Carotenoids comprise a large group of terpenoid pigments synthesized by all plants, algae and cyanobacteria as well as by several nonphotosynthetic bacteria and fungi. Carotenoids fulfill indispensable functions in photosynthesis (Frank et al., 1999), and in plants they also provide colors to flowers and fruits. Dietary carotenoids are essential precursors to vitamin A, and they are implicated in reducing the occurrence of certain cancers and cardiovascular diseases, possibly by serving as antioxidants and free radical scavengers (for review, see Demmig-Adams and Adams, 2002; Fraser and Bramley, 2004). Our understanding of the carotenoid biosynthetic pathway in plants has been advanced greatly in the past decade, mainly due to cloning of many of the genes for enzymes involved in the pathway. The first carotenoid in the pathway is phytoene, a product of condensation of two geranylgeranyl diphosphate (GGDP) molecules. The colorless phytoene undergoes four consecutive dehydrogenation reactions that introduce four double bonds in conjugation, giving rise to the red polyene chromophore of lycopene, the precursor of a diverse group of cyclic carotenoids. In plants they also provide colors to flowers and fruits. Dietary carotenoids are essential precursors to vitamin A, and they are implicated in reducing the occurrence of certain cancers and cardiovascular diseases, possibly by serving as antioxidants and free radical scavengers (for review, see Demmig-Adams and Adams, 2002; Fraser and Bramley, 2004).

Most enzymes in the central pathway of carotenoid biosynthesis in plants have been identified and studied at the molecular level. However, the specificity and role of cis-trans-isomerization of carotenoids, which occurs in vivo during carotene biosynthesis, remained unresolved. We have previously cloned from tomato (Solanum lycopersicum) the CrtISO gene, which encodes a carotene cis-trans-isomerase. To study the biochemical properties of the enzyme, we developed an enzymatic in vitro assay in which a purified tomato CrtISO polypeptide overexpressed in Escherichia coli cells is active in the presence of an E. coli lysate that includes membranes. We show that CrtISO is an authentic carotene isomerase. Its catalytic activity of cis-to-trans isomerization requires redox-active components, suggesting that isomerization is achieved by a reversible redox reaction acting at specific double bonds. Our data demonstrate that CrtISO isomerizes adjacent cis-double bonds at C7 and C9 pairwise into the trans-configuration, but is incapable of isomerizing single cis-double bonds at C9 and C9'. We conclude that CrtISO functions in the carotenoid biosynthesis pathway in parallel with \( \zeta \)-carotene desaturation, by converting 7,9,9'-tri-cis-neurosporene to 9'-cis-neurosporene and 7'-9'-di-cis-lycopene into all-trans-lycopene. These results establish that in plants carotene desaturation to lycopene proceeds via cis-carotene intermediates.

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This study was carried out in order to characterize CRTISO biochemically. For this purpose, we developed an in vitro assay that relied on CRTISO from tomato expressed in E. coli. Activity of cis-to-trans isomerization was analyzed with different carotenoid isomers substrates under various reaction conditions. The results determined that the intermediates of the carotenoid biosynthesis pathway from phytoene to neurosporene are cis-configured. The CRTISO enzyme requires active cell membranes, and its activity is driven by a membrane-bound redox chain.

RESULTS

Functional Expression of CRTISO in E. coli Cells

E. coli cells carrying plasmid pAC-Zeta accumulate \( \zeta \)-carotene (Isaacson et al., 2002). This carotene intermediate typically occurs in three isomeric forms that can be resolved by HPLC system 2 (Fig. 1). To identify the \( \zeta \)-carotene species accumulated in the bacteria, we used synthetic all-trans-\( \zeta \)-carotene as a reference. It was found that the E. coli cells accumulated two types of cis-isomers and only trace amounts of the all-trans-isomer. Based on published data (Clough and Pattenden, 1983; Beyer et al., 1989), we assigned peak 2 in Figure 1 as 9,15,9'-tri-cis-\( \zeta \)-carotene and peak 3 as 9,9'-di-cis-\( \zeta \)-carotene. This identification is corroborated by the typical absorption at 297 nm of the 9,15,9'-tri-cis-\( \zeta \)-carotene and by the fact that the ratio of the peaks 2 and 3 changed from 3:2 in dark-grown bacteria to 2:3 when the bacteria were grown under illumination. This phenomenon is attributed to photoisomerization of the C15 cis-double bond to trans (Beyer et al., 1989). Complementation of CRTISO in E. coli cells that carried plasmids pAC-Zeta and pCRTISO did not change the pattern of \( \zeta \)-carotene isomers, thus demonstrating that cis-\( \zeta \)-carotenones are not substrates of CRTISO. It is important to note that in the same E. coli cells, CRTISO is active in isomerizing 7,9,9'-tri-cis-neurosporene and prolycopene (see below). It is concluded that isomerization of carotenones by CRTISO to form trans-intermediates takes place during or after the desaturation of \( \zeta \)-carotene and not during the phytoene desaturation. For unknown reason, a small increase of phytoene and phytofluene was always observed in cells that expressed CRTISO. This phenomenon could be due to interference of CRTISO with the activity of phytoene desaturase or because of an indirect effect of overexpressing an additional protein in the bacteria.

Substrate Specificity of CRTISO

E. coli cells carrying pCRTISO654 accumulated a polypeptide of an apparent molecular mass of approximately 60 kD, matching the predicted size of the mature CRTISO polypeptide from tomato after cleavage of the predicted transit peptide (Fig. 2). These cells were disintegrated by a French press, and the lysate was centrifuged at 13,000g. Based on SDS-PAGE of polypeptides from the pellet and supernatant, an estimated 5% to 10% of the recombinant protein was in the supernatant while the rest was in inclusion bodies (Fig. 2). The supernatant of the lysate served as a crude enzyme preparation used in vitro to assay the enzymatic activity of CRTISO. The supernatant contained plasma membranes showing active respiration upon NADH addition, as measured with an oxygen electrode (data not shown). Equivalent supernatants of E. coli cells transformed with the empty vector pQE60 were used as controls in all assays.

Figure 1. HPLC analysis of carotenones extracted from E. coli cells accumulating cis-\( \zeta \)-carotene in the absence (A) and presence (B) of CRTISO. To identify the isomers, the sample in B was mixed with a synthetic all-trans-\( \zeta \)-carotene and analyzed by HPLC (C). MaxPlot chromatograms showing each peak at its \( \lambda_{max} \) are presented. Peak 1, All-trans-\( \zeta \)-carotene; peak 2, 9,15,9'-tri-cis-\( \zeta \)-carotene; peak 3, 9,9'-di-cis-\( \zeta \)-carotene; peaks 4a and 4b are cis-isomers of phytofluene; peak 5, 15-cis-phytoene. Absorption spectra of specific peaks are presented to the boxes. 9,15,9'-Tri-cis-\( \zeta \)-carotene is distinguished from the 9,9'-cis-isomer by the typical absorbance at 297.5 nm.
The mixture of cis-carotenes extracted from ripe fruits of the tomato mutant *tangerine* (LA3183) was used as a substrate in the initial experiments. These carotenoids consisted of 30% to 50% 7,9,7,9'-tetra-cis-lycopene (prolycopene), approximately 2% 7,9-di-cis-lycopene, 6% to 11% 7,9,9'-tri-cis-neurosporene, 20% to 30% 9,9'-di-cis-z-carotene, 9,15,9'-cis-z-carotene, and approximately 8% to 10% 9-cis-phytofluene. This composition is consistent with the data given by Clough and Pattenden (1983), who used NMR to determine the various carotenoids. cis-Carotenes applied to the assay in the form of CHAPS micelles or in small amounts of acetone were isomerized very poorly or not at all. When the cis-carotenes were applied to the assay in liposomal membranes made of either *E. coli* lipid extracts or dimyristoyl phosphatidyl choline (DMPC), most of the prolycopene was isomerized by CRTISO into di-cis-lycopene and only a small portion into all-trans-lycopene (data not shown). Therefore, we used poly-cis-carotene substrate solubilized in 0.25% (w/w) Triton X-100 routinely.

Incubation of bacterial supernatant containing the CRTISO with cis-carotenes in the dark at 28°C for up to 2 h resulted in a substantial cis-to-trans isomerization of carotenes (Fig. 3). Time-course analysis of carotenoid conversion showed that prolycopene decreased concomitantly with an increase in di-cis- and all-trans-lycopene (Fig. 3A). Similarly, 7,9,9'-tri-cis-neurosporene decreased concomitantly with an increase of two other isomers of neurosporene, none of which was all-trans-neurosporene, which was identified by running a standard. It is worth noting that similar to the complementation study, the relative proportion of cis-isomers of z-carotene, phytofluene, and phytoene did not change significantly during the incubation. No conversion of cis-carotenes was observed in the control assays (data not shown). To examine whether 7,9,9'-tri-cis-neurosporene alone is a substrate of CRTISO, this proneurosporene was purified from *tangerine* fruits by HPLC system 1 and applied to the in vitro assay (Fig. 4B). Several lines of evidence suggest that the main product of the isomerization of 7,9,9'-tri-cis-neurosporene is 9-cis-neurosporene (peak 3a in Fig. 3). (1) An all-trans-neurosporene standard ruled out this species as a product. (2) In the HPLC analysis used, a longer retention time is correlated with increasing number of cis-double bonds in the molecule of a specific carotenoid. The neurosporene isomer produced by CRTISO was eluted in the HPLC immediately after the all-trans-neurosporene standard and before the two other isomers, one of which was clearly identified as 7,9,9'-tri-cis-neurosporene. This fits an isomer with a single cis-double bond. (3) A

![Figure 2. Expression of CRTISO in *E. coli*. Proteins were separated by SDS-PAGE and stained with Coomassie blue. Lane 1, Insoluble fraction (pellet) of a lysate from *E. coli* carrying pQE60; lane 2, soluble fraction (supernatant) of *E. coli* carrying pQE60; lane 3, insoluble fraction of *E. coli* carrying pCrtISO654; lane 4, soluble fraction of *E. coli* carrying pCrtISO654; lane 5, column-purified CRTISO polypeptide.](image)

![Figure 3. HPLC analysis of cis-carotene substrates and products in a CRTISO in vitro assay. A, Carotenoids at the beginning of the reaction (time 0 min); B, after 90 min. Peak 1a, trans-Lycopene; peak 1b, di-cis-lycopene; peak 2, prolycopene; peak 3a, neurosporene, isomer 1; peak 3b, neurosporene, isomer 2; peak 3c, 7,9,9'-cis-isomer of neurosporene; peak 4a, 9,15,9'-cis-z-carotene; peak 4b, 9,9'-cis-z-carotene isomer; peak 5a and 5b are β-carotene isomers; peak 6, phytofluene; peak 7, phytoene.](image)
7-cis-neurosporene or 7,9-di-cis-neurosporene are excluded because, as was determined with 9,9'-di-cis-z-carotene, the cis-double bond in position 9 is not a substrate for CRTISO. Taken together, these results indicate that CRTISO does not recognize a single cis-double bond at C9 or C9' but utilizes substrates containing 7,9 (or 7',9')-cis-pairs of adjacent double bonds.

Biochemical Analysis of CRTISO Activity in Vitro

A purified His-tagged form of CRTISO was enzymatically inactive with poly-cis-carotenes when present in Triton X-100 micelles or in liposomes prepared from E. coli lipids or DMPC. However, CRTISO-mediated isomerization occurred only when supernatant of E. coli lysate obtained from cells expressing the empty vector (pQE60) was added to the assay system instead of the liposomes or Triton X-100 micelles. This indicated that a component in the bacterial supernatant was essential for CRTISO activity in vitro. Dialysis of the E. coli cells lysate prior to its addition to the in vitro assay abolished this activity, indicating that a dialyzable component in the bacterial lysate supernatant was essential (Fig. 5A). We considered the possibility that the isomerization reaction proceeds via a reversible hydrogen extraction. Such a process could involve an oxidoreduction process that depends on activity of components of the respiratory electron transfer chain of the bacterial membranes still present in the lysate. To test for this possibility, soluble electron donors or acceptors of various nature and/or redox potentials were added to the dialyzed cell lysate referred to as the...

Figure 4. A, Time course of prolycopene isomerization in vitro. A mixture of cis-carotenes from tangerine, at a final concentration of 180 μg mL-1, was used as a substrate in the assay with lysate of E. coli that expressed CRTISO. Aliquots were subjected to HPLC analysis at the indicated time points. Dashed line, Prolycopene; dotted line, 7,9-di-cis-lycopene; solid line, trans-lycopene. B, Reaction carried out with purified 7,9,9'-tri-cis-neurosporene as a substrate. Solid line, 7,9, 9'-Tri-cis-neurosporene; dashed line, neurosporene isomer 1; dotted line, neurosporene isomer 2; dotted-dashed line, trans-neurosporene.

Figure 5. Effect of redox components on the activity in vitro of CRTISO. Time-course experiments were run and plotted as in Figure 4A in the presence of the different effectors. For each condition, the area under the curves was calculated. CRTISO activity in the control condition was taken as 1.0. A, Requirement for respiratory redox intermediates; black bars represent nondialyzed E. coli lysate (supernatant) and gray bars, dialyzed preparations. B, Influence of anaerobic conditions and ferricyanide. C, Effect of various oxidized quinones on CRTISO activity measured in the absence (black bars) or presence (gray bars) of 1 mM NADH. BQ, benzoinoquione; DQ, duroquinone; UQ, ubiquinone; PQ, plastoquinone (1 mM each).
basal assay system. Among these, electron donors such as NADH, NADPH, or succinate restored the activity to the level of the nondialyzed supernatant and even increased to some extent the activity of the enzyme when assayed using the nondialyzed lysate (Fig. 5A).

The promoting effect on CRTISO activity of succinate can be explained in terms of membrane participation and an active succinate:ubiquinone oxidoreductase (succinate dehydrogenase) being present and points to the involvement of ubihydroquinone (UQH$_2$). The CRTISO reaction mechanism possibly depends on active redox chains, such as the respiratory redox chain or parts thereof downstream of UQH$_2$, which are present in E. coli membranes and which are added with the supernatant. To investigate this hypothesis, we applied anaerobic conditions to the assay by using an enzymatic oxygen trap leading to an accumulation of reduced redox-active components in the membrane. Anaerobic conditions enhanced the cis-to-trans isomerization activity of CRTISO (Fig. 5B). This result points to the involvement of a membrane-bound electron donor and excludes the possibility of participation of molecular oxygen in the reaction mechanism. It is in agreement with these findings that the addition of ferricyanide (1–5 mM) as an oxidizing agent inhibited the reaction (Fig. 5B).

To test for the hypothesis of UQH$_2$ involvement, we examined several quinones in their reduced form. Addition of hydroquinones such as benzoquinone, duroquinone, plastoquinone, or ubiquinone had no effect on the reaction (data not shown). However, addition of the above quinones in their oxidized state inhibited the reaction (Fig. 5C). This inhibition points to the participation of hydroquinones and is further corroborated by the fact that this inhibition could be overcome by adding NADH (Fig. 5C).

Antimycin A is a known specific inhibitor of cytochrome b, which acts as an electron acceptor downstream of UQH$_2$ (Izzo et al., 1978). In the presence of antimycin A, cytochrome b can be reduced but not oxidized. No significant effect on isomerization in vitro was observed with antimycin A (data not shown), which excludes the participation of complex III (ubiquinone cytochrome c-oxidoreductase) and IV (cytochrome c-oxygen oxidoreductase) in the phenomenon observed.

Overall, these results suggest that presence of a reduced component(s) of the electron transfer chain is required for the isomerization activity of CRTISO in the basal assay system. The reduced cofactor, acting as an intermediate carrier between succinate dehydrogenase of complex II or NADH ubiquinone oxidoreductase of complex I and cytochrome b of complex III, most probably UQH$_2$, is involved in CRTISO cis-to-trans isomerization. This cofactor undergoing oxidation-reduction is poised at a finely tuned redox potential that serves as intermediate in the isomerization process, possibly via a reversible hydrogen donation/extraction at a specific cis-oriented double bonds.

**DISCUSSION**

**Carotenoid Biosynthesis in Plants Follows the cis-Desaturation Pathway**

Most carotenoids in plants occur in the all-trans-configuration. The carotene intermediates through the four desaturation reactions from phytoene to lycopene have been considered to be in the all-trans-configuration (Spurgeon and Porter, 1980; Britton, 1988; Hugueney et al., 1992). However, occurrence of cis-carotenes was reported in mutants of tomato (tangerine; Zechmeister et al., 1941), Scenedesmus (Humbeck, 1990), and Euglena (Cunningham and Schiff, 1985), in experiments with purified membranes from daffodil (Narcissus pseudonarcissus) chromatplasts (Beyer et al., 1989) and in E. coli cells expressing plant-type biosynthetic enzymes (Bartley et al., 1999; Isaacson et al., 2002; Matthews et al., 2003). Consequently, a poly-cis-pathway of carotene desaturation has been often regarded as being abnormal or as an in vitro artifact. This cis-versus-trans pathway conundrum has remained unsolved for nearly six decades. Characterization of CRTISO that is described here approves the view that cis-configured carotenoids that are found in tangerine represent the legitimate intermediates that normally exist in the plant carotenoid biosynthesis pathway.

It is a well-established fact that the central C15-15’ double bond in phytoene is cis-configured (Britton, 1998). The first desaturation reactions of phytoene catalyzed by PDS can take place with 15-cis-phytoene, producing trans-double bonds at C11 and C11’. This is shown by the existence of 15-cis-phyllofluene and 15-cis-ζ-carotene (Kushwaha et al., 1970; Powls and Britton, 1977; Beyer et al., 1994). However, during ZDS-mediated desaturation, this double bond is found to be trans-configured. Moreover, as demonstrated in this work (Fig. 1), the cis-double bond at C15 is not a substrate for isomerization by CRTISO. It was shown in vitro that ZDS activity was regained only after isomerizing this cis-double bond to trans (Beyer et al., 1989). Since the 15-cis-double bond is known to be very photo labile, this step can be enhanced by light in vitro. How this is achieved in vivo remains unclear. It is evident from the data we present here that CRTISO does not play this part because it leaves the isomeric states of cis-ζ-carotene species unaffected. Thus, the 15-cis-double bond is distinct from other cis-double bonds in carotenoids that are discussed here.

The double bonds at positions 9 and 9’ are not a product of the desaturation sequence but originate from the prenyl transferase reactions that produce GGDP. These double bonds exist in trans in the phytoene found in wild-type as well as in tangerine tomatoes (Clough and Pattenden, 1979, 1983; Britton, 1998). Therefore, there is currently no support for the possibility that the cis-double bonds at positions 9 and 9’ in prolycopene are derived from cis-configured GGDP. Based on NMR analysis of fruit carotenoids from...
the mutant tangerine, Clough and Pattenden (1979, 1983) suggested that during the introduction of the trans-double bonds at position 11 (in phytoene) and 11′ (in phytofluene) by PDS, the double bonds at positions 9 and 9′ undergo specific trans-to-cis isomerization (Clough and Pattenden, 1979, 1983). Accumulation of cis-carotenoids in the mutant C-6 D of the alga Scenedesmus obliquus grown in the dark was explained by impairment of the synthesis of all-trans-β-carotene, possibly due to a mutation in the phytoene desaturase (Sandmann, 1991).

By contrast, our data support the poly-cis-pathway to be the default mechanism of carotenoid desaturation in plants and cyanobacteria. It is evident that CRTISO was unable to isomerize the cis-isomers of β-carotene in the in vitro assay, while it was actively converting tri-cis-neurosporene (proneurosporene) and tetra-cis-lycopene (prolycopene). This is in agreement with CRTISO recognizing only cis-double bonds at positions 9 or 9′ in conjunction with adjacent cis-double bonds at 7 or 7′. This is further substantiated by the fact that, in our assays, CRTISO converted prolycopene into all-trans-lycopene, through the 7,9-di-cis-lycopene intermediate, as has been suggested previously (Beyer et al., 1991). Additional evidence for a cis-desaturation pathway is the fact that the plant-type carotene desaturases produce in E. coli only cis-isomers of β-carotene as well as prolycopene (Bartley et al., 1999). Furthermore, all the β-carotene detected in trace amounts during fruit ripening in the wild-type tomato, and in large amounts in fruits of a mutant that is impaired in β-carotene desaturation, occurs in the 9,9′-cis- and the 9,15,9′-cis-configuration (data not shown). Finally, ZDS is capable of using 9′,9-di-cis-β-carotene as a substrate to produce prolycopene via 7,9,9′-tri-cis-neurosporene (Breitenbach et al., 1999). Since 7,9,9′-tri-cis-neurosporene is efficiently isomerized in vitro by CRTISO, most probably to 9′-cis-neurosporene, we reason that in vivo this proneurosporene isomer is most likely the authentic substrate for isomerization in the normal desaturation pathway. This inference is supported by the fact that we have not been able to detect any prolycopene in wild-type plants under normal conditions while an isomer of di-cis-lycopene could be detected in trace amounts. Therefore, we conclude that CRTISO acts in conjunction with or immediately after ZDS during the desaturation of β-carotene to all-trans-lycopene (Fig. 6).

CRTISO Activity Requires a Redox Mechanism

The findings that reducing reagents such as NADH, NADPH, and succinate accelerate the isomerization reaction in vitro and that oxidizing reagents such as ferricyanide, oxygen, or different oxidized quinones are inhibitory suggest the involvement of the bacterial membrane’s respiratory chain in the process. More specifically, complex I, the NADH ubiquinone oxidoreductase, and complex II, the succinate ubiquinone oxidoreductase, affect the reaction, probably via the quinone pool. These results suggest that, although no net electron transfer occurs in the isomerization reaction, a redox driving force is needed. The isomerization reaction may include a transient reduction of the cis-double bond followed by reoxidation yielding a trans-configuration of the newly formed double bond. The fact that CRTISO harbors a conserved dinucleotide-binding domain at its N terminus supports this possibility. A dinucleotide-binding domain is highly conserved among many of the carotenoid biosynthesis enzymes. PDS from pepper was found to

Figure 6. Proposed pathway of carotenoid biosynthesis in plants. CRTISO, Carotene isomerase; CRTR-B, β-ring hydroxylase, CRTR-E, ε-ring hydroxylase; GGPP, geranylgeranyl diphosphate; LCY-B, lycopene β-cyclase; LCY-E, lycopene ε-cyclase; PDS, phytoene desaturase; PSY, phytoene synthase; ZDS, β-carotene desaturase; ZEP, zeaxanthin epoxidase.
bind flavin adenine dinucleotide (FAD) (Hugueneau et al., 1992), and PDS from daffodil was found to be active only when FAD was added to the medium during its membrane association (Al-Babili et al., 1996). Lycopene β-cyclase was shown to be active in vitro only in the presence of NADH or NADPH (Schnurr et al., 1996), although NADPH does not transfer its hydrogen during the cyclization reaction to the carotene (Hornero-Mendez and Britton, 2002). We have not been able to detect any bound flavin in the His-tagged purified CRTISO. The nature of the redox-active CRTISO dinucleotide cofactor remains to be elucidated. However, it cannot be excluded that its binding occurs only transiently during catalysis. The electrons that participate in the reaction might be transferred back to this cofactor and fed into the electron transfer chain.

The interaction of CRTISO with the electron chain seems rather unspecific. This is because it was assayed with E. coli membranes, while it must be assumed that the photosynthetic electron transport is involved in plants. This is somewhat comparable to PDS function in E. coli (Bartley et al., 1999; this study) that employs quinones and depends on active redox chains (Mayer et al., 1990; Nievelstein et al., 1995). Here again, the photosynthetic electron transport is thought to provide the driving force that is replaced by an alternative redox chain in nongreen plastids. Consistent with this line of reasoning, the involvement of an alternative terminal oxidase was recently shown in Arabidopsis (Carol et al., 1999; Wu et al., 1999; Carol and Kuntz, 2001; Joet et al., 2002) and tomato (Josse et al., 2000). Just like PDS, CRTISO appears to be able to mechanistically link to respiratory as well as photosynthetic redox chains. This again points to the involvement of a benzoquinone as the common redox component in respiration as well as in photosynthesis.

In contrast to carotene desaturation, however, there is no involvement of oxygen in carotene isomerization, as we show here, supporting earlier evidence (Beyer et al., 1991).

### Phytorene Desaturation in Plants and Bacteria

In plants and cyanobacteria, phytorene desaturation to all-trans-lycopene is catalyzed by three enzymes, PDS, ZDS, and CRTISO, whereas in bacteria a single enzyme, CRTI, carries out this function. The bacterial CRTI and plant CRTISO share sequence similarity and probably evolved from a common ancestor (Giuliano et al., 2002; Isaacson et al., 2002; Park et al., 2002). However, CRTISO does not possess any desaturase activity. Although both CRTI and PDS were shown to bind FAD (Al-Babili et al., 1996; Dailey and Dailey, 1998), the mechanism of dehydrogenation of phytorene is evidently different. One aspect relates to the sequence of dehydrogenation in the two enzymes. In plants the desaturation of phytorene by PDS is symmetrical at C11 and C11', to produce a symmetric cis-isomer of ζ-carotene (7,8,7',8'-tetrahydrolycopene), whereas CRTI-type phytorene desaturases apparently dehydrogenate phytorene in one-half of the molecule first to produce an asymmetric isomer of ζ-carotene (7,8,11,12-tetrahydrolycopene). Support for this possibility comes from the detection of 7,8,11,12-tetrahydrolycopene in bacteria (Marshall and Wilmoth, 1981) and the ability of CRTI enzymes to convert phytorene into other asymmetric end products, such as neurosporene (Raisig et al., 1996; Armstrong, 1997) and 3,4-didehydrolycopene (Hausmann and Sandmann, 2000).

The PDS:ZDS:CRTISO desaturation of phytorene appeared in evolution first in cyanobacteria. Since non-oxygenic photosynthetic bacteria that possess CRTI

<table>
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<th>Peak No.</th>
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<tr>
<td>Standard</td>
<td>trans-Neurosporene</td>
<td>420, 445, 474</td>
<td>90</td>
</tr>
<tr>
<td>Standard</td>
<td>trans-ζ-Carotene</td>
<td>380, 401, 425</td>
<td>95</td>
</tr>
<tr>
<td>Standard</td>
<td>trans-β-Carotene</td>
<td>430, 455, 484</td>
<td>21</td>
</tr>
</tbody>
</table>

<sup>a</sup> Absorbance spectra as detected in the HPLC running solvent: a gradient of acetonitrile:water (9:1) and ethylacetate (HPLC system 2).

<sup>b</sup> % III/II, Fine structure of the absorption spectra expressed as the relative heights of the longest wavelength peak (III) to the middle peak (II). Zero means lack of peak III.
preceded cyanobacteria in evolution, the emergence of a three-enzyme desaturation pathway may be associated with the development of oxygenic photosynthesis. The reason for this is yet unknown. One possibility is a biochemical and/or biophysical constraint that necessitates a different enzymatic mechanism of carotenoid desaturation in cyanobacteria and plants in which poly-cis-carotene intermediates are only by-products. Alternatively, the reason for the emergence of a different desaturation pathway is a requirement to maintain the carotenoid intermediates in a cis-configuration. Support for the latter hypothesis comes from the finding that cis-isomers of the early biosynthetic intermediates are more stable than the corresponding trans-isomers (Panthet al., 1992).

MATERIALS AND METHODS

Expression of CRTISO in *Escherichia coli*

Plasmid pAC-Zeta, which carries the genes *crtb* and *criE* from *Erwinia herbicola* and *criP* from *Synechococcus* PCC7942, has been described (Cunningham et al., 1994). All strains of *E. coli* contained plasmids pCB-Ipi that was constructed by inserting the cDNA of Ipi from *Haematococcus pluvialis* (Cunningham and Ganit, 2001; kindly provided by F. Cunningham, University of Maryland, College Park, MD) into the HindIII site of the plasmid vector pGB2 (Churchward et al., 1984). Plasmid pCRTISO, which carries the cDNA of CRTISO from tomato (*Solanum lycopersicum*), has been described (Isaacson et al., 2002). *E. coli* cells of the strain XL-1 Blue carrying plasmids pAC-Zeta and pCRTISO were cotransformed with plasmids pAC-Zeta and pCRTISO and selected on solid Luria-Bertani medium containing 1.5% agar and the appropriate antibiotics: spectinomycin (50 mg/L), ampicillin (100 mg/L), and chloramphenicol (50 mg/L). Cells were incubated overnight at 37°C. Bacterial cells were harvested, resuspended in one-tenth volume of cold incubation buffer (100 mM Tris, 100 mM NaCl, 100 mM imidazole, pH 7). The recombinant protein contained a 6×His tag at its C terminus.

Carotenoid Analysis by HPLC

Carotenoids were separated by HPLC using a Waters system and a Spherisorb ODS2 C18 (5 μm, 4.6 × 250 mm) reversed-phase column (Waters, Milford, MA). In HPLC system 1, acetonitrile was used as the eluent at a constant flow rate of 1.6 mL/min. HPLC system 2 was a gradient. Using acetonitrile:water (9:1; A) and ethylacetate (B), at a constant flow rate of 1.6 mL/min the gradient was: 100% to 80% A during 8 min; 80% to 65% A during 4 min, followed by 65% to 45% A during 14 min and a final segment at 100% B. Carotenoids were analyzed with a Waters 996 photodiode array detector as described previously (Ronen et al., 2000). Carotenones were identified by their absorption spectra and retention time, and in some cases by comparison with authentic reference substances (see Table I for details). A standard of all-trans-neurosporene was obtained by extraction from *E. coli* cells carrying plasmids pAC-NEUR (Cunningham et al., 1994). A standard of trans-′-carotene was purchased from CaroteNature GmbH (Luspingen, Switzerland). The pattern of cis-carotenones extracted from fruit of the tomato mutant *tangerine* was essentially identical to data given by Clough and Pattenden (Clough and Pattenden, 1979, 1983). Quantification was performed by integrating the peak areas using the Millenium chromatography software (Waters).

Extraction of cis-Carotenones from *tangerine* Fruit

Fruits of the *tangerine* tomato mutant were ground in chloroform:MeOH (2:1, v/v). The chloroform phase was collected and carotenones were separated by thin-layer chromatography (Silica gel 60 F254, Merck, Rahway, NJ) with petroleum ether:dichloromethane, 4:1:1. The carotene fraction migrating with the solvent front was scraped off, eluted with acetone, and dried under flow of nitrogen in the dark. This total carotene fraction, when used as substrates, was dissolved in 1% Triton X-100 in sonication buffer (20 mM Tris-HCl, pH 8, 100 mM NaCl) by sonication and further diluted in sonication buffer to a final concentration of 0.25% (v/v; carotene concentration of approximately 180 μg mL⁻¹). In a similar way, carotenones were solubilized in a solution of 1 mM CHAPS. Individual cis-carotene species were isolated from the total carotene fraction by using HPLC system 1. Fractions containing selected cis-carotenones were collected, dried under a stream of nitrogen, and dissolved in 0.25% Triton X-100 as described above.

Carotenoid Extraction from Bacterial Lysate

Supernatants of bacterial lysate were mixed with acetone to a concentration of 80% and centrifuged at 4,000g for 5 min to pellet bacterial debris. The supernatant was collected and 1/3 the volume of petroleum ether was added. The organic phase was collected, dried, and dissolved in 65 μL of acetone and applied to HPLC analysis.

In Vitro Isomerization Assay

Thirty microliters of the cis-carotene preparation in 0.25% Triton X-100 (approximately 180 μg mL⁻¹) were added per milliliter of bacterial extract. Samples were incubated at 30°C in the dark. At various times, 1-mL aliquots were extracted with 4 mL of acetone. Anaerobic conditions were applied in some experiments using an enzymatic oxygen trap according to Lam and Malkin (1982) consisting of catalase, 45 μg mL⁻¹; G6P dehydrogenase, 300 μg mL⁻¹; and G6Cl, 1,500 μg mL⁻¹.

Liposome Preparation

*E. coli* lipids were extracted with chloroform:MeOH (2:1, v/v). Carotenones from *tangerine* tomato fruit dissolved in chloroform were added. The organic phase was collected, dried and dissolved in 65 μL of acetone and applied to HPLC analysis.

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