

DCT4—A New Member of the Dicarboxylate Transporter Family in C_4 Grasses

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Abstract

Malate transport shuttles atmospheric carbon into the Calvin–Benson cycle during NADP-ME C_4 photosynthesis. Previous characterizations of several plant dicarboxylate transporters (DCT) showed that they efficiently exchange malate across membranes. Here, we identify and characterize a previously unknown member of the *DCT* family, *DCT4*, in *Sorghum bicolor*. We show that *SbDCT4* exchanges malate across membranes and its expression pattern is consistent with a role in malate transport during C_4 photosynthesis. *SbDCT4* is not syntenic to the characterized photosynthetic gene *ZmDCT2*, and an ortholog is not detectable in the maize reference genome. We found that the expression patterns of *DCT* family genes in the leaves of *Zea mays*, and *S. bicolor* varied by cell type. Our results suggest that subfunctionalization, of members of the *DCT* family, for the transport of malate into the bundle sheath plastids, occurred during the process of independent recurrent evolution of C_4 photosynthesis in grasses of the PACMAD clade. We also show that this subfunctionalization is lineage independent. Our results challenge the dogma that key C_4 genes must be orthologues of one another among C_4 species, and shed new light on the evolution of C_4 photosynthesis.

Key words: *DCT4*, new transporter gene, grass evolution, C_4 photosynthesis. .

Significance

Dicarboxylate-transporter-2 (*DCT2*) plays a key role during C_4 photosynthesis in *Zea mays*. Its orthologs are assumed to function the same in related species, as *Z. mays* is the main C_4 reference species. We introduce a new gene, *DCT4*, that assumed the role of *DCT2* in *Sorghum bicolor* and other C_4 grass species. By surveying related C_4 species, we propose that different members of the *DCT* family subfunctionalized for photosynthetic malate transport in the BS cells of C_4 grasses of the PACMAD clade. We suggest that rather than being static, biochemical adaptations continued after the divergence of the PACMAD lineages.

Introduction

Three subtypes of C_4 photosynthesis are generally recognized as defined by the primary decarboxylase in the bundle sheath (BS) cells: Chloroplastic NADP-dependent malic enzyme

(NADP-ME); mitochondrial NAD-dependent malic enzyme (NAD-ME); and cytosolic phosphoenolpyruvate carboxykinase (PEPCK) (Hatch and Slack 1966; Hatch 1971; Rathnam and Edwards 1977). Different plant species may contain various

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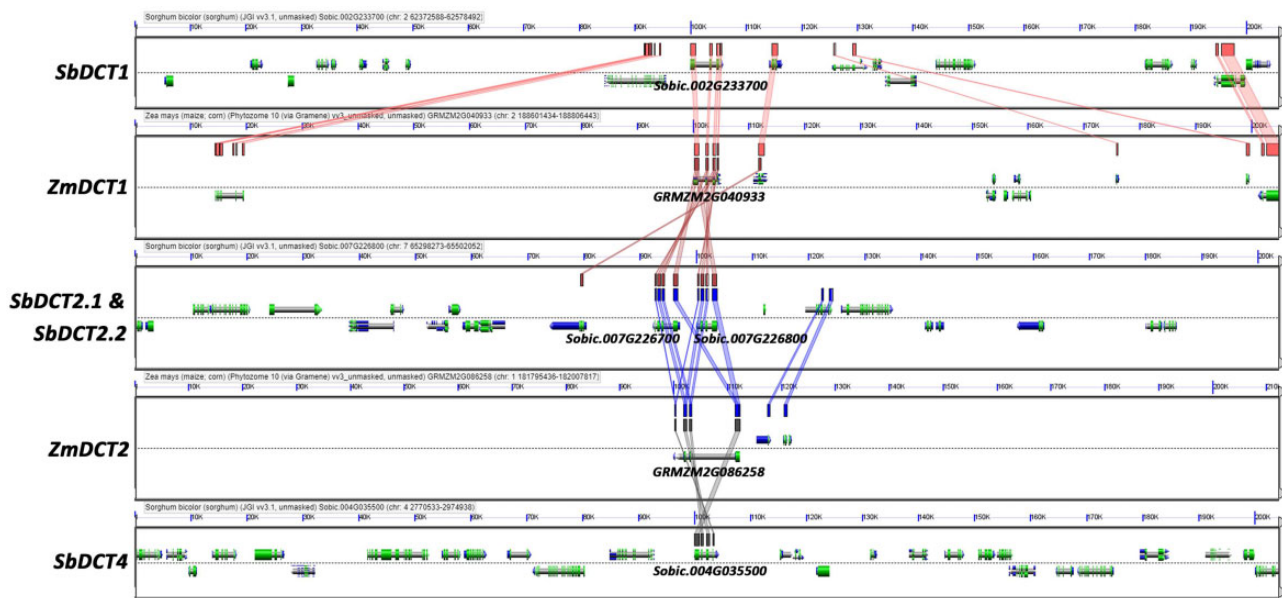


Fig. 1.—CoGe (<https://genomeevolution.org/coge/>) genome viewer screenshots depicting the conservation and genomic contexts of *DCT* genes in *Zea mays* and *Sorghum bicolor*. Colored lines between panels show conserved genes. *SbDCT4* shows high sequence conservation with other *DCT* genes, but is not a syntenic ortholog of *ZmDCT2*, as shown by the lack of conservation in neighboring genes.

combinations of these three subtypes (Hatch 1971; Chapman and Hatch 1979; Furbank 2011; Pick et al. 2011; Wang, Brautigam, et al. 2014). The movement and exchange of malate across membranes, by dicarboxylate transporters (DCTs/ DiTs), plays a significant role during photosynthesis in NADP-ME and NAD-ME C_4 species (Ding et al. 2015). In C_3 plants, DCTs are crucial to nitrate assimilation, such as the GS/GOGAT cycle and photorespiration (Linka and Weber 2010; Kinoshita et al. 2011). Taniguchi et al. characterized several plant DCTs that efficiently exchange malate across membranes (Taniguchi et al. 2002; 2004). The differential expression of C_4 photosynthesis genes in mesophyll (M) and BS cells (John et al. 2014; Tausta et al. 2014; Wang, Czedik-Eysenberg, et al. 2014) suggests that different malate transporters may be needed to move malate out of the chloroplasts of M cells and into the chloroplasts of BS cells. In *Zea mays*, an NADP-ME C_4 grass, dicarboxylate transporter-2 (*ZmDCT2*, GRMZM2G086258) moves malate into the chloroplast of BS cells during C_4 photosynthesis (Weissmann et al. 2016). *ZmDCT2* plays a critical role during C_4 photosynthesis in *Z. mays*, and its absence severely impairs plant growth and development (Weissmann et al. 2016). The role of DCTs in C_4 photosynthesis in other species, however, remains unknown.

Zea mays is the best characterized and functionally annotated C_4 grass species. As such, it is a useful reference for identification of photosynthesis-related genes in poorly characterized C_4 grasses and for resolving orthology (John et al. 2014; Ding et al. 2015; Huang et al. 2017). Microsynteny, the comparison of collinearity among related species, is a reliable approach to determine orthology and predict the function of

a gene (Bennetzen and Freeling 1997; Chen et al. 1997; Tikhonov et al. 1999; Bennetzen 2000; Kumar et al. 2009; Jin et al. 2016). Davidson et al. (2012) showed that syntenic orthologs are likely to have conserved functions and expression patterns across lineages. Here, we identify a new member of the *DCT* family, *DCT4*, which is not syntenic to the photosynthetic gene *ZmDCT2* and is not detected in the maize reference genome. We demonstrate that *S. bicolor DCT4* (*SbDCT4*) efficiently exchanges malate across membranes, consistent with a malate transport role in C_4 photosynthesis. We characterize the diverse expression patterns of *DCT* genes in leaves of multiple grass species. We also propose that subfunctionalization of *DCTs* in grasses of the PACMAD clade (Sanchez-Ken and Clark 2010) occurred during independent recurrent evolution of C_4 photosynthesis.

Results

Identification of *DCT4* in *S. bicolor*

To learn more about C_4 -related DCT in species evolutionarily related to maize, we identified the syntenic ortholog of *ZmDCT2* in *S. bicolor*. Two genes, Sobic.007G226700 and Sobic.007G226800, are present at the predicted syntenic orthologous position on chromosome 7. We refer to them as *SbDCT2.1* and *SbDCT2.2*, respectively (fig. 1). *ZmDCT2* is abundantly expressed (table 1), and its expression is enriched in BS cells of maize leaves (fig. 2) (Li et al. 2010; Tausta et al. 2014; Ding et al. 2015). In contrast, the expression profiles of *SbDCT2.1* and *SbDCT2.2* in *S. bicolor* leaves are low (table 1). *SbDCT2.1* expression is slightly enriched in the M cells

Table 1Genomic Presence or Absence and Whole Leaf TPM Values for Dicarboxylate Transporter Genes in Grass Leaves^a

Species	Presence in Genomic DNA			RNA-seq TPM Values			Photosynthesis	
	<i>DCT1</i>	<i>DCT2</i>	<i>DCT4</i>	<i>DCT1</i>	<i>DCT2</i>	<i>DCT4</i>	<i>OMT1</i>	
<i>Brachypodium distachyon</i>	+	+	–	33	119	N/A	18	C ₃
<i>Oryza sativa</i>	+	+	–	16	11	N/A	101	C ₃
<i>Aristida congesta</i>	+	+	+	393	18	0	871	C ₄ NADP-ME
<i>Eriachne aristidea</i>	+	+	+	1348	89	0	874	C ₄ NADP-ME
<i>Chasmanthium laxum</i>	+	+	–	8	11	N/A	18	C ₃
<i>Danthoniopsis dinteri</i>	+	+	+	0	123	1555	305	C ₄ NADP-ME
<i>Antheophora pubensis</i>	+	+	+	1611	0	0	514	C ₄ NADP-ME
<i>Echinochloa esculenta</i>	+	+	+	1588	0	0	683	C ₄ NADP-ME
<i>Urochloa fusca</i>	+	+	–	50	86	N/A	344	C ₄ PEPCK
<i>Setaria italica</i>	+	+	+	29	41	658	328	C ₄ NADP-ME
<i>Dichantherium oligosanthes</i>	+	+	–	11	13	N/A	13	C ₃
<i>Paspalum vaginatum</i>	+	+	+	591	0	666	354	C ₄ NADP-ME
<i>Arundinella hirta</i>	+	+	+	0	23	481	398	C ₄ NADP-ME
<i>Sorghum bicolor</i>	+	+	+	126	0 ^b	229	483	C ₄ NADP-ME
		+			9 ^c			C ₄ NADP-ME ^d
<i>Zea mays</i>	+	+	–	5	166	N/A	153	C ₄ NADP-ME & PEPCK

^aNote that interspecies comparison is not possible, because expression levels were normalized within each species.^b*SbDCT2.1*.^c*SbDCT2.2*.^dDespite no PEPCK activity detected in *Sorghum*, carbon moves into the BS through aspartate (Chapman and Hatch 1979).

whereas *SbDCT2.2* is enriched in BS cells (fig. 2). We also analyzed the transcript levels of two other *S. bicolor* *DCT*, *SbDCT1* (Sobic.002G233700) and *SbOMT1* (Sobic.008G112300). These genes are the orthologs of the *Z. mays* genes *ZmDCT1* (GRMZM2G040933) and *Zm-oxoglutarate/malate transporter 1* (*ZmOMT1*; GRMZM2G383088), respectively. We found that *SbDCT1* expression, similar to that of *ZmDCT1*, is relatively low (table 1), and only slightly differentially expressed in M cells relative to BS cells (fig. 2). The expression of both *ZmOMT1* and *SbOMT1* is relatively high (table 1), and both are slightly enriched in M cells (fig. 2).

In C₄ species, the expression of many photosynthetic genes is enriched in either BS and M cells (Li et al. 2010; John et al. 2014; Tausta et al. 2014; Weissmann et al. 2016; Rao et al. 2016). In NADP-ME species, two transporters, one within the BS cells and another in M cells, move malate in and out of the chloroplast during C₄ photosynthesis (Brautigam et al. 2008; Weissmann and Brutnell 2012; John et al. 2014; Tausta et al. 2014; Wang, Czedik-Eysenberg, et al. 2014). However, in sorghum leaves, we found only one highly expressed *DCT*, *SbOMT1*, that showed slightly enriched expression in M cells. Therefore, we screened the sorghum genome for additional homologs of known maize *DCTs*. We identified the gene Sobic.004G035500 that showed homology to *ZmDCT1* and *ZmDCT2* but was not syntenic to either gene (fig. 1). *DCT3* is the name of the second transcript of *ZmDCT2* (Taniguchi et al. 2004), so we named this new gene *SbDCT4*. No syntenic ortholog of *SbDCT4* is present in the reference genomes of *Z. mays* or *Oryza sativa*. The absence of syntenic conservation

between *S. bicolor* and *Z. mays* and the lack of direct orthologs in *Z. mays* or C₃ species prevented identification of *DCT4* in a previous bioinformatic screen for C₄ photosynthesis genes (Huang et al. 2017). The expression of *SbDCT4* is moderately abundant (table 1) and strongly enriched in the BS cells of *S. bicolor* leaves (fig. 2).

SbDCT4 is an Efficient Malate Transporter

To verify the ability of *SbDCT4* to transport malate, we cloned coding sequences from the three sorghum *DCT* genes, *SbDCT1*, *SbDCT2*, and *SbDCT4*. We measured the malate transport activities of the recombinant proteins expressed in yeast. *SbDCT4* was an efficient malate transporter (table 2). The *K_m* of *SbDCT4* was similar to that of *SbDCT1*, and the affinity for malate was highest in *SbDCT2* among the three *SbDCTs* (table 2), consistent with the relative malate transport activities reported for maize *DCT1* and *DCT2* (Taniguchi et al. 2004).

Phylogenetic Distribution of *DCT* Genes in Grasses

To understand the relationship of *SbDCT4* to other grass *DCT* genes, we searched the genomes of the grass species *Setaria italica*, *Urochloa fusca*, *Brachypodium distachyon*, and *Dichantherium oligosanthes*. In *S. italica*, an NADP-ME C₄ species, we identified a *DCT*, *Seita.9G375100*, that showed no syntenic orthologous relationship with *DCT* genes in other available grass genomes. Phylogenetic analysis showed that this gene clustered with *SbDCT4* but not with *SbDCT1* and

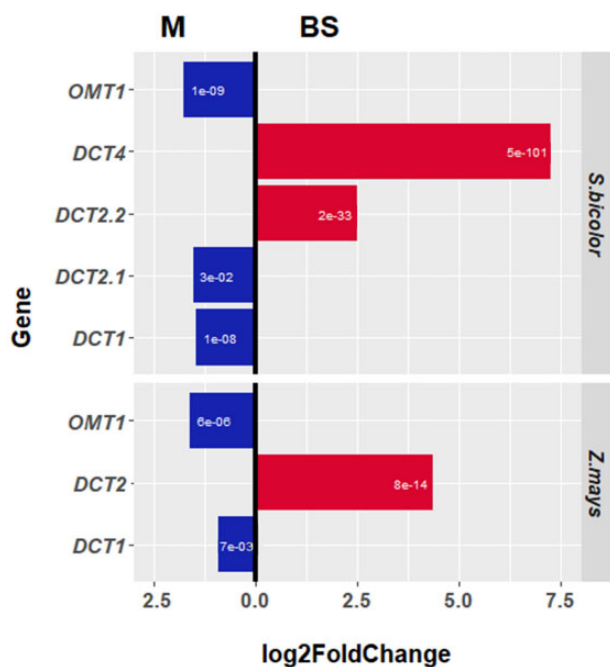


Fig. 2.—Differential expression of malate transporters in *Zea mays*, and *Sorghum bicolor* leaves between the BS and M cells. The genome of *Z. mays* has one copy of *DCT2* and does not contain *DCT4*, and *ZmDCT2* is highly enriched in BS cells. *Sorghum bicolor* has two copies of *DCT2* (*DCT2.1*, and *DCT2.2*) in the syntenic genomic location that are the result of gene duplication. *Sorghum bicolor* also expresses *DCT4*, which is highly enriched in BS cells. Both species express *OMT1* and *DCT1*, which are only slightly enriched in the M cells. Red bars represent enrichment in the BS cells. Blue bars represent enrichment in the M cells. The white numbers inside the bars represent the significance (*P*-value) of the log2(FoldChange).

Table 2

K_m of Malate for Recombinant DCT Proteins^a Demonstrates the Ability of SbDCT4 to Transport Malate Efficiently

K_m (mM)	DCT1	DCT2	DCT4
<i>Sorghum bicolor</i>	1.24 ± 0.14	0.71 ± 0.10	1.13 ± 0.10
<i>Zea mays</i> ^b	1.1 ± 0.1	0.85 ± 0.44	N/A

^aThe values are the means of three independent experiments ± SE.

^bKinetic values from a previous report (Taniguchi et al. 2004).

SbDCT2 (fig. 3). We designated this gene *SiDCT4*. We did not detect orthologs, syntenic or otherwise, in *U. fusca*, a PEPCK C_4 species, or in the two C_3 species. To expand the search for *DCT4* in other grasses currently lacking genome assemblies, we examined leaf-derived transcript assemblies for *Aristida congesta*, *Eriachne aristidea*, *Chasmanthium laxum*, *Danthoniopsis dinteri*, *Anthephora pubensis*, *Echinochloa esculenta*, *Paspalum vaginatum*, and *Arundinella hirta* (Huang P, Mayfield-Jones D, Schnable J, Brutnell T, manuscript in preparation). We then used the predicted coding sequences of the *DCT* genes from available genomes and

from the de novo leaf transcriptome assemblies to generate a phylogenetic tree of the *DCT* family. The resulting phylogeny shows that *DCT4* transcripts form a distinct subclade from the *DCT1* clade (fig. 3). The absence of *DCT4* transcript expression does not rule out the existence of the gene in the genome. We also used polymerase chain reaction (PCR) to survey for *DCT* genes in the genomes of grass species for which whole-genome assemblies were not available. We designed conserved primers (nondegenerate or minimally degenerate) to small regions unique to each of the three *DCT* genes using PrimaClade (Gadberry et al. 2005). We detected *DCT1* and *DCT2* in the genomes of all species tested (table 1, supplemental fig. 1, Supplementary Material online). *DCT4*, however, was detected only in the genomes of NADP-ME C_4 species of the PACMAD clade, excluding *Z. mays* (table 1, supplemental fig. 2, Supplementary Material online).

Expression of Malate Transporter Genes in NADP-ME C_4 Grasses

C_3 species and *U. fusca*, a PEPCK C_4 species, express both *DCT1* and *DCT2* at low levels in leaves (table 1). C_4 NADP-ME species of the PACMAD clade generally express one *DCT* gene in leaves at a high level and also express one or two other *DCT* genes at low levels (table 1). We did not find an apparent lineage-specific pattern for the expression of the predominant *DCT* gene in the NADP-ME species we analyzed. This finding is consistent with random evolutionary processes underlying the subfunctionalization of members of the *DCT* family. Interestingly, *Z. mays* is the only species we examined in which *DCT2* is the predominantly expressed *DCT* gene (table 1). We also examined the expression of the non-*DCT* malate transporter *OMT1* gene in the leaves of grasses (table 1). Interestingly, we found that although *OMT1* expression was generally abundant, there was no consistent pattern of relative expression between the *DCT* and *OMT* genes within the NADP-ME C_4 species (table 1).

Discussion

Evolution of the *DCT* Gene Family in Grasses

We identified *DCT4* as a new member of the *DCT* gene family in the grasses (fig. 1). Our analysis suggests that *DCT4* is present in some C_4 NADP-ME PACMAD grasses. *DCT1* and *DCT2* appear to have originated from a duplication of a single *DCT* gene after the monocot–eudicot split (Taniguchi et al. 2004) and *DCT4* arose from a duplication of *DCT1* at the root of the PACMAD grasses (fig. 3). The expression of *DCT* genes in the grasses that we analyzed exhibited no clear lineage-specific patterns (table 1). Therefore, we propose that different members of the *DCT* family subfunctionalized for photosynthetic malate transport in the BS cells of C_4 grasses of the PACMAD clade.

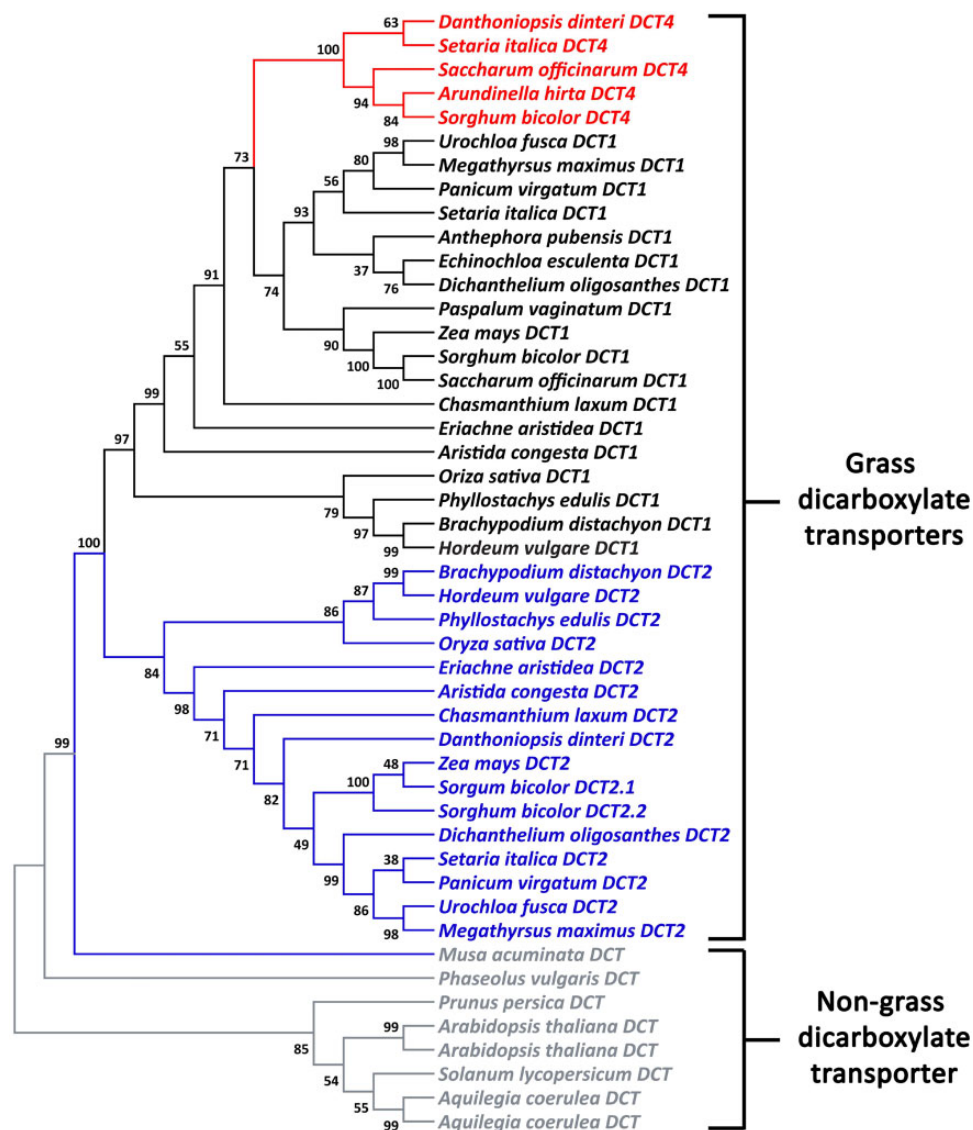


Fig. 3.—A phylogenetic tree of the *DCT* family in the grasses showing that *DCT4* is a subclade of *DCT1*. The *DCT1*, *DCT2*, and *DCT4* gene lineages are black, blue, and red, respectively. The length of the branches represents the evolutionary distance between ancestor to descendent nodes. The numbers represent the confidence level of the specific branch.

This work also challenges the dogma that key C_4 genes must be orthologues of one another, among species, and show that they can be paralogs. This confirms the importance of including syntenic and expression data in assigning orthology across species, and of developing multiple models for C_4 photosynthesis in the grasses. For example, *SiDCT4* was previously misannotated as the ortholog of *ZmDCT2* (John et al. 2014), likely because of the lower expression level of *SiDCT2* (Seita.9G375100) in leaf tissue. The use of different malate transporters, for example, *DCT4* in *S. bicolor* and *S. italica*, or *DCT2* in *Z. mays*, suggests that multiple evolutionary paths resulted in the development of an active C_4 NADP-ME photosynthetic cycle. It is interesting to note that common origins of C_4 photosynthesis are often defined based on the

predominant decarboxylase utilized, thus maize and sorghum are considered to have evolved from a common C_4 ancestor. This analysis suggests that rather than being static, biochemical adaptations continued after the divergence of maize and sorghum lineages. Thus, optimizations of C_4 activities may be continuous as breeding pressures or climate change alters ecological niches of individual species.

Various C_4 Subtype Combinations Have Different Transport Requirements

The variation of expression levels among the different malate transporters within each NADP-ME species (table 1) suggests different transport requirements during C_4 photosynthesis.

This supposition is in agreement with the view that the three subtypes of C_4 photosynthesis are mixed rather than exclusive (Hatch 1971; Chapman and Hatch 1979; Furbank 2011; Pick et al. 2011; Wang, Brautigam, et al. 2014). For example, *Z. mays* utilizes both the NADP-ME (75%) and PEPCK (25%) pathways to fix carbon (Chapman and Hatch 1979; Wingler et al. 1999; Weissmann et al. 2016), and has similar expression levels of *DCT2* and *OMT1* and low expression of *DCT1*. *Sorghum bicolor* moves carbon through both malate and aspartate, although no PEPCK activity was detected in its leaves (Chapman and Hatch 1979). *Sorghum bicolor* has similar expression levels for *DCT4* and *DCT1* and high expression of *OMT1* (table 1). Other grass species may have *DCT* expression ratios that correspond to their unique combination of C_4 subsystems. For example, *OMT1* is highly expressed in *U. fusca*, ~3- to 7-fold higher than *DCT2* or *DCT1*, respectively. *OMT1* transports dicarboxylates, excluding those containing an amino group (Taniguchi et al. 2002, 2004). Thus, in PEPCK C_4 plants, *OMT1* may move oxaloacetate into the mesophyll chloroplast, and 2-oxoglutarate out, to support the high production of aspartate needed to maintain the photosynthetic cycle (Rathnam and Edwards 1977). Interestingly, both *ZmOMT1* and *SbOMT1* are only slightly differentially expressed in the M cells (fig. 2). As the loss of *DCT2* in *Z. mays* prevents movement of malate into the BS chloroplast (Weissmann et al. 2016), *OMT1* cannot be moving malate into the BS chloroplast alongside *DCT2*. But *OMT1* may also have a role in organic acid metabolism in both cell types, such as shuttling reducing equivalents in organelles other than the chloroplast (Pleite et al. 2005).

Conclusions

Our results show that the newly identified member of the *DCT* gene family, *SbDCT4*, is an efficient malate transporter. Based on the expression patterns of malate transporters among the grasses, we suggest that different members of the *DCT* family may have evolved multiple roles in C_4 photosynthesis. Further studies will be needed to verify the subcellular localization of these proteins and to define their specific metabolic functions. Characterizing the various combinations of C_4 photosynthetic subsystems in grasses will facilitate the exploitation of *DCT* genes, through breeding or engineering, to improve the performance of crop plants and increase yield.

Materials and Methods

Identification of *DCT4* Genes in *Sorghum* and Other Grasses

We used QUOTA-ALIGN (Tang et al. 2011) to identify syntenic orthologous regions in grass species with sequenced genomes, following the protocol described in Zhang et al. (2017). To find homologous genes at nonsyntenic locations,

we used two complementary approaches. For species with sequenced genomes, we used LASTZ (Harris 2007) to align the coding sequence of the primary transcript annotated in Phytozome (<https://phytozome-next.jgi.doe.gov>, last accessed January 6, 2021) to the genome assembly. For species without assembled genomes, we used LASTZ to align the coding sequence of the primary transcript from Phytozome to transcript assemblies generated by Trinity (Grabherr et al. 2011).

Measurements of Malate Transport

We cloned each of the three *SbDCT* cDNAs between the promoter and terminator of yeast *GAL2* in the pTV3e vector (Nishizawa et al. 1995). We transformed the plasmids into yeast LBY416 cells and selected transformants on tryptophan-deficient agar plates. We prepared a crude membrane fraction from the selected yeast transformants. We used a freeze-thaw technique to reconstitute liposomes for the measurement of the uptake of [14 C]malate (Taniguchi et al. 2002).

Phylogenetic Analysis of *DCT* Homologs

DCT coding sequences for *Z. mays*, *S. bicolor*, *S. italica*, *B. distachyon*, *O. sativa*, *D. oligoanthes*, and *U. fusca* were from Phytozome (<https://phytozome-next.jgi.doe.gov>, last accessed January 6, 2021). We used BlastN (Altschul et al. 1990) to search *de novo* assembled leaf transcriptomes (Huang P, Mayfield-Jones D, Schnable J, Brutnell T, manuscript in preparation) from the C_4 grass species *A. congesta*, *E. aristidea*, *C. laxum*, *D. dinteri*, *A. pubensis*, *E. esculenta*, *P. vaginatum*, and *A. hirta* with the *DCT* sequences from maize, *Setaria*, and *Sorghum* as queries. We used ProGraphMSA to generate a codon-based sequence alignment (Szalkowski 2012). We used MEGA6 (Tamura et al. 2013), with default parameters and the branch support values based on 1,000 bootstraps, to generate the phylogenetic reconstruction with the maximum likelihood method and based on the nucleotides in the third position of codons (Simmons et al. 2006).

Analysis of Gene Expression for Decarboxylase Transporters in Grasses

For species with published leaf transcriptome profiles (Ouyang et al. 2007; Li et al. 2010; Zhang et al. 2012; Schnable 2014; Wang, Czedik-Eysenberg, et al. 2014; Studer et al. 2016), gene expression levels were calculated and normalized, for each species, as Transcripts Per Million (TPM). For the other species, the normalized TPM values were based on *de novo* transcriptome assemblies (Huang P, Mayfield-Jones D, Schnable J, Brutnell T, manuscript in preparation). The values in table 1 only allow for intraspecies comparisons among the decarboxylase transporters.

Table 3Primers and PCR Conditions for the Amplification of Grass *DCT* Genes

Primer Pair	Forward 5'-3' ^a	Reverse 5'-3' ^a	Cycling Conditions
DCT1	CACCAACGAGGTCATCTGG	AGTAGGTGGCGATDCGGTC	94 °C 3 min, [94 °C 45 s, 58 °C 30 s, 72 °C 1 min] × 30, 72 °C 10 min, 4 °C ∞
DCT2	CVTGGATGTCRAATTGTGTTG	TGGCTTGCAABADATAGTGAA	94 °C 3 min, [94 °C 45 s, 58–52 °C (–0.5 °C/cycle) 30 s, 72 °C 1 min] × 14, [94 °C 45 s, 52 °C 30 s, 72 °C 1 min] × 16, 72 °C 10 min, 4 °C ∞.
DCT4	CTTYGTCAAGTGGCTCGG	GACTTGATGATSGGCAGGA	94 °C 3 min, [94 °C 45 s, 60 °C 30 s, 72 °C 1 min] × 30, 72 °C 10.0 min, 4 °C ∞

^aB = C + G + T, D = A + G + T, R = A + G, S = C + G, V = A + C + G, Y = C + T.

Identification of *DCT4* in Species without Sequenced Genomes

We aligned the coding sequences from each of the *DCT* genes from *Z. mays*, *S. bicolor*, *S. italica*, *U. fusca*, *B. distachyon*, *O. sativa*, *D. oligosanthos*, *A. congesta*, *E. aristidea*, *C. laxum*, *D. dinteri*, *A. pubensis*, *E. esculenta*, *P. vaginatum*, and *A. hirta* using PAL2NAL (Suyama et al. 2006). The resulting multiple sequence alignment enabled the design of non-degenerate or minimally degenerate PCR primers (table 3) using PrimaClade (Gadberry et al. 2005).

Jacob D. Washburn and J. Chris Pires (University of Missouri, Columbia) kindly provided genomic DNA from *A. congesta*, *E. aristidea*, *D. dinteri*, *A. pubensis*, *E. esculenta*, and *A. hirta* (Washburn et al. 2015). We used a CTAB-based method to extract genomic DNA from *C. laxum*, *P. vaginatum*, *Z. mays*, *S. bicolor*, *S. italica*, and *B. distachyon* (Weissmann et al. 2016). *Zea mays* and *B. distachyon* were the negative controls for *DCT4* and the positive controls for *DCT1* and *DCT2*. *Sorghum bicolor* and *S. italica* were the positive controls for *DCT1*, *DCT2*, and *DCT4*.

We conducted amplification of *DCT* genes by PCR using a 25- μ l reaction mix and an ABI 2720 Thermal cycler. The reaction mixture included 2.5 μ l of 10 \times Buffer, 2.5 μ l of 10 μ M solutions of forward and reverse primers, 2 μ l of 2.5 mM dNTP stock, 14 μ l of nuclease-free water, 0.5 μ l of Choice Taq enzyme, and 1 μ l of 100 ng/ μ l DNA. We performed PCR reactions as described in table 3 with 5 μ l of loading dye added to each reaction. Aliquots of 13 μ l were loaded on 3% agarose gels (Invitrogen UltraPure Agarose 1000, 1 \times TAE buffer, Invitrogen SYBR Safe Gel Stain) and electrophoresed for 30 min at 100 volts. We based size estimates on 100 bp and 50 bp DNA markers (GoldBio).

Supplementary Material

Supplementary data are available at *Genome Biology and Evolution* online.

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Data Availability

This work includes no new sequence data.

Literature Cited

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