Individual Maize Chromosomes in the C₃ Plant Oat Can Increase Bundle Sheath Cell Size and Vein Density¹[W][OA]

Ben J. Tolley, Tammy L. Sage, Jane A. Langdale, and Julian M. Hibberd*

Department of Plant Sciences, University of Cambridge, Cambridge CB2 3EA, United Kingdom (B.J.T., J.M.H.); Department of Ecology and Evolutionary Biology, University of Toronto, Toronto, Ontario M5S 3B2, Canada (T.L.S.); and Department of Plant Sciences, University of Oxford, Oxford OX1 3RB, United Kingdom (J.A.L.)

C₄ photosynthesis has evolved in at least 66 lineages within the angiosperms and involves alterations to the biochemistry, cell biology, and development of leaves. The characteristic “Kranz” anatomy of most C₄ leaves was discovered in the 1890s, but the genetic basis of these traits remains poorly defined. Oat × maize addition lines allow the effects of individual maize (Zea mays; C₄) chromosomes to be investigated in an oat (Avena sativa; C₃) genetic background. Here, we have determined the extent to which maize chromosomes can introduce C₄ characteristics into oat and have associated any C₄-like changes with specific maize chromosomes. While there is no indication of a simultaneous change to C₄ biochemistry, leaf anatomy, and ultrastructure in any of the oat × maize addition lines, the C₃ oat leaf can be modified at multiple levels. Maize genes encoding phosphoenolpyruvate carboxylase, pyruvate, orthophosphate dikinase, and the 2′-oxoglutarate/malate transporter are expressed in oat and generate transcripts of the correct size. Three maize chromosomes independently cause increases in vein density, and maize chromosome 3 results in larger bundle sheath cells with increased cell wall lipid deposition in oat leaves. These data provide proof of principle that aspects of C₄ biology could be integrated into leaves of C₃ crops.

The alterations to C₄ leaves allow the initial fixation of HCO₃⁻ by phosphoenolpyruvate carboxylase (PEPC) to be separated from the release of CO₂ in a separate compartment that contains Rubisco (Hatch, 1987). This release of CO₂ around Rubisco can be catalyzed by three different four-carbon decarboxylases. Traditionally, this has led to the classification of C₄ plants into one of three biochemical subtypes, depending on whether they primarily use NADP-dependent malic enzyme (NADP-ME), NAD-dependent malic enzyme (NAD-ME), or phosphoenolpyruvate carboxykinase (Hatch et al., 1975; Hatch, 1987; Leegood, 2002). However, there appears to be reasonable mechanistic flexibility that is determined both temporally and environmentally (Walker et al., 1997; Wingler et al., 1999; Furumoto et al., 1999), which suggests that the distinctions between these three biochemical subtypes are less rigid than originally thought (Furbank, 2011). As well as this diversity within the biochemistry of C₄ photosynthesis, there is also variation in terms of the leaf anatomy associated with the C₄ pathway. Although several examples of C₄ photosynthesis operating in single cells have been described (Reiskind et al., 1997; Casati et al., 2000; Reinfelder et al., 2000; Voznesenskaya et al., 2002, 2003), most C₃ species compartmentalize carboxylation and decarboxylation activities into two distinct cell types. At least 22 variations of two-celled “Kranz” anatomy have been documented (Dengler and Nelson, 1999; Edwards and Voznesenskaya, 2011; Sage et al., 2012) whereby different cell types have been coopted as the photosynthetic carbon assimilation and photosynthetic carbon reduction tissue. In the grasses, nine types of Kranz anatomy have been described (Edwards and Voznesenskaya, 2011; Sage et al., 2012).
Five of these nine types have two sheaths that surround each vascular bundle. The inner sheath has endodermal characteristics and is called the mestome sheath (MS); it is surrounded by a parenchymal bundle sheath (PBS; Brown, 1958). In the other four types, the MS cell layer is absent and the PBS is adjacent to the metaxylem tissue of the vascular bundle (Brown, 1958). In all of these Kranz variants, Rubisco is localized in the PBS, where it is used to fix CO₂.

The molecular basis underpinning the Kranz anatomy of most C₄ leaves remains to be defined. In maize, which has “classical” NADP-ME anatomy, the leaf is organized in repeating units of mesophyll (M)-PBS-vein (V)-PBS-M (Langdale and Nelson, 1991) with no MS cell layer. Although mutants of maize that show perturbations to plastid morphology and expression of photosynthesis genes in M or PBS cells of C₄ leaves have been identified (Langdale and Kidner, 1994; Brutnell et al., 1999; Smith et al., 2001), they have so far provided little insight into the genetic basis of Kranz anatomy.

In addition to our relatively poor understanding of the mechanisms underlying Kranz anatomy in C₄ leaves, it is also unclear whether elements of Kranz anatomy can be introduced into leaves of C₃ plants. As leaves, it is also unclear whether elements of Kranz anatomy can be introduced into leaves of C₃ plants. As leaves, it is also unclear whether elements of Kranz anatomy can be introduced into leaves of C₃ plants.

RESULTS

Maize Chromosomes Can Increase Vein Density in C₃ Oat

Little is known either about genes controlling the development of increased venation in C₄ leaves or whether venation can be modified in C₃ leaves. We used OMA lines generated from two oat parental backgrounds (sun II and starter) to investigate these two fundamental processes. Each maize chromosome was represented by at least one addition line, and in several cases an individual maize chromosome was represented by more than one line, with the lines sharing the same chromosome but differing in their parental backgrounds (Supplemental Table S1). In order to investigate whether the presence of individual maize chromosomes increased venation in oat, distances between the midpoints of adjacent veins were measured in transverse sections of fully expanded third leaves from OMA lines via light microscopy. While veins of maize were separated by approximately 130 μm, in both oat parental lines mean distances between adjacent veins were approximately 500 μm (Fig. 1A; Supplemental Fig. S1). Interven distance (IVD) in OMA lines was variable, indicating that individual maize chromosomes in the oat background exerted an influence on vein spacing (Fig. 1A). In particular, the independent addition of maize chromosomes 1, 5, and 9 was associated with significantly reduced vein spacing compared with oat parents (Fig. 1A). Although none of the lines individually recapitulated maize-like values, additively the differences of OMA 1.36, OMA 5.59, and OMA 9.41 from wild-type oat would produce values similar to those of maize. To determine whether the proximity of adjacent veins in these three lines could be accounted for by a reduction in the number of intervening M cells, the number of palisade M cells separating adjacent vascular bundles was measured. Although the number of M cells between neighboring veins was variable in the OMA lines, none showed values that were significantly different from the oat parental lines (Fig. 1B; Supplemental Fig. S1).

Maize Chromosomes Alter Leaf Anatomy in Oat

C₄ photosynthesis in maize is associated with large PBS cells that contain more chloroplasts than those of C₃ leaves. Since very little is known about loci controlling PBS cell size, the cross-sectional area of the PBS cells in each of the OMA lines was measured and compared with oat and maize controls. Lines possessing maize chromosome 1, 2, 3, 7, or 8 and both lines with maize chromosome 9 had significantly larger PBS areas than either of the oat parental lines. This suggests that these chromosomal additions independently specified larger PBS cells (Fig. 1C; Supplemental Fig. S1).

In some lines, the increases in cell size observed in the OMA lines were not confined to the PBS, with M cells also larger compared with oat parental lines (Supplemental Fig. S2). In order to quantify the extent
to which increases in cell size were specific to the PBS and not a consequence of constitutive cell expansion in the leaf, the area of PBS between the midpoints of two adjacent veins was expressed relative to the total intervening M area between the veins. Line 3.01 had a significantly larger PBS-M ratio than both of the oat parental lines, with a value 70% of the maize PBS-M ratio (Fig. 1D). This indicates that loci controlling PBS cell size are likely located on maize chromosome 3. However, the introduction of maize chromosome 3 was not sufficient to recapitulate other elements of maize PBS morphology, such as high chloroplast numbers and centrifugal chloroplast position, and in fact there was no evidence of increased chloroplast proliferation in any of the OMA lines (Supplemental Fig. S2).

Maize Chromosome 3 Introduction Causes Lipid Deposition in Oat PBS Cell Walls

In order to determine the extent to which the enlarged PBS cells in OMA 3.01 resembled the maize PBS, transverse sections of leaves were stained with Nile red to detect fatty acid deposition. Nile red stains positively for lipids in plants (Fowler and Greenspan, 1985; Greenspan and Fowler, 1985; Greenspan et al., 1985; Pighin et al., 2004; Schmidt and Herman, 2008; Dietrich et al., 2009), fluorescing red in the presence of polar lipids and yellow when staining esterified cholesterol and triacylglycerols (Diaz et al., 2008). This approach showed polar lipids in the PBS of maize, but in the oat parental line, only the MS and some of the veinal cells stained (Fig. 2, A and B). In contrast, in OMA 3.01, the PBS stained positively when examined by fluorescence microscopy (Fig. 2C), indicating that PBS cells in line 3.01 have increased deposition of polar lipids in the PBS cell wall. Transmission electron microscopy (TEM) imaging of the leaves indicated the presence of suberin in the cell walls of the PBS in maize but only in the MS cell layer of OMA 3.01 and wild-type oat (Fig. 2, D–F). Taken together, these data indicate that chromosome 3 likely contains loci that can control PBS cell wall lipid deposition in oat but not the entire suberin deposition pathway.

Figure 1. Alterations to leaf morphology in OMA lines. A and B, Distance between adjacent veins (A) and number of palisade M cells separating adjacent vascular bundles (B). C and D, PBS cell cross-sectional area (C) and PBS-M cross-sectional area ratio (D) in maize, oat, and OMA leaves. Results are expressed as means ± se of three biological replicates with a minimum of 10 measurements per leaf. Starred results are statistically significant relative to oat controls (P < 0.01).

Figure 2. Maize chromosome 3 causes lipid deposition in oat PBS cell walls. Fluorescence (A–C) and TEM (D–F) images of transverse sections of maize (A and D), oat (sun II; B and E), and OMA 3.01 (C and F) leaves are shown. Lipid detection was carried out by staining sections with Nile red (A–C). Suberin detection was by TEM (D–F). Asterisks indicate MS. In TEM images, arrows indicate suberin layers. Bars = 20 μm (A–C) and 500 nm (D–F).
Individual Maize Chromosomes Have Little Impact on Characteristics of Leaf Photosynthesis

Previous work has shown that in OMA lines with maize chromosome 6 or 9, which contain ZmPPDK or ZmPEPC, respectively, there is little impact on CO$_2$ compensation point (Kowles et al., 2008). However, it is possible that alterations to photosynthetic parameters could be induced in oat if a specific maize transcription factor or factors recognized and regulated a significant number of oat photosynthesis genes. In order to determine the extent to which photosynthetic parameters were influenced by the introduction of each maize chromosome, net photosynthesis of the OMA lines, maize, and oat parental lines were measured over a range of external CO$_2$ concentrations and photon flux densities (PFDs; Figs. 3 and 4). In C$_4$ plants, photosynthesis responds more rapidly to CO$_2$ supply than in C$_3$ species, because PEPC displays a higher affinity for HCO$_3^-$ This leads to a steep initial slope (high carboxylation efficiency) and an intercept with the x axis close to zero (low CO$_2$ compensation point).

These characteristics were evident in maize compared with the oat parental lines starter and sun II (Fig. 3). Interestingly the two oat parental lines showed different maximum rates of photosynthesis at high CO$_2$ concentration. When this is taken into account, CO$_2$ responses in most OMAs were typical of C$_3$ photosynthesis and for the most part unaffected by the presence of each maize chromosome, giving similar values to the oat parents (Fig. 3). However, three lines (6.32, 9.41, and 10.23) showed lower saturating levels of photosynthesis, and carboxylation was dramatically reduced at low internal CO$_2$ values in line 9.41 (Fig. 3), indicating that the introduction of chromosome 9 had perturbed carbon assimilation. The introduction of maize chromosome 9 in line 9.01, however, did not show inhibited carbon assimilation.

In C$_4$ leaves, photosynthesis typically saturates at much higher light intensities compared with C$_3$ leaves. In order to test whether any of the maize chromosomes resulted in C$_4$-like light responses, photosynthetic rates of each line were measured over a range of light intensities. Whereas in maize, photosynthesis did not

![Figure 3](image)

**Figure 3.** Photosynthetic assimilation rates are largely unchanged by maize chromosomal introduction. Light-saturated photosynthetic assimilation rates ($A$) over a range of leaf intercellular CO$_2$ concentrations ($C_i$) for leaves of maize, oat, and OMA lines are shown. Each panel represents mean assimilation rates of maize and wild-type oat or chromosomal means of OMA lines. Results are expressed as means of at least three biological replicates ± se. Measurements were taken at 25°C, leaf vapor pressure deficit of less than 1.5 kPa, and PFD of 1,500 µmol m$^{-2}$ s$^{-1}$. 

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saturate below 3,000 \( \mu \text{mol photons m}^{-2} \text{s}^{-1} \), in all OMAs it saturated at values below 1,000 \( \mu \text{mol photons m}^{-2} \text{s}^{-1} \), not significantly different from those of oat parental lines (Fig. 4). Together, these data show that the light and CO\(_2\) responses in leaves from OMAs were typical of C\(_3\) plants.

Several Maize Transcripts Associated with C\(_4\) Photosynthesis Are Detectable in OMAs

RNA blotting has shown that the maize genes encoding PPDK and PEPC are expressed in OMA lines 6 and 9, and the proteins that they encode were detectable by immunoblotting (Kowles et al., 2008). To investigate the extent to which other genes important for the biochemistry of C\(_4\) photosynthesis are expressed in maize, we undertook RNA blotting and then quantitative PCR for eight maize transcripts required for the core C\(_4\) cycle. This showed that in addition to \(ZmPPDK\) and \(ZmPEPC\), the maize 2-oxoglutarate/malate transporter (\(OMT1\)) was expressed in oat. So, three of the eight genes associated with the C\(_4\) biochemical pathway generated transcripts of the correct size in oat, indicating that these genes are expressed and their transcripts are correctly spliced (Fig. 5). To quantify the extent to which \(ZmOMT\), \(ZmPPDK\), and \(ZmPEPC\) were expressed in the oat background, mRNA levels were measured by real-time quantitative (RTq) PCR. For all genes, mRNA expression in OMA leaves was detectable but low compared with that in maize leaves (Supplemental Fig. S3), indicating that although transcripts of the correct size were generated, levels of expression in oat were low compared with those in maize.

Thylakoid Stacking in Chloroplasts of Oat PBS Cells Is Unchanged by Individual Maize Chromosomes

C\(_4\) plants that belong to the NADP-ME subtype often show reduced granal stacking in PBS chloroplasts. In order to determine whether the OMA lines showing either an accumulation of C\(_4\) transcripts or changes to PBS cell development or carboxylation efficiency also
possessed alterations to chloroplast ultrastructure, OMA leaves were examined using TEM. The number of thylakoid stacks per granum in M and PBS chloroplasts in lines 3.01, 6.01, 6.32, 9.01, 9.41, and 10.23 was quantified and compared with oat parental lines (starter and sun II) and maize (B73). Chloroplast morphology was normal in OMA lines (Supplemental Fig. S4). The level of thylakoid stacking in PBS cells was variable in the OMA lines but was not significantly different from that in oat parental lines (Fig. 6). This was also true of M cells, with similar numbers of thylakoids in the grana of both M and PBS cells of OMA lines, in contrast to the dimorphic chloroplast arrangement observed in maize (Fig. 6). This demonstrated that changes to chloroplast and PBS development are uncoupled in line 3.01 and that the reduction in carboxylation efficiency in line 9.41 was unlikely unrelated to granal structure (Fig. 6). Maize also has higher numbers of plasmodesmata between M and PBS cells; however, examination lines of 3.01, 6.01, 6.32, 9.01, 9.41, and 10.23 using TEM showed no evidence of this (Supplemental Fig. S4).

DISCUSSION

Expression of Maize Genes Important for C₄ Photosynthesis in C₃ Oat

The extent to which genes from C₄ species are recognized and expressed in C₃ species is fundamental to our understanding of gene regulation and its conservation between species. It is also important to establish which maize genes are expressed in C₃ leaves to inform attempts being made to introduce characteristics of the C₄ pathway into crops such as rice (Matsuoka et al., 2001; Hibberd et al., 2008; Kajala et al., 2011). Our data extend those of Kowles et al. (2008) to show that, in addition to the maize PPDK and PEPC genes, OMT1 is expressed. For all three genes, the size of the mRNA indicates correct splicing in oat. In the case of PEPC and PPDK, this is consistent with previous work that indicated that full-length maize PEPC and PPDK genes driven by their native promoters are expressed and generate transcripts of the correct size in transgenic rice leaves (Ku et al., 1999; Fukayama et al., 2001; Taniguchi et al., 2008). The most parsimonious explanation is that the same transacting factors recognize maize PEPC and PPDK in these two C₃ grasses. However, it should be noted that transcript levels of PPDK, PEPC, and OMT were lower in each OMA line than in maize and, in the case of PEPC, lower than the endogenous oat PEPC. This suggests either that enhancer elements associated with the maize gene were not recognized in oat or that transcript stability was reduced. Consistent with our findings, Kowles et al. (2008) showed that the activity of PEPC was higher in OMAs than in wild-type oat but significantly lower than that in maize.

Transcripts from a number of genes, such as CARBONIC ANHYDRASE2 (CA2), DICARBOXYLIC TRANSPORTER1 (DCT1) and DCT2, MALATE DEHYRODENASE6 (MDH6), and ME1, important for C₄ photosynthesis in maize were not detectable by RNA blotting in the OMA lines. For example, we were not able to detect significant expression of the maize NADP-ME gene. This contrasts with work on the
maize NADP-ME rice, in which the promoter of the maize NADP-ME gene was able to generate expression of the uidA reporter in rice (Nomura et al., 2005). The difference between the behavior of the maize NADP-ME gene in these C3 crops is interesting and could be due to differences in transacting factors or, alternatively, to large-scale silencing of maize genes in the OMA lines. If large-scale silencing of gene expression occurs in these OMA lines, it may well be possible for these five maize genes important for the C4 cycle to be expressed successfully in oat.

Consistent with the lack of expression of many of the C4 genes in oat, including those encoding transporter proteins important for flux across organelle membranes (Weber and von Caemmerer, 2010), we did not detect substantial alterations to the characteristics of gas exchange of the OMA leaves compared with controls. We did detect reductions in carboxylation efficiency, maximal photosynthetic rate, and stature in OMA lines containing chromosome 9 were stunted; however, we detected no consistent reduction in the rate of photosynthesis, and this may be due to the relatively low accumulation of maize PEPC in these plants.

The Plasticity of Oat Leaf Anatomy

Assessing the impact of individual maize chromosomes on oat allowed us to investigate the plasticity of oat leaves, specifically whether elements of Kranz anatomy can be introduced into them. None of the maize chromosomal introductions showed wholesale changes to leaf anatomy consistent with C4 photosynthesis, but three lines showed reductions in vein spacing, a key trait associated with C4 photosynthesis. In these three lines, IVDs were not as low as those in maize, nor did the cellular arrangement generate the repeated units of M-PBS-V-PBS-M cells typical of maize leaves. However, these data show that vein spacing can be altered in a C3 leaf. Interestingly, only one of two lines with maize chromosome 9 (OMA 9.41) showed reduced IVD compared with the wild type. These data show that maize chromosome 9 is capable of decreasing IVD in oat, but this appears to be dependent on the genomic context. Although maize chromosome 9 is common to both oat recipient lines, they differ in their parental backgrounds (Supplemental Table S1), so it is possible that the loci on maize chromosome 9 encoding this trait are differentially silenced depending on their interaction with the oat genome or that they interact differently with specific oat alleles. Interestingly, two of the lines showing reduced vein spacing are derived from crosses involving oat sun II as the recipient, implying that the genetic background of sun II may be more receptive to alterations in vein spacing. The difficulty of generating stable double and triple OMA lines (Kynast et al., 2001) means that testing the additive effects of chromosomes 1, 5, and 9 is not trivial.

To date, mutant screens have failed to identify genes that control the establishment of Kranz anatomy in leaves of C4 plants. Maize mutants with differential effects on PBS and M cells are mostly defective in the accumulation of photosynthesis proteins rather than Kranz anatomy per se (Langdale and Kidner, 1994; Smith et al., 1998), and mutants such as tangled1 with ectopic PBS cells are associated with pleiotropic effects (Smith et al., 2001) and, therefore, are unlikely to be informative in dissecting the molecular basis of Kranz anatomy. Our results indicate that there are loci that are able to regulate PBS size and cell wall lipid deposition in oat that are located on maize chromosome 3. PBS chloroplast number and development were unchanged in this line, suggesting that these traits are not coupled to cell size, and M cells and chloroplasts develop normally in line 3.01, indicating that M development is independent from PBS development. This is consistent with the lack of mutants identified to date with disruptions in M cell development. The transcription factor Golden2, which controls chloroplast development in maize PBS cells (Langdale and Kidner, 1994; Hall et al., 1998), resides on maize chromosome 3; however, the absence of changes to chloroplast morphology in either M or PBS cells in OMA 3.01 suggests that it is not active in oat. The argentina mutant impacts on the speed with which patterns of the C4 mRNAs develop (Langdale et al., 1987), so our analysis of mature OMA leaves is unlikely to have detected phenotypes relevant to determining whether the wild-type ARGENTIA gene product on maize chromosome 9 is active in oat.

Maize PBS cells are also characterized by a suberized cell wall (O’Brien and Carr, 1970); however, suberin was not detected in the OMA 3.01 PBS cell wall, suggesting that the complete maize suberin deposition pathway was not active. Many of the maize homologs of Arabidopsis genes required for suberin synthesis (Li et al., 2010) are located on other maize chromosomes (Supplemental Table S2), supporting this hypothesis. Interestingly, however, ZmLP1 (for lipid transfer protein 1) is located on chromosome 3 and, therefore, is an interesting candidate for the deposition of lipids in the PBS wall observed in OMA 3.01. Expression of ZmLP1...
in maize leaves is high and confined to the developing transition zone (Li et al., 2010), supporting a role in laying down lipids in the walls of developing PBS cells. Similarly, SN2D2, a transcription factor known to be required for secondary cell wall synthesis in Arabidopsis (Zhong et al., 2008; McCarthy et al., 2009), is also expressed in the developing region of the maize leaf (Li et al., 2010) and is located on maize chromosome 3. In order to obtain a more complete picture of the genes involved in lipid deposition in OMA 3.01, analysis of PBS cell phenotypes in OMA radiation hybrid lines (Kynast et al., 2004) containing fragments of maize chromosome 3 in an oat background would permit finer mapping of this trait. Similarly, a comparison of leaf transcriptomes in maize and OMA 3.01 would reveal candidate genes from maize chromosome 3 for controlling lipid deposition, cell wall synthesis, and expansion.

In summary, we have shown that the introduction of maize chromosomes into an oat background can increase PBS cell size and reduce vein spacing in the C3 oat leaf, two key traits of C4 photosynthesis. In addition, some maize genes involved in the core C4 pathway produce transcripts of the correct size in oat. The polyphyletic distribution of the pathway (Sage et al., 2012) combined with evidence of substantial differences in gene expression and spatial control required for C3–C4 transitions (Bräutigam et al., 2011) have led to the proposal that the expression of C4 traits may be governed by a “master switch” (Westhoff and Gowik, 2010). Because we found that there was no coordinated change to gene expression, cell biology, and leaf anatomy, our results provide no evidence for the existence of a global C4 activator. However, a comprehensive assessment of the transcriptional activity of each maize chromosome in the oat background is needed before this can be confirmed. If a two-celled C4 pathway is to be introduced to C3 crop species, changes to leaf anatomy and gene expression will be necessary. Importantly, this study demonstrates that mechanisms underlying the expression of some key maize genes are conserved in oat and that the anatomy of the monocot C3 leaf can be modified.

**Microscopy**

To quantify anatomical arrangements, leaf material was harvested from the widest points (8–10 cm from the leaf base) of fully expanded third leaves and prepared for light microscopy and TEM (Sage and Williams, 1995). Light microscope images of transverse sections were used to quantify the distance between adjacent veins, the number of palisade M cells between veins, the cross-sectional areas of M and PBS tissue, and the ratio of PBS to M tissue (Supplemental Fig. S7). Morphometric analyses were performed using the software packages Q Capture Pro (Q Imaging) and ImageJ (National Institutes of Health). Three TEM images of PBS and M cells from three representative individuals for each line were used to examine the chloroplast number and quantify the number of thylakoid stacks per granum in M and PBS cells. All anatomical measurements were performed in biological triplicate and subjected to statistical analysis by two-tailed Student’s t test to determine their significance relative to oat parental lines (P < 0.01). Nile red (0.01% in 85% lactic acid saturated with chloral hydrate) was used to determine the presence of lipids using epifluorescence (Greenspan et al., 1985; Waduwar et al., 2008).

**Gas-Exchange Measurements**

The net assimilation rates of fully expanded leaves of oat (starter and sun II), maize (B73), and 20 OMA lines under varying internal leaf [CO2] and incident PFD were analyzed in order to screen for characteristic C4 physiological features. Net assimilation rates were measured over a range of external [CO2] using a LI-6400 portable open gas-exchange system (Li-Cor Biosciences) with a photosynthetic leaf cuvette. Leaf temperature was constant at 25°C, and PFD (provided by a Walz FL440 Fiber Illuminator with a 400-F fiber glass connection and heat-filtering projector) was set at 1,500 μmol m−2 s−1. Leaves were allowed to acclimate to saturating light (1,500 μmol m−2 s−1) at ambient CO2 for 30 min. Air flow through the photosynthetic leaf cuvette was varied between 200 and 300 μmol s−1 to ensure that the leaf vapor pressure deficit remained less than 1.5 kPa. After reaching steady state, photosynthetic assimilation rate (A) as a function of internal CO2 concentration (Ci) was measured by varying external [CO2]. CO2 compensation points and carboxylation efficiencies were calculated by regression analysis of A/Ci (in the linear range), with CO2 compensation point equivalent to the s intercept and carboxylation efficiency corresponding to the initial slope gradient. Photosynthetic responses to a range of external light levels at ambient [CO2] were also measured. Leaf temperature and [CO2] were constant at 25°C and 380 μL L−1, respectively, and leaf vapor pressure deficit was maintained below 1.5 kPa. Leaves were equilibrated at 3,000 μmol photons m−2 s−1 prior to curve initiation (Kubiske and Pregitzer, 1996). CO2 and light response measurements are displayed as means of three biological replicates with 1 st.

**RNA Extractions**

Total RNA was extracted from leaves of 3-week-old oat, maize, and OMA plants using TriPure Isolation Reagent (Roche; Chomczynski and Sacchi, 1987). Approximately 200 μg of leaf tissue was ground in liquid nitrogen using a mortar and pestle. Ground tissue was subsequently transferred to an Eppendorf tube containing 1 mL of TriPure Isolation Reagent and mixed by inversion. Chloroform at 200 μL was added after 5 min of incubation at room temperature, and the sample was vortexed (15 s) and centrifuged at 13,000 rpm at 4°C (15 min). The aqueous phase (approximately 600 μL) was transferred to a clean Eppendorf tube containing 500 μL of isopropanol and centrifuged at 13,000 rpm at 4°C for 10 min. The supernatant was discarded, and the pellet was washed with 1 mL of 75% (v/v) ethanol and centrifuged at 7,500 rpm at 4°C for 5 min. Once again, the supernatant was removed, and the RNA pellet was resuspended in 40 μL of RNasearch (Ambion) preheated to 60°C. To aid RNA resuspension, the sample was incubated at 60°C for 10 min, pulse centrifuged, and transferred to a clean Eppendorf tube for subsequent storage at − 80°C.

The quantity of RNA was determined spectrophotometrically, and the quality of RNA was validated via agarose gel electrophoresis. RNA (1 μg and 5 μL of RNA loading buffer [50% (v/v) glycerol, 1 μL EDTA, 0.4% [v/v] bromphenol blue, 0.005% Triton-borate-EDTA buffer, and 48% [v/v] formamide) were heated to 60°C for 10 min and subsequently run on a 1.5% (w/v) agarose gel containing 0.5 μg mL−1 ethidium bromide using 0.5% Triton-borate-EDTA buffer (44.5 μL Tris, 44.5 μL boric acid, and 1 μL EDTA [pH 8.0]).

**Materials and Methods**

**Plant Material**

Oat (Avena sativa; starter and sun II) and OMA line seeds were dehulled and then germinated on saturated filter paper with the application of 200 μL of 75 mg L−1 solution of GA3 in tap water for 2 to 3 d at room temperature before placing them in 3:1 Levington M3 potting compost (Scotts Miracle-Gro): vermiculite treated with intercrop (200 mg L−1; Scotts Miracle-Gro). Plants were grown in day/night temperatures of 20°C/13°C at relative humidity of 75% and under a photosynthetic PFD of 500 photons mol−1 s−1 for 16 h light period. Leaves were defined as fully expanded on ligule emergence (Greenland and Lewis, 1982).

Maize (Zea mays) B73 was grown from seeds imbied overnight in tap water before planting in soil potting mix and grown at 28°C and 50% relative humidity under a photosynthetic PFD of 500 photons μmol m−2 s−1 for a 16-h light period. Samples for DNA or RNA extraction were taken from fully expanded third leaves flash frozen in liquid nitrogen and stored at − 80°C.

Maize Chromosomes Alter Leaf Anatomy in Oat

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RNA Blotting

Total RNA was separated on the basis of molecular weight via gel electrophoresis on a 1.2% formaldehyde agarose gel and transferred onto a nitrocellulose membrane (Genescreen Plus; Perkin-Elmer) by capillary blotting as described previously (Brown et al., 2005). Maize-specific oligonucleotide probes were amplified by PCR from maize cDNA made using an oligo(dT) primer (Invitrogen Life Technologies) and SuperScript II reverse transcriptase (Promega). Primer sequences are recorded in Supplemental Table S3. Radio-labeled probes were prepared by random priming reactions in the presence of \( ^{32}P \)dATP using T4 polynucleotide kinase (New England Biolabs) after Feinberg and Vogelstein (1983). All probes were used in 10 \( \mu \)M working concentrations. Prehybridization was conducted at 37°C in 3 mL of prehybridization buffer (Ambion). Removal of unincorporated nucleotides and verification of \( ^{32}P \) incorporation into probe 5’ PO4 was carried out by centrifugation through a Microspin S-200 HR column (GE Healthcare) prior to their addition to the prehybridization solution. \( ^{32}P \)dATP cDNA probes were added to the solution, and hybridization was carried out overnight at 42°C. Membrane washes to remove any unbound probe and blot hybridization analysis were carried out after Brown et al. (2005). Blots were analyzed using a Typhoon 8600 Variable Mode Imager (GE Healthcare).

RTq-PCR

Total RNA for RTq-PCR was extracted from maize, oat, and OMA leaves using the RNeasy Plant Mini Kit (Qiagen) according to the manufacturer’s instructions. To remove contaminating genomic DNA, samples were treated with 10 units \( \mu \)L \(^{-1} \)RNase-free DNase (Qiagen) for 30 min at 20°C and for 15 min at 65°C. RNA quality was determined spectrophotometrically. One microgram of RNA was reverse transcribed using oligo(dT) primer and SuperScript II (Invitrogen Life Technologies). The total cDNA volume of 20 \( \mu \)L was stored at -20°C overnight. RTq-PCR was carried out using SYBR Green JumpStart Taq Ready Mix (Sigma-Aldrich) and 5-fold dilution of the template end of the coding region. Primers were designed using Primer 3 software (http://frodo.wi.mit.edu/primer3) to have melting temperatures described previously (Brown et al., 2005). Maize-specific oligonucleotide probe primer sequences.

Supplemental Data

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

Supplemental Figure S1. Leaf anatomical characteristics in the three individual lines of maize used.

Supplemental Figure S2. Leaf morphology in OMA lines.

Supplemental Figure S3. mRNA transcripts of ZmPEPC and ZmOMT are expressed at lower levels in OMA leaves than in maize.

Supplemental Figure S4. Chloroplast morphology in OMA lines.

Supplemental Figure S5. Phenotypes resulting from maize chromosomal introduction to the oat genome in 2-week-old OMA plants.

Supplemental Figure S6. Phenotypes resulting from maize chromosomal introduction to the oat genome in 4-week-old OMA plants.

Supplemental Figure S7. Schematic of vein, MS, PBS, and M cells in oat, and illustration of these cells in transverse sections.

Supplemental Table S1. Plant material used for analysis.

Supplemental Table S2. Chromosomal location of putative maize suberin pathway genes.

Supplemental Table S3. Maize-specific oligonucleotide probe primer sequences.

Supplemental Table S4. RTq-PCR primer sequences.

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