Current Status of Marine-Derived Compounds as Warheads in Anti-Tumor Drug Candidates
Cloning and Functional Characterization of a Lycopene β-Cyclase from Macrophytic Red Alga *Bangia fuscopurpurea*

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Abstract: Lycopene cyclases cyclize the open ends of acyclic lycopene (ψ,ψ-carotene) into β- or ε-ionone rings in the crucial bifurcation step of carotenoid biosynthesis. Among all carotenoid constituents, β-carotene (β,β-carotene) is found in all photosynthetic organisms, except for purple bacteria and heliobacteria, suggesting a ubiquitous distribution of lycopene β-cyclase activity in these organisms. In this work, we isolated a gene (BfLCYB) encoding a lycopene β-cyclase from *Bangia fuscopurpurea*, a red alga that is considered to be one of the primitive multicellular eukaryotic photosynthetic organisms and accumulates carotenoid constituents with both β- and ε-rings, including β-carotene, zeaxanthin, α-carotene (β,ε-carotene) and lutein. Functional complementation in *Escherichia coli* demonstrated that BfLCYB is able to catalyze cyclization of lycopene into monocyclic γ-carotene (β,ψ-carotene) and bicyclic β-carotene, and cyclization of the open end of monocyclic δ-carotene (ε,ψ-carotene) to produce α-carotene. No ε-cyclization activity was identified for BfLCYB. Sequence comparison showed that BfLCYB shares conserved domains with other functionally characterized lycopene cyclases from different organisms and belongs to a group of ancient lycopene cyclases. Although *B. fuscopurpurea* also synthesizes α-carotene and lutein, its enzyme-catalyzing ε-cyclization is still unknown.

Keywords: *Bangia fuscopurpurea*; red algae; lycopene cyclase; carotenoid; metabolism

1. Introduction

Carotenoids are one of the largest groups of natural pigments in land plants, algae, bacteria including cyanobacteria and photosynthetic bacteria, archaea, fungi and animals [1,2]. More than 750 carotenoids, not including their cis- and trans- isomers as distinct compounds, have been isolated and characterized from natural sources [3]. Carotenoids not only serve as essential pigments for photosynthesis and photoprotection in photosynthetic organisms, but also comprise many of the yellow, orange and red colors in flowers and fruits that attract pollinators and seed dispersers in higher plants [4]. Abscisic acid and strigolactones are also derivatives of carotenoids [5].

Higher plants generally have very similar leaf carotenoid profiles, with lutein, β-carotene (β,β-carotene), violaxanthin and neoxanthin being the major components. They may also accumulate specific carotenoids, such as capsanthin in pepper fruits [6], β-carotene in carrot tubers [7] and astaxanthin in *Adonis aestivalis* flowers [8]. In the past 20 years, most of the metabolic reactions in the biosynthesis...
of carotenoids have been deciphered, thanks to the cloning and characterization of genes encoding metabolic enzymes from different organisms [9]. In plants, the plastidic methylethylthiol phosphate (MEP) pathway utilizes glyceradehyde-3-phosphate and pyruvate to synthesize the 5-carbon (C₅) isopentenyl diphosphate (IPP) and its isomer dimethylallyl diphosphate (DMAPP). A geranylgeranyl diphosphate (GGPP, C₂₀) synthase (CrtE) catalyzes the condensation of these two substrates to produce GGPP as a precursor for the biosynthesis of carotenoids and other diterpenoids. Phytoene synthase (CrtB) utilizes two molecules of GGPP to produce phytoene and thus directs the metabolic flux from GGPP into carotenoid metabolic pathway. Phytoene is then converted to acyclic lycopene (Δ,Δ-carotene) by the combination of desaturases CrtP and CrtQ in eukaryotes, but only CrtI in prokaryotes [2,10].

Cyclization of lycopene by lycopene cyclases (LCY) is a crucial bifurcation step in the carotenogenic pathway (Figure 1). Each open end of lycopene can be cyclized by lycopene β-cyclase (LCYB) or ε-cyclase (LCYE) to produce β- or ε-ionone rings, respectively (Figure 1). The formation of two β-ionone rings results in the production of β-carotene and its derivatives (e.g., zeaxanthin and neoxanthin), whereas the combination of β- and ε-ionone rings forms α-carotene (β,ε-carotene) and its derivate lutein (Figure 1). Carotenoids with two ε-ionone rings are uncommon in most plants, with luctucaxanthin in lettuce (Lactuca sativa) being one of the exceptions [11]. Carotenoids with at least one unsubstituted β-ionone ring can be utilized by the human body to produce vitamin A. Therefore, an increase in the activity of LCYB can direct the metabolic flux from lycopene to the production of this group of so-called provitamin A carotenones to increase the nutritional value of crops [12,13].

Cumulative studies on carotenoids in red algae have uncovered broad compositional diversity [2,14–16]. Red algae can also be divided into three groups according to their specific carotenoid compositions [15,16]. Although carotenoid constituents with an ε-ionone ring are missing in some primitive species, β-carotene is found in all red algae, suggesting a ubiquitous distribution of the β-cyclization activity in carotenoid biosynthesis [16,17]. LCYB of Cyanidioschyzon merolae is also the only LCY gene from the entire red algal phylum that has been characterized [16,17].

The red algal genus Bangia Lyngb., although probably not monophyletic, is among the earliest multicellular eukaryotic photosynthetic organisms [18]. Carotenoids with both β- and ε-rings, including β-carotene, zeaxanthin, α-carotene and lutein, are accumulated in Bangia [15,16]. In this work, we isolated a gene encoding an LCY from Bangia fuscospurpurae, and functional characterization revealed that this enzyme has only β-cyclization activity.

2. Results

2.1. BfLCYB Is an Intron-Less LCY Gene

From the alignment of functionally characterized LCYs from different organisms, we designed two pairs of degenerate primers. Nested PCR using these primers resulted in amplicons of different
we successfully amplified a 2218 bp genomic DNA fragment and a 1932 bp complementary DNA (cDNA) encoding a putative protein with 643 amino acid residues, designated BfLCYB in this work (Figure 2A). Sequence alignment demonstrated that BfLCYB does not possess any introns. BfLCYB shares 38% sequence identity with the characterized LCYB from C. merolae [17]. The sequence of BfLCYB has been deposited in GenBank under the accession number KX943552.

C. merolae shares 38% sequence identity with the characterized LCYB from Dunaliella salina (Ds-1, ACA34344.1; Ds-2, ANY98896.1), Oryza sativa (Os-1, XP_015622198.1), Arabidopsis thaliana (At-1, NP_187634.1), and Adonis aestivalis (Aa-1, AAK07430.1), and other conserved LCYs from A. aestivalis (At-2, NP_200513.1) and LCYBs from Dunaliella salina (Ds-1, ACA34344.1; Ds-2, ANY98896.1), Cyanidioschyzon merolae (Cm-1, XP_005536481.1) and Bangia fuscopurpurea (Bf-1, KX943552) were compared.

Figure 2. Sequence analysis of BfLCYB. (A) BfLCYB complementary DNA (cDNA) sequence and its deduced amino acid sequence. Sequence of the predicted N-terminal chloroplast transit peptide is in red. (B–F) Sequence comparison of the NAD(P)/FAD-binding domain (B) and other conserved domains (C–F). (G) Sequence comparison of the six-amino-acid region (in bold) that determines the ring numbers of LCYE and flanking regions. The conserved domains in BfLCYB are in blue and the conserved domains in other LCYs are in red.
Five conserved domains, including an NAD(P)/FAD-binding domain, were previously reported for functionally characterized LCYs [2,10]. These domains were also identified in BfLCYB (Figure 2). However, the NAD(P)/FAD-binding domain in BfLCYB is interrupted by an apparent insertion of 12 additional amino acid residues (Figure 2B). Cunningham and Gantt also identified a six-amino-acid region in LCY that contributes to the determination of ring numbers in lycopene ε-cyclization products [11]. We found that this region is located immediately downstream from the fifth conserved domain in LCYs (Figure 2F,G). However, in BfLCYB, this region has much lower similarity to other LCYs than either its upstream or downstream flanking regions (Figure 2G).

2.2. BfLCYB Localizes in Chloroplasts

From online analysis using TargetP, ChloroP and YLoc, BfLCYB was predicted to have a chloroplast localization [19–21]. Its N-terminal chloroplast transit peptide predicted by TargetP and ChloroP is in square brackets in Figure 2A. To experimentally confirm its localization, the open reading frame (ORF) of BfLCYB was fused in frame to the 5′-end of the yellow fluorescent protein (YFP) in pA7-YFP. Due to the lack of a method to stably transform Bangia, we transiently expressed the BfLCYB-YFP fusion protein in Arabidopsis thaliana leaf protoplasts [22]. The YFP signal of the fusion protein was solely found in chloroplasts and merged nicely with chlorophyll autofluorescence, thus confirming its predicted chloroplast localization (Figure 3).

![Figure 3. Subcellular localization of BfLCYB. The BfLCYB yellow fusion protein (BfLCYB-YFP) was transiently expressed in Arabidopsis thaliana protoplasts, and observed under a confocal microscope. Signals from the fusion protein (A) and chlorophyll autofluorescence (B) were merged (C) for a comparison with the protoplast observed under bright field (D). The transient expression experiment was repeated independently five times and only one representative protoplast is shown here. Scale bar, 5 μm.](image)

2.3. BfLCYB Is a Bicyclic Lycopene β-Cyclase

To determine the enzymatic activity of BfLCYB, we used a bacterial pigment complementation system [23]. The plasmid pAC-LCY contains the genes from Pantoea for geranylgeranyl diphosphate synthase (CrtE), phytoene synthase (CrtB) and phytoene desaturase (CrtI). Escherichia coli transformed with pAC-LCY can utilize endogenous IPP and DMAPP to produce lycopene, which turns the cells pink (Figure 4A). When we co-transformed E. coli with both pMAL-BfLCYB and pAC-LCY, the positive transformants harboring both plasmids had an orange color (Figure 4B). High-performance liquid chromatography (HPLC) analysis revealed an accumulation of both the monocyclic γ-carotene (β,β-carotene) and the bicyclic β-carotene in these orange cells (Figure 4B). We did not find accumulation of either ε-carotene (ε,ε-carotene) or δ-carotene (ε,β-carotene) in the co-transformed bacteria. These results suggested that BfLCYB is able to cyclize one or both ends of lycopene into β-ionone rings but not ε-ionone rings.

We further confirmed the function of BfLCYB by co-transforming pMAL-BfLCYB with pAC-EPSILON. In addition to the genes carried by pAC-LCY, pAC-EPSILON carries the gene from lettuce for LCYE, which catalyzes the ε-cyclization at both ends of lycopene [11]. E. coli cells transformed with pAC-EPSILON accumulated ε-carotene, as expected (Figure 4C). When this plasmid was co-transformed with pMAL-BfLCYB, HPLC analysis revealed the accumulation of both α-carotene and β-carotene, together with a significantly reduced level of ε-carotene and a small
amount of γ-carotene (Figure 4D). This demonstrated that BfLCYB can also β-cyclize the open end of monocyclic δ-carotene.

**Figure 4.** Functional characterization of BfLCYB. (A) Production of lycopene in *Escherichia coli* cells transformed with pAC-LCY. (B) Production of γ-carotene and β-carotene in *E. coli* cells transformed with both pAC-LCY and pMAL-BfLCY1. (C) Production of ε-carotene in *E. coli* cells transformed with pAC-EPSILON. (D) Production of α-, ε-, γ- and β-carotene in *E. coli* cells transformed with both pAC-EPSILON and pMAL-BfLCYB. (E) UV-visible absorption spectra of different carotenoid products. Insets in (A, B) showed different colors of the pelleted bacterial cells accumulating lycopene and β-carotene, respectively.

2.4. BfLCYB Is an Ancient Type of Lycopene Cyclase in Plants

Using sequences of functionally characterized LCYs, we identified LCY homologs in heterokonts and red algae of which full genomes have been sequenced. For heterokonts, in the genome of either *Ectocarpus siliculosus* (brown alga) or *Phaeodactylum tricornutum* (diatom), there is only a single gene encoding an LCY homolog (Figure 5A). However, for the multicellular red alga *Chondrus crispus*, our search identified two genes for putative LCYs in its genome (Figure 5A). Our phylogenetic analysis divided LCYs and their homologs into five major clades. Clade I comprises members from single-cell red algae, including the LCYB from *Cyanidioschyzon merolae* and a homolog from *Galdiera sulphuraria*. Clade II has members from multicellular red algae, including BfLCYB in this...
study and the two homologs from Chondrus crispus. Clade IV has LCY homologs from heterokonts. The other two clades, III and V, are LCYBs and LCYEs, respectively, from the green lineage (green algae and land plants) (Figure 5A).

A recently study by Blatt et al. [24] reported a gene (OluLCY) encoding a lycopene β-cyclase/lycopene ε-cyclase/light-harvesting complex-fusion protein from the green alga Ostreococcus lucimarins. Our sequence analysis of this gene identified two coding regions (OluLCY-B and OluLCY-E) that are tandemly arranged, separated by a short intron (Figure 5B). In our phylogenetic tree, peptides encoded by these two regions are grouped with functionally characterized LCYBs and LCYE s, respectively (Figure 5B).

3. Discussion

In green lineage organisms (green algae and land plants), carotenoids with both β- and ε-rings are found, and separate LCYBs and LCYEs catalyzing the corresponding cyclization steps are also characterized from different species [1,25]. Our phylogenetic analysis clustered LCYBs and LCYE s

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**Figure 5.** Phylogenetic analysis of plant LCYs. (A) Sequences of the functionally characterized LCYs from Zea mays (LCYB, NP_001169155.1; LCYE, NP_001146840.1), Oryza sativa (LCYB, XP_015627234.1; LCYE, XP_015622198.1), Arabidopsis thaliana (LCYB, NP_187634.1; LCYE, NP_200513.1), Ipomoea batatas (LCYB, AGL44392.1; LCYE, BAW34178.1), Adonis aestivalis (LCYB, AAK07430.1; LCYE, AAK07432.1), Marchantia polymorpha (LCYB, BAO27799.1; LCYE, BAO27800.1), Chromochloris zofingiensis (LCYB, CBH31263.1; LCYE, CCG06343.1), Dunaliella salina (LCYB1, ACA34344.1; LCYE2, ANY98896.1), Ostreococcus lucimarinus (LCYB, NP_001169155.1; LCYE, NP_001146840.1), and their homologs from Ectocarpus siliculosus (CBB78004.1), Phaeodactylum tricornutum (XP_002176612.1), Chondrus crispus (1, XP_005713399.1; 2, XP_005715292.1) and Galdieria sulphuraria (XP_005708098.1) are analyzed using MEGA 6. Scale bar, 10% sequence divergence. (B) Structure of the Ostreococcus lucimarins LCY gene that has two tandemly arranged regions for LCYB (OluLCY-B) and LCYE (OluLCY-E) activities.
from these organisms into separate clades (IV and V, Figure 5), showing a divergence of lycopene β- and ε-cyclase activities (Clades IV and V, Figure 5A) before the origination of the green lineage. This is also supported by the identification of the OluLCY gene, which has two tandemly arranged regions for peptides with LCYB and LCYE activities, respectively, suggesting probable gene duplication and divergence.

In red algae and heterokonts, although their carotenoid constituents have been studied, there is not sufficient information on their LCYs [2,26]. In both single-cell red algae and heterokonts, only β-carotene and its derivatives were found. Consistently, LCYs from *Cyanidioschyzon merolae* possess only β-cyclization activity, and studies in diatoms found no sequences showing similarity to LCYE genes either [16,26–28]. Therefore, it is reasonable to postulate that members of Clades I and IV in Figure 5A are all LCYBs.

For multicellular red algae, which accumulates carotenoids with both β- and ε-rings, studies on their LCYs are lacking. In this work, we studied lycopene cyclization in *Bangia fuscopurpurea*, which is considered to be one of the simplest macrophytic red algae. Our molecular cloning and functional characterization results demonstrated that *B. fuscopurpurea* has a functional LCYB (BfLCYB), which is capable of catalyzing β-cyclization at both ends of the acyclic lycopene or at either end of the monocyclic γ-carotene or δ-carotene. By scanning the genome sequence of another multicellular red alga, *Chondrus crispus*, we found two homolog genes for LCYs. These remind us of the possible existence of another as-yet-unidentified LCYE homolog responsible for ε-cyclization in *B. fuscopurpurea*. Cloning and functional characterization of this second LCY from *B. fuscopurpurea* should help to decipher carotenoid metabolism and its evolution in red algae.

### 4. Materials and Methods

#### 4.1. Material and Growth Condition

Thalli of *Bangia fuscopurpurea* (#Bangia-MC1F, Jiangsu Institute of Oceanology and Marine Fisheries, Nantong, Jiangsu, China) were initially collected from cultivated populations in Putian, Fujian Province, China (24°59′ N, 118°48′ E) [29] and then subcultured in sterilized seawater in our laboratory. Ambient air filtered through a 0.45 µm filter was gently supplied to the culture. The growth conditions were 16 °C with a 12 h/12 h light/dark regime under 80 µmol m−2 s−1 irradiance. Seawater was changed every week.

#### 4.2. Sequence Analysis

Sequences of functionally characterized LCYs from *Zea mays*, *Oryza sativa*, *Arabidopsis thaliana*, *Ipomoea batatas*, *Adonis aestivalis*, *Marchantia polymorpha*, *Chromochloris zofingiensis*, *Dunaliella salina*, *Ostreococcus lucimarinus* and *Cyanidioschyzon merolae* were downloaded from GenBank, and their homologs in *Ectocarpus siliculosus*, *Phaeodactylum tricornutum*, *Chondrus crispus* and *Galdieria sulphuraria* were searched in GenBank using the blastP algorithm with the sequences of known LCYs as queries. Sequence alignment was performed using ClustalW in MEGA 6 [30]. A bootstrap (1000 replicates) neighbor-joining phylogenetic tree was also generated using MEGA 6.4.3. Molecular manipulation.

Total RNA was isolated from *Bangia fuscopurpurea* material as previously described [31] and treated with DNase (Promega, Madison, WI, USA) to eliminate any genomic DNA contamination. For rapid amplification of cDNA ends (RACE), the SMARTer RACE cDNA Amplification Kit (TaKaRa, Shiga, Japan) was used to synthesize a full-length cDNA pool. According to the alignment of the functionally characterized LCYs, two pairs of degenerate primers (DF1/DR1, DF2/DR2, sequences of all primers used in this work are listed in Table 1) were designed from the conserved regions [32], and used for amplifying a corresponding fragment from the *B. fuscopurpurea* cDNA pool. After two rounds of nested PCR, amplified products were subcloned and sequenced by GenScript (Nanjing, China). The sequences of the cloned fragments were queried against GenBank using the tblastX algorithm to analyze their similarities to known or predicted LCY genes.
Table 1. Primers used in this work.

<table>
<thead>
<tr>
<th>Primer</th>
<th>Sequence (5′-3′)</th>
</tr>
</thead>
<tbody>
<tr>
<td>DF1</td>
<td>MMNAAYTAYGGNKKNTGGBWNAGYR</td>
</tr>
<tr>
<td>DR1</td>
<td>GGRTRGNACSDADNHSNCGNGYNSECNCCNANWNSC</td>
</tr>
<tr>
<td>DF2</td>
<td>ATGBTNYTNTAGGAYKDNMNGGA</td>
</tr>
<tr>
<td>DR2</td>
<td>RNVTNNCCNCCANAYNGDATNWW</td>
</tr>
<tr>
<td>RR1</td>
<td>GCGCCCTCAATGGGGATCAGACAAAACTC</td>
</tr>
<tr>
<td>RR2</td>
<td>CAGACAAAAACTCCTCAGGCAGCATCGG</td>
</tr>
<tr>
<td>RR3</td>
<td>CACCCGCCTCAGGGCATGGAAGG</td>
</tr>
<tr>
<td>RF1</td>
<td>GGTGCTGATGGACTACCGGGATGG</td>
</tr>
<tr>
<td>RF2</td>
<td>GGGATGGGCACATGCAGGGGGAC</td>
</tr>
<tr>
<td>RF3</td>
<td>GGACGCCGCGGGGCGGGCGAGTC</td>
</tr>
<tr>
<td>HF</td>
<td>ATGGAGGCGTTCATCCCCGCTC</td>
</tr>
<tr>
<td>ER</td>
<td>CTACGCCTCGTCCTCTGAGGTG</td>
</tr>
<tr>
<td>gHF</td>
<td>AATCAGTCACTTCTCGTGATCTTCTT</td>
</tr>
<tr>
<td>gER</td>
<td>CCGCCGAAACGCGCTACCCCT</td>
</tr>
<tr>
<td>SpeI-LCY-F</td>
<td>GACTAGTATGGAGGCCTTCATCCGCCCTC</td>
</tr>
<tr>
<td>SpeI-LCY-R</td>
<td>CGGACTAGCTACGGCTCTGTCCTGAGCAGGG</td>
</tr>
<tr>
<td>BamHI-LCY-F</td>
<td>GGATCGATGGAGGCGCTTCATCCCCGCCCTC</td>
</tr>
<tr>
<td>BamHI-LCY-R</td>
<td>GGAGATCCCTACGCGCTTCGTCTGAGGAGGG</td>
</tr>
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</table>

1 Codes for degenerate nucleotides are M (A/C), R (A/G), W (A/T), S (C/G), Y (C/T), K (G/T), V (A/C/G), H (A/C/T), D (A/G/T), B (C/G/T) and N (any nucleotide); 2 Restriction sites are underlined.

Using the sequences that had high similarity to known LCYs, three pairs of primers (RF1/RR1, RF2/RR2 and RF3/RR3) were designed for 5′- and 3′-RACEs, respectively, using the SMARTer RACE kit. After three rounds of nested PCR, two overlapping fragments were amplified from the cDNA pool, subcloned and sequenced. The sequences were assembled into a putative cDNA sequence that contained both translation initiation and stop codons. We then designed an additional pair of primers (HF and ER) from the assembled sequence to amplify the full-length ORF from the cDNA pool for confirming the existence of this transcript (named BfLCYB).

To study the genomic sequence of BfLCYB, genomic DNA was extracted according to Yang et al. [31]. Primers HF and ER were used to amplify its coding region. A Genome Walking Kit (TaKaRa) was used to amplify the 5′ and 3′ flanking sequences. All amplicons were purified, subcloned, sequenced and assembled. Two primers (gHF and gER) were used to amplify from the genomic DNA for confirming the assembled genomic DNA sequence of BfLCYB.

4.3. Subcellular Localization

Prediction of the subcellular localization of BfLCYB was performed using TargetP [19], YLoc [20] and ChloroP [21].

For studying the subcellular localization of BfLCYB, protoplasts isolated from Arabidopsis thaliana mesophyll were transiently transformed [22]. The full-length ORF of BfLCYB was amplified with primers HF-Spel and ER-Spel to incorporate restriction sites. The amplicon was purified, digested with SpeI, and subcloned into pA7-YFP (kindly provided by Dr. Hongquan Yang). The transformed protoplasts were cultured in 24-well plates at room temperature for 16 h in the dark. Fluorescence signal from the BfLCYB-YFP fusion protein was observed using a confocal laser scanning microscope (Olympus Fluoview FV1000, Tokyo, Japan). The transient expression experiments were repeated independently at least three times.

4.4. Functional Characterization

To assess the catalytic activity of BfLCYB, we used the pigment complementation system [23]. Two vectors, pAC-LCY and pAC-EPSILON were provided by Dr. Cunningham. The full-length ORF of BfLCYB was amplified with primers HF-BamHI and ER-BamHI to incorporate restriction sites.
The amplified product was subcloned into pMAL-C5X (New England BioLabs, Ipswich, MA, USA) to produce pMAL-BfLCYB. pMAL-BfLCYB was co-transformed with pAC-LCy, or pAC-EPSILON, into E. coli BL21(DE3) cells (New England BioLabs, Ipswich, MA, USA). After overnight growth at 37 °C on Luria-Bertani (LB), plants containing 34 µg mL⁻¹ chloramphenicol and 100 µg mL⁻¹ ampicillin, positive co-transformed single colonies were picked and used to inoculate 8 mL LB medium containing the same antibiotics. One mL of the overnight culture was transferred to 100 mL of the same medium and grown at 37 °C until reaching an OD600 of 0.30. Protein expression was induced by adding isopropyl β-thiogalactopyranoside (IPTG) to a final concentration of 50 µmol L⁻¹. After growing for 16 h at 18 °C, bacterial cells were harvested by centrifugation and resuspended in 80% (v/v) acetone to extract the pigments.

4.5. Carotenoid Analysis

Separation of carotenoid components was performed by reverse-phase HPLC [33]. A Waters 2695 separation module and 2998 photodiode array detector (PDA) were used with a Spherisorb ODS2 column (5 µm, 4.6 mm × 250 mm, Waters, Milford, MA, USA). A 45 min gradient of ethyl acetate (0% to 100%) in acetonitrile-water-triethylamine (9:1:0.01, v/v/v) at a flow rate of 1 mL min⁻¹ was used, and the absorbance of the eluent at 440 nm was monitored [34]. The identity of the carotenoid in each peak was further confirmed by their UV/visible spectra recorded by the PDA detector.

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Author Contributions: T.-J.C., Y.-Y.D. and S.L. conceived and designed the experiments; T.-J.C., X.-Q.H., Y.-Y.D., Y.-Y.Q. and Z.Z. performed the experiments; T.-J.C., X.-Q.H. and S.L. analyzed the data; S.L. wrote the paper.

Conflicts of Interest: The authors declare no conflict of interest.

References


