMZM2G129513</td>
<td>NADP-MDH</td>
<td>2,621</td>
<td>95</td>
</tr>
<tr>
<td>GRMZM2G383088</td>
<td>OMT</td>
<td>476</td>
<td>22</td>
</tr>
<tr>
<td>GRMZM2G131286</td>
<td>PPDK-RP</td>
<td>988</td>
<td>74</td>
</tr>
<tr>
<td>GRMZM2G085309</td>
<td>NADP-ME</td>
<td>87</td>
<td>12,192</td>
</tr>
<tr>
<td>GRMZM2G086258</td>
<td>DCT2</td>
<td>7</td>
<td>545</td>
</tr>
<tr>
<td>GRMZM2G001696</td>
<td>PCK</td>
<td>21</td>
<td>7,801</td>
</tr>
<tr>
<td>GRMZM2G07820</td>
<td>rbsS-4</td>
<td>87</td>
<td>4,256</td>
</tr>
<tr>
<td>GRMZM2G113033</td>
<td>rbsS-2</td>
<td>109</td>
<td>5,280</td>
</tr>
</tbody>
</table>

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microspores (Guan et al., 2008) and secondary wall formation in xylary and cotton fibers (Ranocha et al., 2010), respectively. These genes code for transmembrane proteins involved in the cell wall development of specialized plant cells that require stringent impermeability to specific substances (e.g. oxygen). Therefore, the counterparts in maize leaves might be involved in the development of the BS cell wall, which plays a key role in preventing the leakage of CO₂ released from C₄ acid decarboxylation in order to maintain a high CO₂ concentration in BS cells (Hatch et al., 1995). At present, little is known about the biochemical composition of the C₄ BS cell wall. For the first time, to our knowledge, our data suggest that C₄ plants may have adopted the exine composition of microspore and pollen for its impermeability to a number of substances, such as CO₂. Further analysis on the expression profile of these genes along the development of the Kranz anatomy in C₄ leaves should provide insights into their functional roles.

Cell Type-Enriched Transcription Factors

Among the 2,056 annotated transcription factor (TF) genes in maize, 654 and 820 were expressed in M and BS cells, respectively (Table II). The median RPKM for all expressed TFs in the two cell types were 5.7 and 7.1, respectively (Supplemental Data S4). We identified 53 TF genes as M enriched and 214 TF genes as BS enriched (Table II), which, according to the TF family database PlantTFDB version 2 (Zhang et al., 2011), belong to 16 families (Table II), respectively because of two different types of DNA-binding domains in a protein. Three TFs, MYB59 (GRMZM2G130149), MYB111 (GRMZM2G305856), and another MYB family protein (GRMZM2G002128), belong to both MYB and MYB-related families. The other three TFs are AGL12 (GRMZM2G105387), which belongs to both M-type and MIKC families, RPL (GRMZM2G154641), which belongs to both HB-other and TALE families, and a zinc finger (B-box type) family protein (GRMZM2G075562), which belongs to both CO-like and DBB families. Interestingly, in 11 of the families with cell type-enriched TF genes, all members showed BS enrichment, whereas none has all members showing M enrichment (Fig. 1E). This observation suggests that many TF families have a cell type preference, indicating a differentiation in regulatory network for the functional differentiation of the two photosynthetic cell types in maize leaves. Comparison of the distributions of cell type-enriched TF families between this study and Li et al. (2010) shows some similarities but many differences. For example, Li et al. (2010) identified only two ARF genes enriched in M cells and none enriched in BS cells (Fig. 5d in Li et al., 2010), while we identified seven ARF genes enriched in BS cells but none enriched in M cells (Fig. 1E). Cross-contamination of isolated M and BS cells will lead to underestimation of cell-specific genes.

We searched the literature to identify the functions or pathways where a TF might be involved (Table IV). Most M-enriched TFs are involved in the pathways of responses to environmental stimuli. For example, the three paralogs of DREB1A (GRMZM2G380377, GRMZM2G069146, and GRMZM2G069126) are related to the function of cold acclimation (Novillo et al., 2007). The two ERF TFs, CBF4 (GRMZM2G124037) and ERF7 (GRMZM2G089995), are involved in the pathways of responses to drought and abscisic acid (Haake et al., 2002; Song et al., 2005), respectively. In addition, three TFs in the NAC family and two in the WRKY family are induced by wounding and pathogens (Collinge and Boller, 2001; Xu et al., 2006; Zheng et al., 2006), respectively. Two other M-enriched TFs were GLK1 (GRMZM2G026833), which is involved in pathways of chloroplast development specific to M cells (Nakamura et al., 2009), and ARR10 (GRMZM2G013612), which is related to protoxylem differentiation (Yokoyama et al., 2007; Table IV).

Very different from the M-enriched TFs, most BS-enriched TFs are related to development or the biosynthesis of cells and organelles. These functions include chlorophyll biosynthesis, secondary cell wall biosynthesis, cell division, cell differentiation, or development of chloroplast and leaves. The observations also match some functional preferences of BS-enriched genes: categories of development and cell (division, organization, vesicle transport, etc.; Fig. 1D). Three TFs, MYB20 (GRMZM2G055158), MYB52 (GRMZM2G455869), and MYB54 (GRMZM2G077147), in the MYB family, and AITA1 (GRMZM2G179049) in the NAC family, are related to secondary cell wall biosynthesis (Zhong et al., 2006, 2008). Two TFs, TCP4 (AC205574.3_FG006) and TCP5 (GRMZM2G120151), in the TCP family are related to cell division (Cubas et al., 1999; Aggarwal et al., 2011). The GRAS TFs, RGA1 (GRMZM2G023872) and SHR (GRMZM2G132794), are related to asymmetric cell division (Truong et al., 1997; Heidstra et al., 2004). Two bHLH TFs, PIL5 (GRMZM2G165042) and FMA (GRMZM2G162450), are related to chlorophyll biosynthesis and cell differentiation (Huq et al., 2004; Gray, 2007), respectively. Three ARF TFs, ARF3 (GRMZM2G056120), ARF6 (GRMZM2G317900), and ARF8 (GRMZM2G078274), are related to leaf or floral organ development (Finet et al., 2010; Tabata et al., 2010; Table IV).

Interestingly, the ERF family was the only one that showed a larger number of M-enriched TFs than BS-enriched TFs. Many of these M-enriched ERF TFs have been shown to play a role in response to environment stimuli, but none of the BS-enriched ERF TFs are known to play such a role. Our data also showed that GOLDEN2 (GLK2; GRMZM2G087804) was preferentially expressed in BS cells of mature maize leaves at a moderate level (RPKM = 29) with R = 0.78 (Supplemental Data S1). GLK2 has been shown to be involved in maize chloroplast development in the BS cells independent of light, but it is not involved in rbcS accumulation (Hall et al., 1998; Cribb et al., 2001). On
Previously Known Genes Are Expressed in M and BS Cells Than the Other Hand, GLK1 Was Highly and Preferentially Expressed in M Cells (R = 0.89). These Data Suggest That GLK1 May Regulate Plastid Development in M Cells, While GLK2 May Regulate Plastid Development in BS Cells of C₄ Leaves.

More Genes Are Expressed in M and BS Cells Than Previously Known

Recently, Li et al. (2010) Examined the Transcriptomes of M and BS Cells Isolated from Fully Expanded Third Maize Leaves by LCM. About 41 Million 32-bp Sequencing Reads from the Two Cell Transcriptomes Were Obtained. In This Study, We Generated Approximately 370 Million 120-bp Reads from M and BS RNA Samples (Approximately 177 and Approximately 193 Million Reads, Respectively). Our Data Allowed the Identification of 18,482 Expressed Genes in Total, While Li et al. (2010) Detected Only 16,638 Expressed Genes. Another Possible Reason for the Above Difference Between the Two Studies Is That Li et al. (2010) Used an Earlier Annotation Version (APGv1) and Only Included Genes with Complementary DNA (cDNA) Support, While We Used APGv2. As a Result, the Total Number of Expressed Genes Obtained from Our Data Is Higher Than That Obtained by Li et al. (2010).

In Addition to the Expressed Genes in the FGS, We Detected an Additional 2,224 Expressed Genes in the WGS (Fig. 1A), With the Exclusion of Transposable Elements, Gene Fragments, and Pseudogenes. Many Genes in This Additional Gene Set Were Highly Expressed, With RPKM > 100 for 85 M-enriched Genes and 86 BS-enriched Genes. The Highest RPKM Values in This Set Were 4,900 (GRMZM5G809350; a Paralog of Lhcb1; CAB2) and 31,561 (GRMZM2G142891; a Paralog of β-glucosidase Aggregating Factor) in M and BS Cells, Respectively. Furthermore, We Detected 178 M-enriched and 610 BS-enriched Genes in the WGS. Although There Is No MapMan Annotation for the Genes in the WGS, 263 of the 2,224 Expressed Genes in the WGS Are Found to Have One or More Gene Ontology Terms (http://www.geneontology.org/); the Five Most Abundant Terms in Molecular Functions Are ATP Binding (23%), Ice Binding (13%), Zinc Ion Binding (12%), DNA Binding (10%), and Protein Binding (10%). Three M-enriched Genes (GRMZM2G411457, GRMZM2G030616, and GRMZM2G152576) Are Related to Light Reaction in Photosynthesis (Supplemental Data S1).

Notably, Our Data Reveal More Genes Expressed in BS Than in M Cells in Mature Maize Leaves. Expressed Genes From Vascular Tissues (E.g. Sieve Element) in the BS Cell Preparation Might Have Contributed to the Category of BS-expressed Genes, as the BSS Isolated in

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Table IV. Selected cell type-enriched TFs and related functions or pathways

<table>
<thead>
<tr>
<th>Cell</th>
<th>TF Family</th>
<th>Gene Identifier</th>
<th>M</th>
<th>BS</th>
<th>R</th>
<th>Gene Name</th>
<th>Function or Pathway</th>
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<tbody>
<tr>
<td>M</td>
<td>G2-like</td>
<td>GRMZM2G026833</td>
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<td>0.81</td>
<td>ARR10</td>
<td>Protoxylem differentiation</td>
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</tr>
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<td>DREB1A</td>
<td>Cold acclimation</td>
</tr>
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<td>0.92</td>
<td>DREB1A</td>
<td>Cold acclimation</td>
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<td>0.94</td>
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<td>Induced by wounding</td>
</tr>
<tr>
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<td>0.92</td>
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<td>M</td>
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<td>1.73</td>
<td>0.91</td>
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<tr>
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<td>GRMZM2G165042</td>
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<td>41.36</td>
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</tr>
<tr>
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<td>42.49</td>
<td>0.97</td>
<td>TCP5</td>
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</tr>
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<td>1.00</td>
<td>TCP4</td>
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</tr>
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<td>Cell differentiation</td>
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<td>0.95</td>
<td>ARF3</td>
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<tr>
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<td>1.00</td>
<td>RGA1</td>
<td>Asymmetric cell division</td>
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<td>2.29</td>
<td>0.99</td>
<td>SHR</td>
<td>Asymmetric cell division</td>
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</tbody>
</table>

*TF family names are as follows: G2-like, GOLDEN2-like; ARR-B, type B phospho-Accepting Response Regulator; WRKY, conserved WRKY domains, about 60 amino acid residues with the WRKYQK sequence followed by a C2H2 or C2HC zinc finger motif; ERF, Ethylene-Responsive Element Binding Factor; NAC, NAM (no apical meristem), ATAF, and CUC (cup-shaped cotyledon) TFs; bHLH, basic helix-loop-helix proteins; MYB, myeloblastosis virus; TCP, TEOSINTE BRANCHED1 from maize, CYCLOIDEA from snapdragon (Antirrhinum majus), and PROLIFERATING CELL FACTOR1 and -2 from rice (Oryza sativa); ARF, Auxin Response Factor; GRAS, GAI, RGA, and SCR genes.
this study contained the vascular tissue (Supplemental Table S4) and sieve elements are active cells for transport function. Thus, as discussed above, a small portion of the RNA from vascular tissues might have contributed to our BS transcriptome. However, it is most likely that more genes are expressed in BS cells than in M cells because of functional necessity, in agreement with the observation that the BS cells in C_4 plants play much more important roles than the M cells in various physiological processes (Majeran et al., 2005; Leegood, 2008). In addition to genes related to C_4 acid decarboxylation, expression of specific genes related to the metabolism of photorespiration, nitrate, sulfate, ammonium, starch, ion transport, secondary metabolites, and cell wall structure is largely confined to BS cells.

**Light Reaction, Carbon Fixation Pathway, and Related Transporters**

Many of the maize genes coding for the PSI components are highly expressed in both cell types, but with higher expression levels in M cells (Supplemental Data S3). In contrast, many of the maize genes coding for the PSII components are highly and predominantly expressed in M cells, with \( R_i > 0.9 \) for most of the genes. Transcripts for the two genes coding for ATP synthase, respectively. These data are in agreement with the observations that maize BS cells possess a higher capacity for cyclic electron transport (Ivanov et al., 2007).

Based on the expression profiles of C_4-specific genes (Table III; Supplemental Data S1 and S3), we propose the partitioning of the key biochemical reactions of the C_4 photosynthesis pathway between M and BS cells of maize, an NADP-ME subtype C_4 plant (Fig. 2). Hydration of CO_2 to bicarbonate, generation of PEP from pyruvate, carboxylation of bicarbonate to OAA, and its subsequent transamination to Asp or reduction to malate in the chloroplast all take place in the M cells. In contrast, decarboxylation of OAA to PEP and CO_2 and decarboxylation of malate to pyruvate and CO_2 occur in the cytosol and chloroplast of BS cells, respectively. The high expression levels of NADP-ME and PCK suggest that the PCK decarboxylation system plays a substantial role in maize C_4 photosynthesis, consistent with previous biochemical studies (Furumoto et al., 1999, 2000; Wingler et al., 1999; Furbank, 2011). The dual decarboxylation system in some NADP-ME subtype C_4 plants, such as maize, probably makes them the most efficient photosynthetic plants.

High rates of photosynthesis by C_4 plants imply high fluxes of metabolites intercellularly and intracellularly. In C_4 plants, most of the photosynthetic metabolites are known to be transported symplastically between M and BS cells through plasmodesmata. However, within each cell type, transporters on the chloroplast envelope must be active to substantiate the heavy traffic of related metabolites between cytosol and chloroplast. A maize gene coding for a 2-oxoglutarate/malate translocator (ZmpOMT1; GRMZM2G383088) was highly and predominantly expressed in M cells, and a dicarboxylate transporter gene (ZmpDCT1; GRMZM2G040933) was also preferentially expressed in M cells at a moderate level. Thus, these two transporters may be responsible for the uptake of OAA and the export of malate by M chloroplasts, respectively (Taniguchi et al., 2004). In contrast, the transcript for another maize gene coding for DCT (ZmpDCT2; GRMZM2G086258) was highly and almost exclusively expressed in BS cells, which may be responsible for the import of malate by BS chloroplasts, as suggested previously by Taniguchi et al. (2004).

In C_3 plants, the chloroplast phosphate translocator triose phosphate/phosphate translocator (TPT) is thought to have a central role in the partitioning of fixed carbon between starch in the chloroplast and Suc in the cytosol. Inorganic phosphate is taken up by the translocator in exchange for triose phosphate, which is then metabolized to Suc in the cytosol. In NADP-ME subtype C_4 plants, BS chloroplasts are deficient in PSII, so the triose phosphate produced must be transported to M chloroplasts for reduction. Consistent with this role, the transcript of a maize TPT gene (APE2; GRMZM2G070605) was expressed with very high RPKM values of 2,069 and 1,551 in M and BS cells, respectively (Supplemental Data S3).

In C_4 plants, pyruvate produced from malate decarboxylation in the BS chloroplasts must be returned to M chloroplasts for regeneration of PEP via PPTD with the input of ATP, and PEP is subsequently exported to the cytosol for carboxylation by PEPC. Earlier biochemical and molecular studies suggested that to accommodate these new transport functions in C_4 plants, TPT has been adopted and modified for the exchange of phosphate and PEP (PPT; Flugge, 1999), and two types of light-dependent pyruvate transporter have been employed by the M chloroplasts for the uptake of pyruvate in a wide range of C_4 plants, one sodium dependent (Ohnishi et al., 1990; Ohnishi and Kanai, 1990; Aoki et al., 1992) and one proton dependent (Aoki et al., 1992). The NADP-ME subtype C_4 *Flaveria* spp. use the sodium/pyruvate symporter (Gowik et al., 2011), but maize, a NADP-ME subtype monocot, uses the proton/pyruvate symporter (Ohnishi et al., 1990; Ohnishi and Kanai, 1990; Aoki et al., 1992). The gene coding for the sodium-dependent pyruvate transporter BASS2, a bile acid:sodium symporter family protein 2, has been cloned recently from C_4 *Flaveria* spp. (Furumoto et al., 2011), but the gene(s) coding for the proton-dependent one remains elusive. Transcripts of several maize PPTs were detected in this study. One of them (GRMZM2G174107) was highly enriched in
M cells, whereas a second one (GRMZM2G066413) was moderately enriched in BS cells. Thus, the GRMZM2G174107 gene may specifically code for the exchanger of triose phosphate and PEP in the M chloroplasts of maize leaves. In addition, our transcriptome data showed that for the three MEP (for methyl erythritol phosphate) genes coding for the inner envelope transporter proteins, MEP3 (GRMZM2G138258) was expressed exclusively in BS cells (Ri = 0.99) at an extremely high level (RPKM = 2,734), GRMZM2G305851 was moderately expressed with a preference in M cells, whereas MEP2 (GRMZM2G077222) was equally expressed in both cell types at relatively low levels. The exact physiological roles of MEP transporters in maize have not been established, but all three MEP transporters have plastidial transporters. Since pyruvate is one of the two crucial substrates of the MEP pathway in plastids, we propose that the two MEP3 paralogs may code for the inner chloroplast envelope proteins responsible for pyruvate export from BS chloroplasts and import of pyruvate by M chloroplasts, respectively, whereas MEP2 may mainly function to supply pyruvate to the MEP pathway, as in C₃ plants. In agreement with our data, Friso et al. (2010) suggested earlier that MEP2,4 and MEP3/4 may function as pyruvate transporters in M and BS chloroplasts, respectively, based on proteomic studies of isolated maize M and BS chloroplasts.

Although photorespiration is not apparent in C₄ plants under ambient atmospheric conditions, measurable photorespiratory activity can be detected at low CO₂ conditions (de Veau and Burris, 1989; Dai et al., 1993) and in young leaves (Dai et al., 1993, 1995). Consistent with the localization of Rubisco in the BS chloroplasts of C₄ plants, the expression of genes coding for most of the key photorespiratory enzymes exhibited a BS specificity (Supplemental Data S3). High levels of transcript were detected in the BS cells for glycolate oxidase (GRMZM2G129246), Gly decarboxylase (GRMZM2G399183 and GRMZM2G104310), and Ser hydroxymethyltransferase (GRMZM2G135283), with R values ranging from 0.94 to 1.00. However, expression of two of the photorespiratory pathway enzymes, chloroplast phosphoglycolate phosphatase and peroxisomal NAD-hydroxypyruvate reductase,
Development, Cell Wall, and Plasmodesmata Formation

Kranz leaf anatomy is essential for most C₄ plants to concentrate CO₂ around Rubisco in the chloroplasts of BS cells to suppress its oxygenase and the associated photorespiratory activity. Consequently, the BS cell wall in many C₄ plants is thickened and specialized with a suberin layer to prevent CO₂ leakage (Evans and Von Caemmerer, 1996; Leegood, 2002). Our data depict the expression of many cell wall-associated genes specifically expressed in BS cells, which include genes for cell wall-associated kinase (GRMZM2G135291 and GRMZM2G062471) and several highly expressed nodulin-type proteins (MtN3 and MtN21). Therefore, these genes may be related to the differentiation of C₄ BS cell walls. In contrast, only one cell wall-related gene coding for expansin (GRMZM2G094990) is specifically expressed in M cells.

Intercellular transport of metabolites between M and BS cells (Evert et al., 1977; Robards and Lucas, 1990) and between the BS and the adjacent vascular parenchyma cells (Botha et al., 2000) in C₄ leaves is mainly by diffusion through interconnecting plasmodesmata, which are concentrated along the cell wall between these cell types. Interestingly, our data reveal that the genes coding for plasmodesmata-located proteins (signaling receptor kinases), plasmodesmata callose-binding proteins (β-1,3-glucan hydrolases), and callose synthase are almost exclusively expressed in BS cells (Supplemental Data S3), suggesting that genes related to plasmodesmata development may be controlled by the regulatory network of BS cells. Callose, which differs from cellulose in consisting of a β-1,3-glucan chain, is known to be made by a few cell types at a specific stage during cell wall development, such as microspore and pollen. It is also ectopically deposited over the cell wall between the BS and vascular parenchyma cells in maize leaf minor veins, thereby occluding the plasmodesmata (Botha et al., 2000). Therefore, we speculate that a similar cell wall composition may be shared by nodule, microspore, pollen, xylary fiber, and C₄ BS cells. C₄ plants regulate the expression of these cell wall-related genes in an organ- and cell-specific manner to function in the CO₂-concentrating mechanism.

CONCLUSION

The C₄ physiology is an integrated syndrome of developmental, anatomical, cellular, and biochemical traits that must rely on regulatory networks that control the functional differentiation of M and BS cells. Our transcriptome profiling of the two photosynthetic cell types of mature maize leaves reveals the cell differentiation of gene expression at both regulatory and functional levels. To our knowledge for the first time, genes coding for the transporters of several C₄ metabolites and the development of the BS cell wall have been identified that express in a cell-specific manner. Clearly, both cell types are specialized in different metabolic functions but are also coordinated to provide a high efficiency in carbon, mineral, and water use. In addition, many TF genes have been shown to express in a cell-specific manner. Thus, the comprehensive data sets from this study have laid a solid foundation for further study of the regulation of the expression and evolution of C₄-specific and -related genes.

Although C₄ plants represent only approximately 3% of plant species, they contribute to about 30% of terrestrial carbon fixation, and 60% of C₄ plants are grasses, which include some of our most important food crops, such as maize, sugarcane (Saccharum officinarum), millet (Pennisetum americanum), and sorghum (Sorghum bicolor) of the NADP-ME subtype. Despite this scarcity, some of the most productive crops and noxious weeds on earth belong to the C₄ plants, which make this photosynthetic property a desirable trait for introduction into the less productive C₃ crops (Hibberd et al., 2008). However, at present, very little is known about the molecular regulatory networks underlying the development of Kranz leaf anatomy and the functional differentiation of its photosynthetic cells in C₄ photosynthesis and other metabolism. A comprehensive understanding of the C₄ syndrome is required before we can effectively transfer the C₄ traits into C₃ crops, such as rice (Oryza sativa), wheat (Triticum aestivum), and soybean (Glycine max).

A thorough and systematic examination of the regulatory networks involved in C₄ syndrome from comparative C₃, C₃-C₄ intermediate, and C₄ transcriptome study (Gowik and Westhoff, 2011) and from developmental and cellular transcriptome profiling of C₄ leaves (Li et al., 2010; this study) should streamline this process.

MATERIALS AND METHODS

Plant Growth Conditions and Leaf Sample Collection

Seeds of maize (Zea mays 'White Crystal'), a glutinous cultivar, were germinated and cultivated in the greenhouse in June 2011 under natural sunlight conditions (13–14 h) with a maximum photosynthetic photon flux density of 1,600 μmol m⁻² s⁻¹, day/night temperatures of 28°C to 32°C/25°C to 28°C, and 60% to 70% relative humidity. The second newly expanded leaves of 9-d-old plants were harvested around 9 AM for isolation of M and BS cells in the laboratory and extraction of total RNA.

Enzymatic Isolation of M Protoplasts

MP were isolated from the mid section (5 cm) of newly matured second leaves by digesting leaf segments in a cell wall digestive medium, according to
the procedures of Kanai and Edwards (1973), Sheen (1995), Markelz et al. (2003), Sawers et al. (2007), and Sharpe et al. (2011) with minor modifications. All solutions and glassware were sterilized either by filtration or by treatment with diethyl pyrocarbonate water followed by autoclave. The digestive medium, containing 1.5% (w/v) cellulose (Onozuka RS), 0.1% (w/v) macer- ozyme (Onozuka R-10), 20 mM MES (pH 5.8), 0.6 M sorbitol, 5 mM β-mercaptoethanol, 1 mM CaCl₂, 0.1% bovine serum albumin, and the medium was then heated to 55°C for 10 min and cooled to room temperature before the addition of sorbitol and digestive enzymes. The medium was then heated to 55°C for 10 min to solubilize the enzymes as well as to inactivate DNase, and other components were added. Several leaf blades were stacked and cut perpendicularly to the long axis into 0.5- to 1-mm slices, were repeated. BS cells that passed through the 500-μm mesh was resuspended in an 80-μm nylon net to retain undigested tissue and BS, while MP (15-35 μm diameter) in the filtrate was collected by centrifugation at 300 × g for 5 min. The crude MP pellet was gently resuspended in 4 to 6 mL of 13% (w/v) dextran-Suc solution (13% dextran, 0.6 M Suc, 5 mM HEPES-KOH, pH 7.0, 100 mM β-mercaptoethanol, 1 mM CaCl₂ to release more MP. The filtrate was then filtered through an 80-μm nylon net to retain undigested tissue and BS, while MP, which was then washed with the cold isolation medium to remove attached M tissue and retained on an 80-μm nylon net. The purity of BS cells was monitored with sterilized razor blades and then quickly transferred into a 50-mL sterile beaker containing 20 mL of digestive medium. The medium with leaf slices was vacuum infiltrated twice and incubated at 23°C under low light (100 μmol m⁻² s⁻¹) for 3 h without shaking.

At the end of digestion, the mixture was first filtered through a 500-μm mesh, and the materials retained on the mesh were gently mixed with 100 mL of sterile sorbitol wash medium (0.6 M sorbitol, 5 mM HEPES-KOH, pH 7.0, 100 mM β-mercaptoethanol, and 1 mM CaCl₂) to release more MP. The filtrate was then filtered through an 80-μm nylon filter to release more MP. Leaf slices, prepared as described above, were quickly added to 50 mL of cold BS cell isolation medium and homogenized twice (10 s each) at high speed in a Waring blender. After blending, the mixture was first filtered through a 500-μm mesh, and the crude tissue retained on the 500-μm mesh was resuspended sequentially with 2 mL of 10.7% (w/v) dextran-Suc solution and 2 mL of sorbitol wash medium. The gradient was centrifuged at 300g in a swing-bucket rotor for 20 min, and pure MP was collected from the interphase between the sorbitol wash medium and 10.7% dextran-Suc solution and diluted in a small volume of sorbitol wash medium before isolation of RNA. The purity of isolated MP was examined with a light microscope.

Mechanical Isolation of BSS

To avoid damage to BS cells during enzymatic digestion, a mechanical method was adopted for rapid isolation of BS cells. The isolation medium contained 0.6 M sorbitol, 50 mM Tris-HCl (pH 8.0), 5 mM EDTA, 0.5% polyvinylpyrrolidone-10, 10 mM DTT, and 100 mM β-mercaptoethanol. Leaf slices, prepared as described above, were quickly added to 50 mL of cold BS cell isolation medium and homogenized twice (10 s each) at high speed in a Waring blender. After blending, the mixture was first filtered through a 500-μm mesh, and the crude tissue retained on the 500-μm mesh was resuspended in another 50-mL isolation medium, and the blending and filtration processes were repeated. BS cells that passed through the 500-μm mesh were filtered and retained on an 80-μm nylon filter. The purity of BS cells was monitored with a light microscope, and if deemed necessary, the collected BS cells were further washed with the cold isolation medium to remove attached M tissue and the epidermal tissue floating on top of the isolation medium. The whole isolation procedure was completed within 10 min.

Enzyme Assay and Immunoblot Analysis

The activities of NADP-ME and PEPC were assayed at 30°C using a spectrophotometer according to Kanai and Edwards (1973). The procedure of Murphy et al. (2007) was followed for immunoblot analysis using specific antibodies.

RNA Isolation, Library Preparation, and Deep Sequencing

Total RNA from isolated cells was extracted using Trizol reagent (Invitrogen) and purified using acid phenol-chloroform extraction. The quality of extracted RNA was examined by gel electrophoresis and by BioAnalyzer (Agilent). Using the Illumina mRNA-seq kit, 10 to 15 μg of a total RNA sample was purified for poly(A) RNA and subjected to fragmentation for 3 min at 94°C. The mRNA fragments were then converted into single-stranded cDNA, and the ends were modified and ligated to the paired-end adaptors. The ligation products were size selected on agarose gels (independent slices of 250-350 and 350-450 bp), amplified by 15 cycles of PCR, cleaned up using Ampure beads (Beckman Agencourt), and examined by Qubit (Invitrogen) and BioAnalyzer High Sensitivity DNA chip (Agilent). Paired-end 2 × 120-nucleotide se- quencing was performed on Illumina Genome Analyzer IIx, and two lanes of data were obtained for each cell type by sequencing two independently sized selected libraries. Approximately 177.4 million and 193.4 million reads were obtained from the M and BS RNA samples, respectively.

Deep-sequencing Data Processing, Mapping, and Expression Level Calculation

We processed and mapped raw reads to the maize genome (B73) using the mapping tool Tophat version 1.3.3 (http://tophat.ccb.org/; http://tophat.ccb.org/) with default parameters (maximum allowed mismatches, two; maximum number of hits for a read, 10). Based on the AFCoV annotation downloaded from http://ftp.maizesequence.org/; we calculated the expression levels of each annotated gene in RPKM (Mortazavi et al., 2008) using the expression calculation tool Cufflinks version 1.2.1 (http://cufflinks.cbcb.umd.edu/).

Estimation of Cell Type Specificity

As described in “Results,” to estimate the degree of differential expression of genes between the two cell types, we defined the degree of cell specificity of gene i as Ri = 1 - mi(1 - b)/(max(mi, b)), where mi and b represent the RPKM of gene i in the M and BS RNA samples, respectively. Applying this measure to the transcriptome data, one can study the gene expression differences between the M and BS cells.

Estimation of Contamination Rates

We can use the transcriptome data to obtain an approximate estimate of the contamination rate in the M (or BS) cells isolated. Let ci be the percentage contamination of the isolated M cells by BS cells. That is, ci is the proportion of BS cells in the sample of M cells. For gene i, let x and y be the expected expression levels in M cells and BS cells, respectively, under no contamination. Then, the expected expression level of gene i in the isolated M cells, denoted by mi, should be given by

\[ m_i = c_1 x + c_2 y. \]

Equation 1 cannot be solved because there are two unknowns, c1 and c2. However, if gene i is supposed not to be expressed in M cells, we can assume x = 0 and obtain c1 = m_i/y. Since using one gene to estimate c1 may not be reliable, we estimate c by

\[ c_i = \frac{\sum m_i}{\sum y_i}, \]

where the summation is over the genes that are supposed to be expressed only in BS cells but not in M cells. We can take the estimate as an upper estimate because one or more of the genes used may not be absolutely only expressed in BS cells.

Similarly, let c2 be the rate of contamination of BS cells by M cells and let b be the observed level of expression in the transcriptome data of the BS cells. Then, we have

\[ b = c_2 x + (1 - c_2) y, \]

\[ c_2 = \frac{\sum b}{\sum x}, \]

where the summation is over the genes that are supposed to be expressed only in M cells but not in BS cells.

Functional Annotation

The identified cell type–enriched genes were annotated with biological functions and metabolic pathways by MapMan (Thimm et al., 2004; http://mapman.gabipd.org/). The TF information was downloaded from Plant TFDB version 2.0 (Zhang et al., 2011; http://planttfdb.cbi.pku.edu.cn/). The read data have been submitted to the National Center for Biotechnology Information Short Read Archive under accession number SRP009063.

Supplemental Data

The following materials are available in the online version of this article.

Transcriptomes of Maize Leaf Cells
Chang et al.

Supplemental Figure S1. Micrographs of isolated M and BS cells.

Supplemental Figure S2. Immunoblot analysis in mature leaves and isolated M and BS cells (two preparations).

Supplemental Figure S3. Median depth of read-mapping coverage along the cDNA.

Supplemental Table S1. Activities of PEPC and NADP-ME in the whole leaf and in the isolated M and BS cells of maize.

Supplemental Table S2. Validation of cell type-specific gene expression.

Supplemental Table S3. List of quantitative reverse transcription-PCR primers, Universal Probe Library probe, and PCR efficiency.

Supplemental Table S4. RPKM of selected genes of vascular and epidermal tissues.

Supplemental Data S1. RPKM values of all genes with at least one read in one of the two transcriptomes.

Supplemental Data S2. List of genes with R > 0.99 in the two cell types.

Supplemental Data S3. Differential expression between M and BS cells of genes.

Supplemental Data S4. List of cell type-enriched TFs.

Supplemental Materials and Methods S1.

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Transcriptomes of Maize Leaf Cells


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