Characterization of Tocopherol Cyclases from Higher Plants and Cyanobacteria. Evolutionary Implications for Tocopherol Synthesis and Function¹

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Tocopherols are lipophilic antioxidants synthesized exclusively by photosynthetic organisms and collectively constitute vitamin E, an essential nutrient for both humans and animals. Tocopherol cyclase (TC) catalyzes the conversion of various phytyl quinol pathway intermediates to their corresponding tocopherols through the formation of the chromanol ring. Herein, the molecular and biochemical characterization of TCs from Arabidopsis (VTE1 [VITAMIN E 1]), Zea mays (SXD1 [Sucrose Export Deficient 1]) and Synechocystis sp. PCC6803 (slr1737) are described. Mutations in the VTE1, SXD1, or slr1737 genes resulted in both tocopherol deficiency and the accumulation of 2,3-dimethyl-6-phytyl-1,4-benzoquinone (DMPBQ), a TC substrate. Recombinant SXD1 and VTE1 proteins are able to convert DMPBQ to γ-tocopherol in vitro. In addition, expression of maize SXD1 in a Synechocystis sp. PCC6803 slr1737 knockout mutant restored tocopherol synthesis, indicating that TC activity is evolutionarily conserved between plants and cyanobacteria. Sequence analysis identified a highly conserved 30-amino acid C-terminal domain in plant TCs that is absent from cyanobacterial orthologs. That TC activity is evolutionarily conserved between plants and cyanobacteria. Sequence analysis identified a highly conserved 30-amino acid C-terminal domain in plant TCs that is absent from cyanobacterial orthologs.

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bined molecular characteristics allow tocopherols to protect polyunsaturated fatty acids from lipid peroxidation by scavenging lipid peroxyl radicals that propagate lipid peroxidation chain reactions in membranes (Burton et al., 1986; Liebler, 1993). Though direct evidence is lacking, tocopherols are thought to play similar roles in protecting the polyunsaturated fatty acid-rich plastid membrane from lipid peroxidation.

Recent studies in mammalian systems have demonstrated additional biological activities of tocopherols that are independent of their antioxidant functions. The underlying mechanisms for these effects are the modulation of signal transduction pathways by specific tocopherol-binding proteins (Brigelius-Flohe and Traber, 1999; Sen et al., 2000; Chan et al., 2001; Ricciarelli et al., 2001; Yamauchi et al., 2001; Clement et al., 2002; Nobata et al., 2002). Modulation of the protein kinase C signaling cascade and eicosanoid synthesis are two well-characterized examples of the antioxidant-independent effects of tocopherols in mammalian systems (Greenberglevy et al., 1993; Tran et al., 1996; Azzi et al., 2002). Although direct experimental evidence is lacking for antioxidant-independent tocopherol activities in plants, these data raise the possibility that tocopherols may also have roles in plants that extend beyond their proposed antioxidant functions.

Though the functions of tocopherols in plants remain an open question, much has been learned about tocopherol synthesis and the pathway enzymes during the past 5 years (Norris et al., 1998; Shintani and DellaPenna, 1998; Collakova and DellaPenna, 2001; Schledz et al., 2001; Porfirova et al., 2002; Savidge et al., 2002; Shintani et al., 2002). Tocopherol synthesis draws substrates from two separate metabolic pathways, aromatic amino acid metabolism and isoprenoid synthesis. Homogentic acid, an intermediate in aromatic amino acid degradation and the head group of tocopherols, is produced from \( \text{p-hydroxyphenylpyruvate} \) by the cytosolic enzyme \( \text{p-hydroxyphenylpyruvate dioxygenase} \) (HPPD; Garcia et al., 1997, 1999; Norris et al., 1998; Dahnhardt et al., 2002). The isoprenoid-derived phytol tail of tocopherol is a product of the plastid-localized 1-deoxy-D-xylulose-5-phosphate pathway (Eisenreich et al., 1998; Lichtenthaler, 1998). The remaining steps in tocopherol synthesis occur within the inner envelope of the chloroplast and include a phytol transferase, two different methyltransferases, and a ring-producing enzyme, the tocopherol cyclase (TC; Soll et al., 1980, 1985; Arango and Heise, 1988a).

The TC adds a second oxygen-containing ring at the junction between the aromatic head group and phytol tail to create a two-ring structure known as a chromanol ring (Fig. 1), which is essential for resonance stabilization of tocopheroxyl radicals after single-electron transfer. Previous work has characterized TC activity in chloroplasts and chromoplasts of higher plants and in cyanobacteria (Soll, 1979; Soll and Schultz, 1980; Soll et al., 1987; Stocker et al., 1993, 1994, 1996; Arango and Heise, 1998b). The primary substrate of the TC is reduced (quinol form) 2,3-dimethyl-6-phytyl-1,4-benzoquinone (MPBQ), which is converted to \( \gamma \)-tocopherol by TC (Soll, 1979; Soll and Schultz, 1980; Soll et al., 1985). However, the enzyme characterized from \( \text{Anabaena} \) sp. PCC7120 has also been shown to cyclize other 6-prenyl-1,4-benzoquinol substrates in vitro (Stocker et al., 1996). In this report, we describe the isolation and functional characterization of TCs from Arabidopsis, maize (\( \text{Zea mays} \)), and \( \text{Synechocystis} \) sp. PCC6803 and discuss the evolutionary implications of tocopherol cyclization for both tocopherol synthesis and function. The identification and characterization of TC from Arabidopsis were recently reported by Porfirova et al. (2002).

**RESULTS**

**Isolation and Characterization of vte1 Mutants**

To further understand the tocopherol pathway in higher plants, an HPLC-based screen of Arabidopsis
leaf tissue was developed to isolate mutants with tocopherol profiles that differ from wild type. Arabidopsis leaves accumulate approximately 10 ng α-tocopherol mg⁻¹ fresh weight and 0.2 ng γ-tocopherol mg⁻¹ fresh weight under standard growth conditions (see “Materials and Methods”). Numerous mutants were identified from an ethyl methanesulfonate (EMS)-mutagenized population, including two mutants that were devoid of tocopherols in leaf tissue (Fig. 2A). Genetic complementation tests confirmed that the mutants were allelic (data not shown). The mutants were designated vte1-1 and vte1-2 (vitamin e).

The visible phenotypes of both vte1 mutants did not significantly differ from wild type when grown under normal laboratory conditions (see “Materials and Methods”). Although several possibilities could result in a tocopherol-deficient phenotype, the two most likely are a loss of HPT activity or a loss of TC activity (Fig. 1). Assuming no genetic redundancy, a mutation disrupting either gene would result in a tocopherol-deficient phenotype, but the two classes of mutations should be readily distinguishable by the intermediates that accumulate. A defect in the TC should result in the accumulation of the DMPBQ, whereas a mutation in HPT would not accumulate tocopherol pathway prenyl quinone intermediates (Fig. 1). To understand the biochemical basis of vte1-1 and vte1-2, prenyl quinones were isolated from each mutant and analyzed by HPLC (Fig. 3A). A novel peak with a retention time and spectrum consistent with the prenyl quinone DMPBQ (Hutson and Threlfall, 1980; Marshall et al., 1985; Johnson et al., 2000) was observed in the vte1-1 and vte1-2 mutants but not in wild type (Fig. 3A). This compound was purified by HPLC and a mass of 415 D, the mass of DMPBQ, was determined by fast atom bombardment mass spectroscopy (data not shown). These combined characteristics indicate that the novel compound that accumulates in vte1-1 and vte1-2 mutants is DMPBQ, the substrate of the TC.

In addition to green tissues, seeds also contain tocopherols, but instead of α-tocopherol predominating as in leaves, γ-tocopherol accumulates due to low γ-TMT activity (Shintani and DellaPenna, 1998). Lipids were extracted from the seeds of wild-type and vte1 mutants, and the tocopherols were analyzed by HPLC (Fig. 4). Tocopherols were absent in vte1-1 seed, but vte1-2 seeds contained approximately 25% of the tocopherol level in wild type (84.9 ± 0.8 versus 322.5 ± 7.4 ng total tocopherols mg⁻¹ seed in vte1-2 and wild type, respectively), suggesting vte1-2 is a weaker allele than vte1-1.

A map-based cloning approach was undertaken to isolate the gene encoding the TC from Arabidopsis. vte1-1 was crossed to Landsberg erecta, and 1,100 individuals from an F₃ population were used to map the VTE1 locus to a 140-kb interval on the bottom of chromosome 4. Analysis of the genes within this interval identified At4g32770, encoding an unknown protein of 488 amino acids that contains a putative N-terminal chloroplast transit peptide of 68 amino acids. Previously, At4g32770 and the Synechocystis sp. PCC6803 protein slr1737 were identified as homologs of SXD1 (Suc Export Defecient 1) from Maize (Provencher et al., 2001). slr1737 is a protein of unknown function and is in the same operon as slr1736, which encodes HPT (Collakova and DellaPenna, 2001; Schledz et al., 2001; Savidge et al., 2002), another enzyme of the tocopherol biosynthetic pathway (Fig. 1). At4g32770 was fully sequenced in both mutants, and each was found to contain a nonsense mutation. The vte1-1 mutation creates a premature

![Figure 2. HPLC analysis of tocopherols in wild-type and mutant Arabidopsis, maize, and Synechocystis sp. PCC6803.](https://www.plantphysiol.org/)
stop codon at amino acid 237 (Trp to stop), whereas the vte1-2 mutation creates a premature stop codon at amino acid 465 (Gln to stop). Both mutations are base transitions, G to A and C to T, respectively, which is consistent with the mutagenic properties of EMS. The identification of At4g32700 as the Arabidopsis TC concurs with a recent report by Porfirova et al. (2002).

Sequence Analysis and Evolutionary Origin

BLAST searches revealed that SXD1, VTE1, and slr1737 share a high degree of amino acid sequence similarity (Table I) with other proteins in the nonredundant GenBank database: three proteins of unknown function in the cyanobacteria Anabaena sp. PCC7120, N. punctiforme, and Synechococcus sp. PCC7002. SXD1 is a chloroplast-targeted protein of unknown function that had been identified previously based on a mutation causing a defect in symplastic photosynthate transport near the site of phloem loading within the minor veins of maize leaves (Russin et al., 1996; Provencher et al., 2001). When the SXD1 and VTE1 chloroplast transit peptides are removed, SXD1, VTE1, and the four cyanobacterial proteins share long stretches of amino acid identity.

The four cyanobacterial proteins are assumed to be orthologs of VTE1 because the cyanobacterial genomes each contain obvious orthologs of the four other known genes of the tocopherol pathway: HPPD, HPT, MPBQ methyltransferase, and γ-TMT. VTE1 and the cyanobacterial orthologs exist as single genes within their respective sequenced genomes. There are several other cyanobacteria whose genomes also have been sequenced: Prochlorococcus marina...
rinus MED4, P. marinus MIT9313, Synechococcus sp. PCC7002 WH8102, and Thermosynechococcus elongatus BP-1. The genomes of these organisms lack obvious VTE1 orthologs and obvious orthologs for HPPD, HPT, and γ-TMT (refer to pathway in Fig. 1). Thus, it appears likely that only a subgroup of cyanobacteria have evolved the ability to synthesize tocopherols.

VTE1, SXD1, and the four cyanobacterial orthologs lack any previously described protein motifs. There are numerous plant expressed sequence tags (ESTs) in the public database that share high similarity with VTE1 and SXD1, and full-length sequences of the M. truncatula and barley VTE1 orthologs were obtained from EST assemblies. These four representative plant sequences are more conserved than the four cyanobacterial sequences (Table I). Sequence alignment of the plant and the cyanobacterial protein sequences identified a highly conserved 30-amino acid carboxyl domain in the plant VTE1 orthologs (starting at Thr-458 of Arabidopsis VTE1) that is absent from the cyanobacteria proteins (Fig. 5). The last five amino acids of this carboxyl domain (KPPGL) are invariant among the plants represented, which include the bryophyte P. patens, monocots, and dicots. With the exception of vascular and nonvascular VTE1 orthologs, this 30-amino acid domain was not found in other proteins in the nonredundant database. Interestingly, the C. reinhardtii VTE1 ortholog has a shortened version of the carboxyl domain (Fig. 5) and lacks the last 12 amino acids (starting at Leu-477 of Arabidopsis VTE1), including the invariant KPPGL motif. The vte1-2 mutation causes premature termination of VTE1 and deletion of 24 amino acids of the conserved carboxyl domain.

**TC Function in Arabidopsis, Maize, and Synechocystis sp. PCC6803**

To confirm that the VTE1 orthologs are required for tocopherol synthesis in plants other than Arabidopsis, lipids were isolated from leaves of the sxd1 mutant and analyzed for tocopherols by HPLC. As with vte1-1 and vte1-2, leaves of the sxd1 mutant lack tocopherols, whereas wild-type maize leaves contain both α- and γ-tocopherols (Fig. 2B). Prenyl quinones from leaves of wild-type maize and the sxd1 mutant were also analyzed by HPLC. This analysis indicated that like the vte1 mutants, sxd1 contained a prenyl quinone that was absent from wild type (Fig. 3B) with an absorbance spectrum and retention time identical to the DMPBQ that accumulated in vte1 mutants.

To show that this gene family has an identical function in cyanobacteria and plants, an insertional mutant, slr1737, was created in the slr1737 open reading frame (ORF) of *Synechocystis* sp. PCC6803.

### Table I Pairwise comparisons of VTE1 orthologs from plants and cyanobacteria

Pair-wise comparisons were performed using ClustalW and are expressed as percentage amino acid similarity. The predicted mature plant protein sequences were used for alignments. Anabaena, *Anabaena* sp. PCC7120 (all0245); Nostoc, *Nostoc punctiforme* (506-74); Synecho, *Synechococcus* sp. PCC7002.

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**Figure 5.** Alignment of the carboxy termini of TC orthologs from plants and cyanobacteria. The asterisk above the At4g32770 protein sequence denotes the position of the vte1-2 mutation. Anabaena, *Anabaena* sp. PCC7120 (all0245); Chlamy, *Chlamydomonas reinhardtii*; Medicago, *M. truncatula*; Nostoc, *N. punctiforme* (506-74); Physcomitrella, *Physcomitrella patens*; Synecho, *Synechococcus* sp. PCC7002. The *P. patens*, rice (*Oryza sativa*), and wheat (*Triticum aestivum*) sequences are partial sequences obtained from ESTs. Dark shading, Amino acid identity; light shading, Amino acid similarity. The threshold for amino acid consensus identity or similarity is 51%.
Figure 6. TC activity of proteins expressed in E. coli. E. coli cell lysates from cells overexpressing the empty pET vector or pET engineered to express TC proteins from Arabidopsis, maize, and Synechocystis sp. PCC6803 were incubated with radiolabeled 2,3-methyl-6-phytyl-1,4-benzoquinol (3 methyl $^{14}$C) for 4 h as described in “Materials and Methods.” Total lipids were extracted, separated by TLC, and radiolabeled products were detected by phosphor imager analysis. Products were identified by comigration with standards. The $^{14}$C incorporation into γ-tocopherol was quantified densitometrically and expressed as pixels per microgram of total protein.

that SXD1 is able to functionally complement Δslr1737. Hence, these cyanobacterial and plant genes have not only high sequence similarity but also functional equivalency, suggesting a common evolutionary ancestry.

To determine the activity of the VTE1 protein and its maize and cyanobacterial orthologs, we expressed VTE1, SXD1, and slr1737 in Escherichia coli using the pET expression system. Lysates from E. coli expressing either VTE1 or SXD1 were able to convert $^{14}$C2,3-dimethyl-6-phytyl-1,4-benzoquinol into γ-tocopherol (Fig. 6). This result conclusively demonstrates that both genes encode an enzyme with TC activity. Activity was not observed with the slr1737 protein expressed in E. coli for reasons that are unknown.

Carbohydrate Assimilation in vte1

Although SXD1 and VTE1 have similar enzymatic activities (Fig. 6) and primary biochemical phenotypes (tocopherol deficiency and DMPBQ accumulation, Figs. 2 and 3), sxd1 was initially isolated because of a secondary phenotype, a Suc transport defect (Russin et al., 1996; Provencher et al., 2001). To determine whether vte1 caused a similar Suc transport defect, Glc, Suc, and starch levels were analyzed in mature leaves from 4-week-old vte1-1 plants. The leaves were sampled at the end and at the beginning of the photoperiod. There were no significant differences between wild-type and vte1-1 leaves for Glc, Suc, and starch at the beginning or end of the photoperiod (Table II). Thus, unlike the sxd1 mutant, carbohydrate metabolism in mature leaves appears unaffected by the vte1 mutation. The molecular and biochemical similarities of the VTE1 and SXD1 proteins suggest the difference in sxd1 and vte1 carbohydrate metabolism phenotypes in maize and Arabidopsis reflects additional roles of tocopherols or the tocopherol pathway beyond antioxidant chemistry, rather than simply a difference in the enzymatic activity of the SXD1 and VTE1 proteins.

DISCUSSION

In this report, we have shown that the three proteins, VTE1, SXD1, and slr737 from a dicot, monocot,
and cyanobacterium, respectively, function as TCs. Mutations in the TC gene from each organism result in identical primary biochemical phenotypes, a block in tocopherol synthesis, and accumulation of DMPBQ, the endogenous substrate for the TC. In addition, the SXD1 and VTE1 proteins expressed in E. coli were able to convert DMPBQ to γ-tocopherol. Finally, expression of maize SXD1 was sufficient to complement the tocopherol-deficient phenotype of the Synechocystis sp. PCC6803 slr1737 deletion mutant (Δslr1737). This result demonstrates that slr1737 and SXD1 are functionally equivalent and that the biochemical activity of TCs has been evolutionarily conserved between plants and cyanobacteria. Our finding that At4g32770 encodes a functional TC in Arabidopsis concurs with a recent report by Porfirova et al. (2002).

Sequence Analysis and Evolutionary Implications of the TC Family

The TCs (VTE1, SXD1, and slr1737) share significant amino acid similarity with each other and define an evolutionarily conserved gene family that includes putative orthologs in a large number of other plants and cyanobacteria. VTE1 orthologs were not identified in databases of fungal, animal, or non-photosynthetic bacterial species, none of which are known to produce tocopherols. Full-length sequences of two additional VTE1 orthologs from plants (barley and M. truncatula) and three from cyanobacteria (N. punctiforme, Anabaena sp. PCC7120, and Synechococcus sp. PCC7002) were identified in the public databases. Although all VTE1 proteins share a high degree of amino acid similarity, they are devoid of any previously described protein motifs, with the exception of ubiquitous phosphorylation and myristolation motifs. All VTE1 orthologs are hydrophobic proteins with low pIs and a high number of conserved Trp residues (Provencher et al., 2001). These characteristics are consistent with the TC activity characterized in Anabaena variabilis being membrane associated (Stocker et al., 1993, 1994, 1996).

Although plant and cyanobacterial TCs exhibit a high degree of protein sequence similarity, plant orthologs have additional N- and C-terminal domains that are absent in the four cyanobacterial TCs. The N-terminal domains of plant VTE1 orthologs are poorly conserved and are predicted to encode chloroplast transit peptides that would target each protein to the chloroplast. The N-terminal sequence of SXD1 has been demonstrated experimentally to be required for import into plastids (Provencher et al., 2001). HPT and γ-TMT, two other tocopherol biosynthetic enzymes, are also predicted to be chloroplast targeted (Shintani and DellaPenna, 1998; Collakova and DellaPenna, 2001; Schledz et al., 2001; Savidge et al., 2002), and chloroplast localization of the TC is consistent with the reported localization of TC activity and tocopherol synthesis in plastids (Soll et al., 1985; Arango and Heise, 1998a).

In contrast to the N-terminal domain of plant TCs, the 30-amino acid C-terminal domain is highly conserved between angiosperms and the moss P. patens. This evolutionary conservation suggests an important function for this domain in tocopherol synthesis in plants, whereas the absence of the sequence from the four cyanobacterial VTE1 orthologs suggests that the domain is not an absolute requirement for TC enzymatic activity per se. The restriction of this C-terminal domain to vascular and nonvascular plants suggests it arose relatively recently in progenitors of land plants rather than in the endosymbiotic cyanobacteria that gave rise to plastids (Goksoy, 1967). The shortened C-terminal domain present in the C. reinhardtii TC further suggests evolution took place in an ancestor common to C. reinhardtii and plants (i.e. before the split of Chlorophyta and Chlorophyta; Karol et al., 2001). The vte1-2 mutation causes deletion of the majority of the conserved C-terminal domain, and tocopherols fail to accumulate in vte1-2 leaf tissue but reach 25% of wild-type levels in seeds, indicating the truncated protein retains at least partial activity in vivo. The vte1-2 phenotype suggests the C-terminal domain plays a more significant role in TC activity/function in leaf chloroplasts than in the plastids of seeds. The complementation of Δslr1737 by SXD1 suggests that the presence of the C-terminal domain does not affect TC activity and function in cyanobacteria. The relevance of this highly conserved carboxyl domain for TC function in plants requires further investigation.

The sxd1 and vte1 Phenotypes

A surprising phenotype of the sxd1 mutants is a block in Suc export from leaves and an accumulation of anthocyanins and starch in leaf blades (Russin et al., 1996). This pleiotropic phenotype results from aberrant plasmodesmata at the interface between the bundle sheath cells and the vascular parenchyma cells surrounding the minor veins. These defective plasmodesmata block symplastic transport of Suc to the phloem and, hence, cause the Suc export defect. Unlike sxd1, vte1-1 and vte1-2 do not accumulate anthocyanins in leaves, do not accumulate starch in cells surrounding the leaf veins, and do not have stunted growth (data not shown). In addition, the similar Suc, Glc, and starch levels in mature leaves of vte1-1 and wild type (Table II) indicate that a functional Suc export pathway is present in vte1. However, this most obvious difference in phenotype between sxd1 and vte1 is not entirely unexpected considering the anatomical and physiological differences between C3 and C4 plants. Maize bundle sheath cells contain chloroplasts that differ morphologically and physiologically from the chloroplasts of mesophyll cells (Evert et al., 1977a; Fahn, 1990).
Maize bundle sheath plastids have few grana, little PSII activity, high NADP-malate decarboxylase activity, and accumulate starch (Evert et al., 1977a, 1977b, 1978; Fahn, 1990). Arabidopsis lacks a physiologically equivalent cell type to the C4 bundle sheath cell. Chloroplasts within the analogous cells surrounding the minor veins in Arabidopsis do not differ significantly from the chloroplasts of mesophyll cells (Hari-tatos et al., 2000). Mutants of the Arabidopsis SUC2 gene, which encodes a Suc-H+ symporter required for apoplastic phloem loading, share striking similarities to sxd1 mutants (Gottwald et al., 2000). suc2 mutants accumulate anthocyanins and starch in their cotyledons, have stunted growth, and are seedling lethal in soil (Gottwald et al., 2000). Thus, the absence of the sxd1 Suc export phenotype in vte1 suggests that disruption of VTE1 in Arabidopsis either does not affect Suc export, most likely because of the fundamentally different mechanisms of Suc export in maize and Arabidopsis, or that the effect is too small to be observable as an analogous whole-plant phenotype in Arabidopsis.

In sxd1 mutants, the link between the production of aberrant plasmodesmata in the BS parenchyma cells and the defect in symplastic transport of Suc was straightforward and easy to rationalize (Provencher et al., 2001). However, the mechanistic link between the disruption of a gene encoding a chloroplast protein of unknown function (SXD1) and defective Suc transport was not so obvious. Provencher et al. (2001) raised the possibility that the sxd1 mutation exerts its pleiotropic phenotype by disrupting or altering a signal from the chloroplast to the nucleus. The nature of this signal was unknown, and cloning of the SXD1 locus did not provide further insight. Although the precise nature of this signal remains unclear, the demonstration that SXD1 encodes a TC and determination of the primary sxd1, vte1, and Δs1r1737 biochemical phenotypes now greatly limits the possibilities.

The absence of tocopherols in vte1 and Δs1r1737 does not cause a pleiotropic phenotype analogous to sxd1. vte1 mutants are indistinguishable from wild type under normal growth conditions, and Δs1r1737 and Δls1r737 mutants grow at rates identical to wild type (Collakova and DellaPenna, 2001). These data suggest that the sxd1 phenotype is not simply due to the absence of tocopherols as lipophilic antioxidants because similar effects would be observed in the vte1, Δls1r1737, and Δls1r1736 mutants, which also lack tocopherols. In addition, the DMPBQ that accumulates in vte1, sxd1, and Δls1r1737 mutants can still act as an antioxidant by donating a pair of electrons and then being recycled by reduction back to the quinol form of DMPBQ (Kruk et al., 1994; Kruk and Strzalka, 1995; Liebler and Burr, 2000). Thus, even in the absence of tocopherols, the sxd1 mutant is not entirely deficient in membrane-associated antioxidants.

Influencing membrane fluidity is another potential function of tocopherols, and this could be related to the plasmodesmatal defect in sxd1 mutants. However, the consensus from cell fractionation studies is that tocopherols are localized exclusively in plastid membranes (Lichtenhaler et al., 1981; Fryer, 1992; Kruk and Strzalka, 1995), and although it cannot be excluded, it is unlikely that tocopherols or DMPBQ would be present in plasma membranes to directly impact plasmodesmatal development. However, the bundle sheath cell is a specialized cell type, and direct physical connections between plasmodesmata and chloroplasts through the endoplasmic reticulum have been reported in maize bundle sheath cells (Evert et al., 1977a, 1977b). This potential association between the chloroplast and plasmodesmata would provide a means for tocopherols or prenyl quinones to impact membrane fluidity at the plasma membrane. Still, it is difficult to envision a mechanism whereby membrane fluidity would alter plasmodesmata development. If diminished antioxidant capacity or altered membrane fluidity is not the cause of the sxd1 phenotype, the question still remains: How does disruption of TC activity result in the pleiotropic sxd1 phenotype?

In addition to the well-defined role of tocopherols as antioxidants, specific tocopherols, tocotrienols, and their oxidized products have been demonstrated to have biological activities in mammalian systems that are independent of their antioxidant functions. The unifying theme for these antioxidant-independent activities is the modification or modulation of various signal transduction pathways (Brigelius-Flohe and Traber, 1999; Ricciarelli et al., 2001; Clement et al., 2002). The effects of tocopherols on the protein kinase C signaling cascade and the synthesis of eicosanoids in mammals have been well characterized. α-Tocopherol posttranslationally inhibits the activity of protein kinase C in several mammalian systems (Chan et al., 2001; Azzi et al., 2002; Clement et al., 2002). Plant genomes also contain protein kinase C homologs and other components of this signaling pathway. Tocopherols also have been shown to posttranslationally inhibit the activity of phospholipase A2 (Tran et al., 1996; Chandra et al., 2002), cyclooxygenase (COX-2; Jiang et al., 2000; Wu et al., 2001), and lipoxygenase-5 (Greenbergley et al., 1993; Wang et al., 2000; Ricciarelli et al., 2002). These enzymes are involved in the production of eicosanoid signaling molecules (prostaglandins, prostacyclins, thromboxanes, and leukotrienes) from polyunsaturated fatty acids in mammals (Ohuchi and Levine, 1980; Tran et al., 1996; Kim et al., 2001). Like eicosanoids in animals, jasmonic acid and other oxylipins in plants are synthesized from polyunsaturated lipids by the action of lipoxygenase(s) and phospholipase(s) (Blee, 2002; Howe and Schilmiller, 2002). A third example of α-tocopherol-dependent, antioxidant-independent signal transduction in mammals is the transcriptional activator TAP.
(tocopherol-associated protein). Upon the binding of \( \alpha \)-tocopherol by TAP, the complex is translocated into the nucleus, where it has been shown to activate the transcription of transgenes (Yamauchi et al., 2001).

Although tocopherols have not yet been shown to affect protein kinase C signaling, transcriptional regulation, or the synthesis of jasmonic acid or other oxylipins in plants, studies in mammalian systems suggest plausible mechanisms whereby the absence of TC activity (and, hence, tocopherols) could affect signaling and result in the pleiotropic sxdI phenotype. Thus, it is possible that many tocopherol functions will be universal, including the roles tocopherols play in modulating signal transduction pathways or acting as signals themselves. Although the downstream events of signal transduction pathways would not necessarily be evolutionarily conserved between plants and mammals, many of the core components and biochemical motifs of signal transduction pathways are. We suggest that the sxdI phenotype is the first evidence that tocopherols act as signaling molecules or modulators of signaling in plants. Tocopherols, tocopherol derivatives, or tocopherol pathway intermediates may provide or modulate signals required for the development of maize bundle sheath vascular parenchyma plasmodesmata, analogous to the effects of tocopherols in mammalian signaling. Alternatively, the DMPBQ that accumulates in sxdI may interfere with an endogenous signaling pathway required for the process. Several groups have provided evidence that the redox status of the chloroplast, which is monitored through the plastoquinone (PQ) pool, regulates nuclear-encoded photosynthetic gene expression (Pfannschmidt et al., 1999a, 1999b, 2001; Alfonso et al., 2000; Allen and Pfannschmidt, 2000; Kujat and Owttrim, 2000; Li and Sherman, 2000; Pursiheimo et al., 2001; Trebitsh and Danon, 2001; Yang et al., 2001). DMPBQ has the same 2,3-dimethyl-1,4-benzoquinone head group as PQ and could interfere with redox signaling through the PQ pool. Another more remote possibility is that the SXD1 protein has an unknown substrate in addition to DMPBQ, and this product is required for signal transduction in maize. Although none of these models can be excluded based on the present data, they do provide a framework for future study.

The observation that Arabidopsis vte1 mutants do not exhibit phenotypes analogous to sxdI suggests that the downstream signal transduction events impacted by tocopherol deficiency differ between monocots and dicots. The pathways leading to maize bundle sheath vascular parenchyma plasmodesmata formation may either be absent or not equivalent in Arabidopsis, or the effects are too subtle to be observed at the whole-organism level as in sxdI. Experiments to assess the whole-genome responses of Arabidopsis vte1 mutants are under way.

### MATERIALS AND METHODS

#### Growth Conditions and Seed Stocks

Arabidopsis plants were grown at 22°C under a 12-h photoperiod (120 \( \mu \)E) in a vermiculite and potting soil mixture. M3 EMS-mutagenized Arabidopsis seeds (Columbia ecotype) were purchased from Lehle Seed (Round Rock, TX). vte1-1 was backcrossed to wild type three times, and vte1-2 was backcrossed twice. Maize (Zea mays) plants were grown under greenhouse conditions in the same soil mixture and fertilized biweekly with 20-20-20 fertilizer. The maize sxd1-2 allele used in this publication was isolated through the Trait Utility System for Corn (Pioneer Hybrids, Johnston, IA). The sxd1-2 Mu insertion site and mutant phenotype were described previously (Provencher et al., 2001). *Synechocystis* sp. PCC6803 was grown on BG-11 media photautotrophically or photomixotrophically (BG11 media containing 15 mm Glc) on plates or in liquid culture at 30°C and 50 to 70 \( \mu \)E light.

#### Tocopherol Analysis

For tocopherol analyses, total lipids were extracted from 30 to 35 mg of Arabidopsis or maize leaf tissue and a 500-mL culture of *Synechocystis* sp. PCC6803 cells (Bligh and Dyer, 1959; Collakova and DellaPenna, 2001), and dissolved in 100 \( \mu \)L of methanol or hexane. Methanol extracts (50 \( \mu \)L) were subject to HPLC (Agilent 1100 series, Agilent, Wilmington, DE) on a Spherisorb ODS-2 5-\( \mu \)m, 250- \( \times \) 4.6-mm reverse phase column (Column Engineering, Ontario, CA) at 28°C with a flow rate of 2 mL min\(^{-1}\) with 95% (v/v) methanol and 5% (v/v) isopropanol. Hexane extracts (50-\( \mu \)L volume) were subjected to HPLC on a RelaSil Silica 5-\( \mu \)m, 250- \( \times \) 4.6-mm normal phase column (Column Engineering) at 42°C with a flow rate of 2 mL min\(^{-1}\) with 83% (v/v) hexane and 17% (v/v) isopropanol ether. Tocopherols were detected by fluorescence using 290-nm excitation and 325-nm emission.

#### Analysis of Prenyl Quinones

One gram of Arabidopsis or maize leaf tissue and a 500-mL culture of *Synechocystis* sp. PCC6803 (OD\(_{660}\) = 0.8) were harvested and total lipids extracted (Collakova and DellaPenna, 2001). Prenyl quinones were purified by TLC as described by Pennock (1985) and eluted with diethyl ether. After drying, the samples were resuspended in 500 \( \mu \)L of hexane. A 100-\( \mu \)L aliquot was dried, resuspended in 75 \( \mu \)L of isopropanol, and subjected to HPLC on a reverse phase column (described above) under conditions previously described by Johnson et al. (2000). Alternatively, 60 \( \mu \)L of hexane extract was subjected to HPLC on a normal phase column (described above), at 30°C with a 1 mL min\(^{-1}\) flow rate using 0.1% (v/v) dioxane in hexane. Prenyl quinones were detected by A\(_{254}\) using a diode array detector.

#### Map-Based Cloning of vte1

PCR-based markers were designed using INDEL or SNP from the Cereon Arabidopsis Polymorphism and Landsberg erecta Sequence Collection (Cereon Genomics LLC, Cambridge, MA; Jander et al., 2002). DNA was extracted from 1- to 2-mm developing leaves using Plant DNazol (Invitrogen, Carlsbad, CA) following the protocol of the manufacturer. Alternatively, DNA was isolated from 1- to 2-mm developing leaves in 1.1-mL tubes arrayed in a 96-well format. Two 4-mm glass beads were added to tubes along with 200 \( \mu \)L of 10 mm Tris (pH 8.0) and 200 \( \mu \)L of chloroform. The tubes were shaken with a commercial paint shaker for 5 min and centrifuged at 3,750 rpm for 10 min. One microliter of the aqueous phase or resuspended DNA from DNazol extractions was used in a 20-\( \mu \)L PCR reaction.

#### Sequence Analysis

All DNA sequences other than vte1 mutant alleles were obtained from public databases using BLAST: wheat (*Triticum aestivum*; BQ199591), rice (*Oryza sativa*; AU031770), *Physcomitrella patens* (BJ164674), *M. truncatula* (BF641471 and TC40011), barley (*Hordeum vulgare*; TC33553 and TC32886), Arabidopsis (AF302188), and maize (AF302187). A TC prefix denotes sequences obtained from The Institute for Genomic Research. All others have GenBank accession numbers. The cyanobacteria and algae sequences were...
obtained from their respective genomic sequencing projects. Synechocystis sp. PCC6803 and Anabaena sp. PCC7120 sequences were obtained from Cyanobase (http://www.kazusa.or.jp/cyano/cyano.html). Chlamydomonas reinhardtii and Nostoc sp. Ndo25 sequences were obtained from the Joint Genome Institute (http://www.jgi.doe.gov). The Synechococcus sp. PCC7002 sequence was obtained from National Center for Biotechnology Information (www.ncbi.nlm.nih.gov). Alignments were performed using MacVector 7.0 (Oxford Publishing, London), which includes the ClustalW algorithm.

Construction of slr1737, SXD1, and VTE1 Protein Expression Vectors

Primers 5′-CATATGACCCCTAATTTACCTCTCCTTGTT-3′ (F1) and 5′-CATATGGAGAATTCCTCGGAAAGTCTGC-3′ (F2) were selected by spectinomycin resistance, and replacement of the slr1737 ORF with the slr1737 ferase was inserted as a BsiWI site separating the 5′- and 3′-flanking portions. The PCR product was cloned into the pBscript II SK+ digested with Smal and NdeI primers) into the pBluescript II SK+ digested with Smal and NdeI sites of the pET30b expression vector (Novagen, Madison, WI).

Primers 5′-CATATGAAAAATTTACGCGCGCATAGGTGTAC-3′ (F3) and 5′-GGATCCGCTCGAGAAACCAAGC-3′ (R3), and turbo Pfu polymerase (Stratagene, La Jolla, CA) were used to PCR amplify the VTE1 cDNA from a seed cDNA library (a gift from Dr. John Ohlrogge, Michigan State University, East Lansing). F2 was used to amplify the full-length cDNA, and F3 was used to amplify a truncated version of the cDNA that encodes a version of the protein lacking chloroplast transit peptide. PCR products were cloned into the Smal site of pBluescript II SK+. The 5′ ends of the VTE1 cDNA were cloned as Ndel-Xhol fragments (sites underlined in primers) into Ndel and XhoI sites of the pET30b expression vector (Novagen, Madison, WI).

Construction of Δslr1737, SXD1, and VTE1 Protein Expression Vectors

Primers 5′-CTGTTGATATTCTTGCCGTC-3′ (F3) and 5′-GGGATTGAGAGAATTCCTCCTGAG-3′ (R3) were used to PCR amplify the 5′ region flanking slr1737 from Synechocystis sp. PCC6803 genomic DNA. The PCR product was cloned into the EcoRV site of pBluescript II SK+. The slr1737 gene was cloned as Ndel-BamHI fragments (sites underlined in primers) into Ndel and BamHI sites of the pET30b expression vector (Novagen).

Primers 5′-TCTCATATGCGACCGCCACATAGCCTACC-3′ (F4) and 5′-TCTGCGCGAGCTTTGTTACATTCTGCTTG (R4) were used to amplify the SXD1 gene from Synechocystis sp. PCC6803 genomic DNA. The PCR product was cloned into the pET30b expression vector (Novagen, Madison, WI) without a chloroplast transit peptide. The PCR product was cloned into the Ndel and NotI sites (sites underlined in primers) of pET24d (Novagen). The fidelity of all constructs was confirmed by sequencing.

Generation of Δslr1737

Primers 5′-CTTGGTATATTCTTGCCGTC-3′ (F4) and 5′-GGGATTGAGAGAATTCCTCCTGAG-3′ (R4) and Pfu polymerase (Stratagene) were used to PCR amplify the 5′ region flanking slr1737 from Synechocystis sp. PCC6803 genomic DNA. Primers 5′-ATAAATACCTCAGCTCGGATGAAAATTACGCGGCATAGGTGTAC-3′ (F5) and 5′-ACGGTCTTACCTAACGACAGTGC-3′ (R5) and Pfu polymerase (Stratagene) were used to PCR amplify the 3′ region flanking slr1737 from Synechocystis sp. PCC6803 genomic DNA (underlined nucleotides indicate an added BsiWI restriction site). The 5′- and 3′-flanking PCR products were joined together through re-amplification with Pfu polymerase using Primers F5 and R6 to generate a chimeric 1,441-bp fragment containing a BsiWI site separating the 5′- and 3′-flanking portions. The PCR product was cloned into the pPCR-Script AMP vector (Stratagene) to generate pSLR1737-5′-flank. The aidA gene encoding spectinomycin adenyltransferase was inserted as a BsiWI fragment into the corresponding site of pSLR1737-5′-flank. The resulting plasmid was then used to replace ORF slr1737 with homologous recombinant (Williams, 1988). Recombinant lines were selected by spectinomycin resistance, and replacement of the slr1737 ORF with the aidA gene was confirmed by PCR.

Complementation of Δslr1737 by Expression of the Maize SXD1 cDNA

A vector, designated pSy6Exp2, was used to express SXD1 in Synechocystis sp. PCC6803, pSy6Exp2 was derived from pPCR-Script AMP (Stratagene) and contained the Synechocystis sp. PCC6803 psbA2 promoter linked to a multicloning site to Tn9, which encodes chloramphenicol acetyltransferase. To facilitate homologous recombination, the promoter and multicloning site were flanked by S12- and 429-bp sequences from the slr2699 locus. Primers 5′-TTTTTTTTTTGTAGCGACGCGCGCATAGCGGTTACCC-3′ (F6) and 5′-TTTTTTTTTTGTAGCGACGCGCGCATAGCGGTTACCC-3′ (R6) and Pfu polymerase were used to PCR amplify the coding sequence of the mature maize SXD1 polypeptide from the SXD1 cDNA. The PCR product was cloned into the Ndel and BstI restriction sites of the multicloning site behind the psbA2 promoter. The resulting plasmid was used to transform Δslr1737 through homologous recombination of the slr2699 locus. Recombination events were selected by chloramphenicol resistance, and the introduction of the maize SXD1 cDNA into slr1737 knockout mutants was confirmed by PCR analysis of genomic DNA isolated from chloramphenicol- and spectinomycin-resistant (selection for Δslr1737) resistant cell lines.

TC activity from Escherichia coli-Expressed Proteins

C45 (DE3) cells containing the relevant pET vectors engineered to express TCs from the three organisms were grown at 30°C in 50 mL of Luria-Broth culture (50 μg mL⁻¹ kanamycin) to mid-log phase (0.4-0.6 OD595 nm) then 1 mM isopropyl-β-D-galactoside was added and grown at 15°C for 18 h (Moiroux and Walker, 1996). Cells were harvested by centrifugation and washed with 100 mM KH2PO4 (pH 7.8) and 4 mM MgCl2. The cells were resuspended in 1 mL of 100 mM KH2PO4 (pH 7.8), 4 mM MgCl2, 4 mM diithiothreitol, and 4 mM glutathione, and sonicated six times for a duration of 10 s at 40 Hz. Dodecyl maltoside was added to 0.8 mM and Triton X-100 to 0.08 mM, and the cell lysate was shaken gently for 1 h at 4°C. Radiolabeled DMNPQ substrate was prepared by enzymatic labeling of DMNPQ with 14C-adenosyl-1-Met (Amersham, Piscataway, NJ) using heterologously expressed Synechocystis sp. PCC6803 DMNPQ methyl transferase (Shintani et al., 2002). The labeled DMNPQ was purified from the reaction by TLC (Pennock, 1985). Each 100-μL TC assay contained 4.5 × 10⁻⁴ μCi of 14C-3,5-dimethyl-6-phytyl-1,4-benzoquinol. The substrate was reduced with 2 mg of NaBH4 immediately before the assay. Each reaction contained 100 mM KH2PO4 (pH 7.8), 4 mM MgCl2, 0.8 mM diithiothreitol, 0.8 mM glutathione, 0.8 mM dodecyl maltoside, 20 mM ascorbic acid, 1 mM cycloextrin, and 20 μL of cell lysate from E. coli expressing SXD1, VTE1, or slr1737. The dodecyl maltoside and cycloextrin were added first to solubilize the substrate. The reactions were sparged with nitrogen gas and gently shaken for 4 h at room temperature. Total lipids were extracted in extraction buffer containing 1 mg mL⁻¹ butyldihyroxyxylolene and unlabeled 20 μg mL⁻¹ γ-tocopherol as a carrier. The prenyl lipids were separated by TLC, and the presence of the unlabeled prenyl quinone and tocopherol standards was determined by staining with Emmeric-Engel reagent (Pennock, 1985). 14C-labeled compounds were detected by a 4-d exposure to a low-energy Phosphorimage screen (Molecular Dynamics, Piscataway, NJ) and analyzed using a PhosphorImager (Molecular Dynamics).

Analysis of Glc, Suc, and Starch in Leaves

Mature leaves from 4-week-old Arabidopsis plants were harvested at the end and the beginning of the photoperiod. Sugars were extracted from leaf tissue in 80% (v/v) ethanol at 80°C for 30 min. Starch was also extracted from the cleared leaf tissue using 0.2 M KOH at 95°C for 45 min and neutralized with 1 M acetic acid to pH 5.0. The sugars and starch levels were measured enzymatically through the conversion of NAD to NADH by Glc-6-phosphate dehydrogenase and observed at 340 nm as described (Stitt et al., 1989).

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LITERATURE CITED
