Enhancement of carotenoids biosynthesis in *Chlamydomonas reinhardtii* by nuclear transformation using a phytoene synthase gene isolated from *Chlorella zofingiensis*

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**Abstract** The isolation and characterization of the phytoene synthase gene from the green microalga *Chlorella zofingiensis* (*CzPSY*), involved in the first step of the carotenoids biosynthetic pathway, have been performed. *CzPSY* gene encodes a polypeptide of 420 amino acids. A single copy of *CzPSY* has been found in *C. zofingiensis* by Southern blot analysis. Heterologous genetic complementation in *Escherichia coli* showed the ability of the predicted protein to catalyze the condensation of two molecules of geranylgeranyl pyrophosphate (GGPP) to form phytoene. Phylogenetic analysis has shown that the deduced protein forms a cluster with the rest of the phytoene synthases (PSY) of the chlorophycean microalgae studied, being very closely related to PSY of plants. This new isolated gene has been adequately inserted in a vector and expressed in *Chlamydomonas reinhardtii*. The over-expression of *CzPSY* in *C. reinhardtii*, by nuclear transformation, has led to an increase in the corresponding *CzPSY* transcript level as well as in the content of the carotenoids violaxanthin and lutein which were 2.0- and 2.2-fold higher than in untransformed cells. This is an example of manipulation of the carotenogenic pathway in eukaryotic microalgae, which can open up the possibility of enhancing the productivity of commercial carotenoids by molecular engineering.

**Keywords** Carotenoids · *Chlorella zofingiensis* · Phytoene synthase · Transgenic microalgae · *Chlamydomonas reinhardtii*

**Introduction**

Carotenoids are isoprenoids synthesised by all photosynthetic organisms and by some fungi and non-photosynthetic bacteria. In photosynthetic organisms, carotenoids bind to integral membrane proteins of the thylakoid where they participate in light harvesting and in the protection of the photosynthetic apparatus against the photo-oxidative damage (Frank and Cogdell 1996; Varkonyi et al. 2002). Carotenoids are high-value compounds, being used as nutraceuticals and as natural dyes and additives in food, feed, aquaculture, and cosmetic industries. They are considered effective agents for the prevention of a variety of age-related, degenerative and chronic diseases such as cataracts, macular degeneration, atherosclerosis, cardiovascular diseases, and some types of cancer (Dweyer et al. 2001; Demming-Adams and Adams 2002; Guerin et al. 2003; Olmedilla et al. 2003; Stahl and Sies 2005).

In chloroplasts of plants and algae, the carotenoids precursor, geranylgeranyl pyrophosphate (GGPP), is formed by the action of GGPP synthase from isopentenyl pyrophosphate and dimethylallyl pyrophosphate, which are derived from deoxyxylulose 5-phosphate pathway. GGPP is not only the precursor for carotenoids but also participate in
the synthesis of other terpenoid compounds, such as phytol, plastoquinones, and tocopherols. The condensation of two GGPP molecules produces the first carotene, phytoene, catalyzed by PSY (Fig. 1). Phytolene is desaturated by phytolene and \( \zeta \)-carotene desaturases (PDS and ZDS) and isomerized by 15-cis-\( \zeta \)-carotene isomerase (Z-ISO) (Chen et al. 2010) and carotene isomerase (CRTISO) to form the linear all trans-lycopene. The cyclation of lycopene by lycopene \( \epsilon \)-cyclase (LCYe) and lycopene \( \beta \)-cyclase (LCYb) introduces \( \epsilon \) - and \( \beta \)-ionone end groups, respectively, yielding \( \alpha \) - and \( \beta \)-carotenoids. \( \alpha \) - and \( \beta \)-carotene are hydroxylated into lutein and zeaxanthin, respectively. Two P450 hydroxylases (P450b-CHY and P450e-CHY) are active in \( \alpha \)-carotene hydroxylation. P450b-CHY is also active in \( \beta \)-carotene hydroxylation, together with two non-nucleotide di-iron hydroxylases (CHYb). Zeaxanthin epoxidase (ZEP) and violaxanthin de-epoxidase (VDE) catalyze the interconversion of zeaxanthin and violaxanthin (Kim et al. 2009). A limited number of organisms including some green algae as Haematococcus pluvialis and Chlorella zofingiensis can synthesize astaxanthin from \( \beta \)-carotene by the action of a ketolase/oxygenase (BKT) and the hydroxylase (CHYb) (Fan et al. 2009; Sandmann et al. 2006). A limited number of plants and microalgae. Phytoene synthase (PSY) catalyzes the first step in the carotenoid specific pathway, which leads the carbon flux towards carotenoids and xanthophylls production. IPP isopentenyl pyrophosphate, DMAPP dimethylallyl pyrophosphate, GGPP geranylgeranyl pyrophosphate synthase, PDS phytoene desaturase, Z-ISO 15-cis-\( \zeta \)-carotene isomerase, ZDS \( \zeta \)-carotene desaturase, CRTISO carotene isomerase, LCYb lycopene \( \beta \)-cyclase, LCYe lycopene \( \epsilon \)-cyclase, P450b-CHY cytochrome P450 \( \beta \)-hydroxylase, P450e-CHY cytochrome P450 \( \epsilon \)-hydroxylase, CHYb carotene \( \beta \)-hydroxylase, BKT \( \beta \)-carotene oxygenase, ZEP zeaxanthin epoxidase, VDE violaxanthin de-epoxidase

![Fig. 1 Schematic diagram of the carotenoid biosynthetic pathway in plants and microalgae. Phytolene synthase (PSY) catalyzes the first step in the carotenoid specific pathway, which leads the carbon flux towards carotenoids and xanthophylls production. IPP isopentenyl pyrophosphate, DMAPP dimethylallyl pyrophosphate, GGPP geranylgeranyl pyrophosphate synthase, PDS phytoene desaturase, Z-ISO 15-cis-\( \zeta \)-carotene isomerase, ZDS \( \zeta \)-carotene desaturase, CRTISO carotene isomerase, LCYb lycopene \( \beta \)-cyclase, LCYe lycopene \( \epsilon \)-cyclase, P450b-CHY cytochrome P450 \( \beta \)-hydroxylase, P450e-CHY cytochrome P450 \( \epsilon \)-hydroxylase, CHYb carotene \( \beta \)-hydroxylase, BKT \( \beta \)-carotene oxygenase, ZEP zeaxanthin epoxidase, VDE violaxanthin de-epoxidase](image)

In the present work, we report the isolation and characterization of the PSY gene from C. zofingiensis, as well as its ability to convert two GGPP molecules into phytolene. Moreover, this novel gene has been inserted in an adequate vector and expressed in C. reinhardtii. This is an example of the overexpression of an exogenous gene (PSY) in an eukaryotic microalgae, which can be an interesting tool for the massive production of carotenoids in transgenic microalgae by genetic engineering.
Materials and methods

Strains and culture conditions

The green microalga strain C. zofingiensis SAG 211–14 was obtained from the Culture Collection of Göttingen University (SAG, Germany). This microalga was grown photoautotrophically in Arnon medium (Arnon et al. 1974) modified to contain 4 mM K$_2$HPO$_4$ and 20 mM NaNO$_3$, at 25°C under continuous illumination (50 μmol photons m$^{-2}$ s$^{-1}$). The light intensity was measured at the surface of the flasks using a LI-COR quantum sensor (model LI-1905B, Li-Cor, Inc., Lincoln, NE, USA). The liquid cultures were continuously bubbled with air supplemented with 1% (v/v) CO$_2$ as the only source of carbon. C. reinhardtii cell-wall-deficient strain 704 was kindly provided by Dr. Roland Loppes (Loppes et al. 1999) and cultured mixotrophically in either liquid or agar solidified Tris-acetate phosphate (TAP) medium (Gorman and Levine 1965) at 25°C under a continuous irradiance of 50 μmol photons m$^{-2}$ s$^{-1}$. Escherichia coli DH5α and BL21 (DE3) strains were used as the hosts for DNA manipulation and for heterologous expression of PSY gene, respectively.

For the analysis of transformants, cells were grown in Erlenmeyer flasks of 100 mL capacity at 25°C under continuous illumination (50 μmol photons m$^{-2}$ s$^{-1}$) in liquid TAP medium.

Genomic DNA and RNA isolation and cDNA preparation

DNA and total RNA were isolated using DNaseasy Plant Mini Kit and RNasey Plant Mini Kit (Qiagen, Düsseldorf, Germany), respectively. For genomic DNA isolation for PCR screening of transformants from C. reinhardtii, a loopful of cells was scrapped from a plate and resuspended in 150 μL of cold distilled water and 350 μL of a buffered solution containing 50 mM Tris–HCl, pH 8, 0.3 M NaCl, 5 mM EDTA, and 2% SDS. The DNA isolation was performed by phenol–chloroform–isoamyl alcohol (50:48:2) extraction and selective precipitation with ethanol, according to previously described protocols (Anwaruzzaman et al. 2004). For quantitative real-time PCR analysis (qRT-PCR), first-strand cDNA synthesis was obtained from total RNA treated with DNase as recommended by the manufacturer, by using the SuperScript First-Strand Synthesis System (Invitrogen, Barcelona, Spain) primed with oligo(dT)$_{18}$ according to the manufacturer’s instructions.

Cloning of C. zofingiensis PSY cDNA and genomic gene

For isolating the cDNA clone coding for the C. zofingiensis PSY homologue, amino acid sequences deduced from previously cloned PSY genes from different kinds of algae, cyanobacteria, and plants were aligned. Highly conserved regions were identified, and different pairs of degenerated primers were designed. The PCR product was cloned in the pGEM-T vector (Promega, Madison, WI, USA) according to the manufacturer’s manual and then sequenced. The cDNA fragment obtained corresponding to partial PSY clone provided sequence information for the designing of gene-specific primers for amplification of 5’ and 3’ cDNA ends by RACE-PCR. All reactions were performed with kits according to the manufacturer’s instructions (Smart RACE cDNA Amplification Kit, Clontech, Mountain View, CA, USA). 5’ and 3’ RACE products were cloned into pGEM-T vector and sequenced. Specific primers were synthesized for genomic DNA amplification based on cDNA sequence. The primers sets used in this study are listed in Table 1.

Nucleotide sequence accession numbers

The CzPSY cDNA and genomic DNA sequences have been registered in the EMBL database (EMBL, FR670783 and EMBL, FR670784, respectively).

Sequencing and phylogenetic analysis

The deduced amino acid sequence of the C. zofingiensis PSY was compared with other PSY sequences of algae, plants, cyanobacteria, and bacteria. The sequence analysis and alignments were done with CLUSTAL W software. The deduced amino acid sequence was subjected to the ProtParam application at the ExPASy server (Gasteiger et al. 2005) for physical and chemical parameters, program Predotar v. 1.03 (Caboche 2003) for the prediction of possible plastid localization, and ChloroP 1.1 server (Emanuelsson et al. 1999) for the identification of a chloroplast transit peptide. For transmembrane analysis, ProtScale (Gasteiger et al. 2005) and TopPred (von Heijne 1992) servers were used. The construction of a phylogenetic tree was performed in MEGA4 (Tamura et al. 2007) using the UPGMA method.

Southern blot analysis

Genomic DNA was digested with HincII and HindIII, which showed one and no recognition site, respectively, in the probed region of the PSY gene. The probe was prepared by amplifying genomic DNA with the primers Czpsy-S-F and Czpsy-S-R, resulting in a 752-bp fragment of CzPSY gene. The digested DNA was transferred to a Hybond-N membrane (GE Healthcare, Little Chanfont, UK) by capillary transfer and hybridized with the $^{32}$P labelled DNA probe at both low and high stringency overnight. After hybridization, the radioactivity of the membrane was monitored by the Cyclone Phosphor System (Perkin-Elmer, Waltham, MA, USA).
Table 1 Nucleotide sequences of primer pairs used for PCR amplification

| Primer | Sequence (5′→ 3′) |
|--------|-------------------|
| Partial PSY fragment | |
| psy-1F | GAYATGATGARGGNATG |
| psy-1R | AARTTRCRTARTCRTT |
| 5′and 3′ RACE | |
| GSP-F | ATGAATTTAGTCAAGTCAGG |
| GSP-R | TTTGGGACATAAGCAGG |
| NGSP-F | CTGGATGAAGGCAAGG |
| NGSP-R | CCGGCCTTCTACCCAG |
| Genomic DNA amplification | |
| psy-2F | ACATGGGGGCGAATTTGTGTT |
| psy-2R | CCCGTTGCTGTATAAGTCAT |
| psy-3F | ATGCCGTCGTTACAGTGA |
| psy-3R | ACCGTGACTATCTTGGT |
| psy-4F | ACAGCAGCAGATAGAAGG |
| psy-4R | CTCTCAAACTTGAGTGAAC |
| psy-5F | TTGTTACATTTACCTGGAGT |
| psy-5R | AGCACGACATATAGGCTC |
| PCR for probe preparation | |
| Czpsy-S-F | ATGGATTAGTCAAGTACGG |
| Czpsy-S-R | TTTGGGCACATAAGCAGG |
| Genetic complementation-pQE-80L | |
| pQE-psy-F | ggtacctAGGGCGCTGTTGACACAG |
| pQE-psy-R | ggtacctTTAGTCTGTGTTGCACG |
| PCR for Chlamydomonas reinhardtii transformation | |
| CzpsyCr105-F | cgcctcggATAGGGCGCTGTTAGCACCCAG |
| CzpsyCr105-R | cgccgattCTAGTCTGTGTTGCACG |
| Czpsy expression | |
| RT-Czpsy-F | CACCGATTGTCAGATCCCA |
| RT-Czpsy-R | ACTAGTTGTTGTGCTAICT |
| CrPSY expression | |
| RT-Crpsy-F | CACCGTGCCAAGCAGCAGCAG |
| RT-Crpsy-R | CACCGGCAAGCAGCACCATC |
| CBLP expression | |
| RT-cblp-F | CGCCACCGAACCTCCACATCAAG |
| RT-cblp-R | CTAGGGCGCGCTGGCATTAC |

F′, forward; R, reverse

*a BamHI and KpnI sites (lowercases underlined) were added for cloning the gene into the corresponding cut sites of pQE-80L vector.

*b XhoI and EcoRI sites (lowercases underlined) were added for cloning the gene into the corresponding cut sites of pSI105 plasmid.

Functional analysis of CzPSY cDNA by heterologous expression in E. coli

The CzPSY ORF was amplified by PCR with the primers pQE-psy-F and pQE-psy-R, which were designed to contain BamHI and KpnI restriction sites, respectively, and cloned between the BamHI and KpnI restriction sites of the pQE-80L expression vector (Qiagen), resulting in plasmid pQE-CzPSY, which carries ampicillin resistance. Plasmid pAC-85b, a gift from Prof. Cunningham, contained the carotenoid pathway genes responsible for the synthesis of β-carotene except PSY gene (crtE, crtI, and crtY of Erwinia herbicola) (Cunningham and Gantt 2007). Transformation of E. coli BL21 (DE3) with either pAC-85b (carrying chloramphenicol resistance) or pAC-85b and one of the two plasmids, pQE-CzPSY or pQE-80L (empty vector), was made by electroporation. Transformed cells were plated on Luria–Bertani solid medium (Sambrook et al. 1989), supplemented with 100 μg mL⁻¹ ampicillin and/or 40 μg mL⁻¹ chloramphenicol, and grown at 37°C for 1 day. The inducer isopropyl-β-D-1-thiogalactopyranoside was added at a final concentration of 1 mM.

Chlamydomonas nuclear transformation

The complete coding region of CzPSY was amplified by PCR with primers CzpsyCr105-F and CzpsyCr105-R, which were designed to contain XhoI and EcoRI restriction sites respectively, and cloned into pGEM-T vector (Promega), which carries ampicillin resistance. The PCR product was digested with XhoI and EcoRI and cloned in the pSI105-Tp1 vector, resulting in the plasmid pSI105-Tp1-psy that was used to transform Chlamydomonas cells. The plasmid pSI105-Tp1 is based on the plasmid pSI104-PLK (León et al. 2007), derived from the pSI103 (Sizova et al. 2001), in which the aphVIII gene from Streptomyces rimosus, coding for an aminoglycoside 3′phosphotransferase that confers resistance to the antibiotic paromomycin, is expressed under the control of the strong constitutive promoters rbcS2 and hsp70A and terminated by the 3′ untranslated region of rbcS2. The construction pSI105-Tp1 also carries a second expression cassette driven by the same two constitutive promoters and terminator region. This second cassette carries the transit peptide of RuBiSCO small subunit (rbcS2) to target the final peptide into the chloroplast stroma and also carries a polylinker region in which the cDNA from C. zofingiensis PSY was subcloned in frame with the transit peptide sequence.

Nuclear transformation was carried out using the glass beads method of Kindle (1990) with some modifications (León et al. 2007). C. reinhardtii cells grown to about 10⁷ cells mL⁻¹ were harvested by centrifugation and resuspended in fresh TAP medium to obtain a 100-fold concentrated cell suspension. The concentrated cell suspension (0.6 mL) was added to a conical tube containing 0.3 g of sterile glass beads (Ø 0.4–0.6 mm), 0.2 mL of 20% polyethylene glycol (MW8000) and 1 μg of the desired plasmid. Cells were vortexed and resuspended in 50 mL of fresh sterile TAP medium where they were incubated in the dark overnight. After this incubation, cells were centrifuged.
and spread onto solid TAP medium with paromomycin (30 μg mL⁻¹). Transformed colonies were visible after 4 to 5 days.

Quantitative RT-PCR

The mRNA relative abundance of endogenous C. reinhardtii PSY and foreign C. zofingiensis PSY was examined by qRT-PCR on an IQ5 Real-Time PCR Detection System (BioRad, Hercules, CA, USA), according to Cordero et al. (2010). In each experiment, a series of standard dilutions containing a specific concentration of a PCR fragment or a cDNA template was amplified in 20 μL of reaction containing 1×SYBR Green PCR Master Mix (Quantimix Easy SYG kit, BioTools B&M Labs, Madrid, Spain) and corresponding primers for either PSY from C. zofingiensis or C. reinhardtii (Table 1). After heating at 95°C for 10 min, cycling parameters were: 40 cycles of 95°C for 30 s, 60°C for 30 s, and 72°C for 30 s. Finally, the specificity of the qRT-PCR products was confirmed by performing a melting temperature analysis at temperatures ranging from 55°C to 95°C at 0.5°C/min and also by electrophoresis on a 2% agarose gel. Data were captured as amplification plots. Transcription levels of the target genes were calculated from the threshold cycle by interpolation from the standard curve. To standardize the results, the complete experiments (RNA isolation, cDNA synthesis followed with qRT-PCR) were repeated twice independently, and the data were averaged.

Analytical methods

Cell concentration and dry weight determinations

Cell number was determined with a Neubauer hemocytometer. For dry weight measurements, aliquots (5 mL) of the cell culture were filtered through Whatman GF/C paper (Whatman plc, Kent, UK), washed three times, and dried at 80°C for 24 h.

Carotenoid extraction and HPLC analysis

Total pigments were extracted with 80% of acetone (v/v) according to León et al. (2005). Then the samples were centrifuged and analysed by HPLC using a Waters Spherisorb ODS2 column (4.6×250 mm, 5 μm particle size) (Waters, Mildford, MA, USA). The chromatographic method described by Cordero et al. (2010) was used. Pigments were eluted at a flow rate of 1.0 mL min⁻¹ and were detected at 440 nm using a Waters 2996 photodiode-array detector. Identification of carotenoids was achieved by comparison of the individual characteristic absorption spectrum and the retention time with known standards. Quantification was performed using a calibration curve generated with commercially available carotenoids standards from Sigma-Aldrich (St. Louis, MO, USA) and DHI (Holsholm, Germany).

Results

Isolation and characterization of the PSY gene and deduced protein from C. zofingiensis

Different pairs of degenerate primers were designed on the basis of the conserved motifs present in PSY from microalgae, cyanobacteria, and plants. A partial cDNA fragment of 1,337 bp was isolated by PCR amplification using degenerate primers (psy-1F and psy-1R) (Table 1). A complete BLAST homologous search in the Genbank database showed that this fragment had enough similarity with the PSY gene from other species, and provided sequence information for designing specific primers for rapid amplification of 5′ and 3′ cDNA ends (RACE-PCR).

This analysis generated a full-length cDNA of 2,944 bp, which contained an ORF of 1,263 bp, 310-nucleotides of 5′-untranslated region (UTR), and a long 3′ UTR of 1,340 nucleotides. A typical algal polyadenylation signal TGTAAA (Gruber et al. 1992) was present in the 3′ UTR at 18 nucleotides upstream from the beginning of the poly(A) tail. The predicted protein has 420 amino acids residues, with an estimated molecular weight of 47.64 kDa, a theoretical isoelectric point of 8.53 and an instability index of 48.41 (data obtained with ProtParam program).

The differences between the C. zofingiensis PSY gene and the cDNA sequence were compared and revealed the presence of four exons and three introns (Fig. 2).

To determine the copy number of PSY gene in the genome of C. zofingiensis, genomic DNA was digested with two different restriction enzymes (either HinCl or HindIII) and subjected to Southern blot analysis at different conditions of stringency. Using a 752-bp fragment of CzPSY as a probe, strong hybridization signals have been obtained with both digestions. The digestion with HinCl enzyme, which cuts once inside the probe sequence, showed two bands, while digestion with HindIII, with no restriction site in the probe, exhibited only one band (Fig. 3). These results have suggested the presence of a single copy of the PSY gene in the genome of C. zofingiensis.

The BlastP search results demonstrated that the cloned CzPSY showed the highest overall homology sequence with other PSY from green algae, such as Dunaliella salina and Dunaliella sp. (identity, 74% and similarity, 84%) and
C. reinhardtii (identity, 70% and similarity, 84%). The GC content of the CzPSY coding region was 53%, which was lower than that of D. salina (64%) or of C. reinhardtii (59%). The phylogenetic analysis of the PSY from microalgaes, cyanobacteria, plants, and bacteria is illustrated in Fig. 4. Analysis was conducted in MEGA4 (Tamura et al. 2007) using the UPGMA method. The predicted CzPSY forms a cluster with the rest of the microalgae studied, which are phylogenetically closely related to PSY of plants (between 65% and 70% of identity and around 75–80% similarity). The degree of homology was lower with cyanobacterial phytoene synthases (55–60% identity and 65–70% similarity). As other algal PSY, CzPSY was distantly related to bacterial PSY (CRTB), sharing with them only a few conserved motifs and less than 40% identity.

Since in microalgae and plants PSY is located in the chloroplast, we analysed the CzPSY sequence with two different programs to determine the presence of a signal peptide. The Predotar v. 1.03 program predicted putative plastid localization for the CzPSY, and ChloroP 1.1 server identified an N-terminal chloroplast transit peptide at position 45. Analysis with ProtScale and TopPred servers identified a deduced transmembrane domain of CzPSY located between amino acids 236 and 256.

Functional analysis of the CzPSY in E. coli

The use of E. coli engineered to produce different carotenogenic substrates is an efficient and frequently used method for the functional characterization of the enzymes of the carotenoid biosynthetic pathway (Cunningham and Gantt 2005; Cunningham et al. 1996), due to the complexity of determining the activity of these membrane-associate and low-abundant enzymes.

In order to check the functionality of the recently isolated gene, the full-length ORF of CzPSY was amplified and cloned into pQE-80L vector under the control of the β-galactosidase promoter. The resulting plasmid (pQE-CzPSY) was introduced in E. coli that carried the plasmid pAC-85b, which contained the carotenogenic genes responsible for the synthesis of β-carotene as final product except PSY gene. HPLC analysis of carotenoids extracted from E. coli showed that cells containing pAC-85b and pQE-CzPSY produced β-carotene (Fig. 5). As negative controls, E. coli co-transformed with pAC-85b or pAC-85b and empty pQE-80L were used, resulting in no accumulation of carotenoids. These results confirmed the functionality of CzPSY gene product, which could catalyze the conversion of two molecules of GGPP into phytoene, as most PSY previously isolated.

Nuclear transformation of C. reinhardtii with PSY gene from C. zofingiensis and screening of transformants obtained

The complete coding region of CzPSY gene was amplified by PCR and cloned between the XhoI and EcoRI restriction sites of the Chlamydomonas expression vector pSI105-Tp1, resulting in the plasmid pSI105-psy. C. reinhardtii cells amplified by PCR. A plasmid containing the PSY gene was used as a positive control (lane 3). b HincII and HindIII restriction sites present in the CzPSY gene. The black bar indicates the probe location.
were transformed with plasmid pSI105-psy and the transformants were firstly selected on the basis of their paromomycin resistance. The colonies obtained after transformation that showed resistance to paromomycin were screened for the insertion of CzPSY cDNA in their genome by PCR test. Figure 6 shows some of the positive transformants analysed exhibiting a band of 147 bp, which correspond to the CzPSY cDNA integrated in their genome. The primers used for PCR analysis to confirm this integration were the same used for the expression analysis of CzPSY (Table 1). More than 100 colonies resistant to paromomycin were analysed and 80% of them were found positives. Some of these positive transformants selected by PCR were grown in liquid TAP medium and their carotenoids content analysed by HPLC and the relative mRNA levels of C. reinhardtii PSY (CrPSY) and CzPSY genes determined by qRT-PCR.

Analysis of carotenoids content and mRNA levels of CrPSY and CzPSY genes in both parental and CzPSY-transformed strains of C. reinhardtii

The mRNA relative abundance of endogenous (CrPSY) and exogenous (CzPSY) phytoene synthase genes of parental and CzPSY-transformant strains of C. reinhardtii was monitored by qRT-PCR, and changes in cellular carotenoids content were determined in order to correlate transcript levels with the biosynthesis of those specific carotenoids. Figure 7 shows carotenoids content and mRNA levels of CrPSY and CzPSY genes in parental and six selected CzPSY transformants. The carotenoid profile of parental and CzPSY transformed C. reinhardtii cells were very similar, being violaxanthin the major carotenoid, followed by lutein and β-carotene, α-carotene showing the lowest levels. However, some of the transformants (approximately 10% of the positives found) exhibited a violaxanthin and lutein content of 1.8- to 2.0- and 2.0- to 2.2-fold higher than the parental strain, whereas β-carotene and α-carotene levels were virtually identical to those of the wild type strain. T10 and T11 are two representative examples of these violaxanthin and lutein hyperproducing mutants (Fig. 7a). Other transformants showed only a slight increase

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**Fig. 4** UPGMA tree analysis of the indicated plant, algal, cyanobacterial, and bacterial PSY amino acid sequences. Analysis was performed in MEGA4 (Tamura et al. 2007). The GenBank accession numbers for other species are as follows: Dunaliella salina, AAT46069; Dunaliella sp., ABE97388.1; Dunaliella bardawil, AAB51287.1; Haematococcus pluvialis, AAW28851.1; Chlamydomonas reinhardtii, XP_001701192.1; Triticum turgidum, ACQ59152.1; Sorghum bicolor, AED78689; Salicornia europaea, AAX19898.1; Solanum lycopersicum, ABM45873.1; Tagetes erecta, AAG10427.1; Synechocystis sp. PCC 6803, BAA17848.1; Anabaena variabilis PCC 29413, YP_325286.1; Nostoc sp. PCC 7120, BAB73532.1; Gloeobacter violaceus PCC 7421, NP_924690.1; Synechococcus sp. JA-3-3-Ab, YP_473801.1; Brevibacterium linens, AAF65581.1; Corynebacterium glutamicum, AAK64298.1. The tree is drawn to scale, with branch lengths in the same units as those of the evolutionary distances used to infer the phylogenetic tree. The evolutionary distances correspond to the number of amino acid substitutions per site and were computed using the Poisson correction method. Numbers at nodes indicate bootstrap values calculated over 500 replicates.

**Fig. 5** HPLC elution profile of carotenoids extracted from cultures of E. coli carrying plasmids pAC-85b+pQE-CzPSY. The absorption spectrum of β-carotene is also shown. E. coli BL21 (DE3) colonies transformed with the indicated plasmids were isolated in the presence of chloramphenicol+ampicillin. Peak identification, β-carotene.
in the levels of violaxanthin and lutein, although in all of them the exogenous PSY gene was adequately transcribed and high levels of CzPSY mRNA were found.

Regarding the expression analysis, in all the transformants analysed the levels of CzPSY mRNA were higher than those of the endogenous PSY, since exogenous CzPSY was constitutively expressed under the control of the strong promoters rbcS2 and hsp70A. Transformants T1 and T10 showed the highest CzPSY levels, which reached 140-fold the level of the endogenous PSY. Expression levels of endogenous PSY in transformants were quite similar to that of the wild type cells. Levels of PSY transcripts were standardized respect to the housekeeping control gene (CBLP) and expressed as numbers of normalized molecules.

Discussion

Phytoene synthase catalyzes the first step of the carotenoid biosynthetic pathway and is considered as a rate-limiting key enzyme in this pathway (Cunningham and Gantt 1998) and as the branching enzyme that determines the carbon flux towards carotenoids production (Shewmaker et al. 1999). Therefore, PSY has been the target of genetic manipulation in many crop plants to increase carotenoid biosynthesis (Fray et al. 1995; Fraser et al. 2002; Lindgren et al. 2003). The CzPSY gene isolated in this work, as well as the known PSY genes of eukaryotic microalgae, have a smaller number of exons than the common five or seven exons in plants. Southern blot analysis has demonstrated that in C. zofingiensis, as in other microalgae as C. reinhardtii and D. salina (McCarthy et al. 2004; Lohr et al. 2005; Yan et al. 2005), only a single gene coding for PSY is present. However, recently, it has been described that other algae as Dunaliella bardawil and Micromonas pusilla contain either multiple paralogous or orthologous copies of the PSY, which could be expected that analogous to the diversity of PSY genes and their differential expression in higher plants, some algae also differentially could regulate expression of their multiple copies of PSY gene. The comparative analyses of various algal genomes for PSY in combination with phylogenetic analyses have suggested an ancient gene duplication creating two classes of PSY (Tran et al. 2009). Our results indicate that only one class of PSY seems to be present in C. zofingiensis. Alignments at the protein level have indicated that CzPSY has the essential characteristics of both classes of PSY including predicted substrate-Mg²⁺-binding sites (Aspartate-rich regions) and catalytic residues (data not shown). Moreover, the alignments have showed that the sequence differences between plant and bacterial PSY are mainly found at the N terminus due to the presence of a signal.
peptide responsible for the localization of these enzymes in chloroplasts and chromoplasts (Cunningham and Gantt 1998). The high homology degree found between the predicted protein encoded by the new gene isolated from C. zofingiensis and plant phytoene synthases, specially with the known PSY of other green microalgae, would have been probably enough to consider this new gene as a PSY, but functional analysis has definitely confirmed this hypothesis. It has been shown by functional complementation in E. coli that this gene encodes phytoene synthase. Although the environment and the processing of the putative PSY gene in the prokaryotic system probably differ from those in the algal original system, the obtained gene product was functional and able to catalyze the synthesis of phytoene, complementing plasmid pAC85 and resulting in the accumulation of β-carotene (Fig. 5). This confirms that the isolated gene is responsible for the synthesis of phytoene from two GGPP molecules.

Overexpression of bacterial or plant PSY genes in crop plants has resulted in increases in total carotenoid content of about 1.8–6.3-fold in carrot roots (Hauptmann et al. 1997), tomato fruit (Fraser et al. 2002), and potato tubers (Ducreux et al. 2005). Higher carotenoid increases have been reported in plant or tissues with no carotenoids or with very original levels (Farré et al. 2010), such as a 50-fold increase in canola seeds (Shewmaker et al. 1999). Microalgae combine the fast and easy growth of bacteria with an active isoprenoid metabolism that ensures enough precursors for carotenogenic pathway and adequate storage capacity. The unicellular microalgae C. reinhardtii is the first and best studied transformed chlorophyta. It grows at high rates under photoautotrophic, heterotrophic or mixotrophic conditions and its nuclear genetic manipulation is easy and well established. All these reasons make C. reinhardtii a good candidate to express foreign carotenogenic genes, for both carrying out basic metabolic and regulatory studies of the pathway as well as for the biotechnological production of commercially interesting carotenoids. As the sequence of the PSY from C. zofingiensis shares a 76% of identity with that reported for Chlamydomonas PSY, the expression of such a gene in the nuclear genome of this microalgae is a step forward both in the characterization of the gene and in the increased production of interesting carotenoids in microalgae. In this study, the nuclear transformation of C. reinhardtii with CzPSY gene resulted in a significant accumulation of violaxanthin and lutein content that reached 2.0- and 2.2-fold, respectively, as compared with the parental level (Fig. 7a). Moreover, transformants showed an overexpression of C. zofingiensis PSY gene due to the strong promoters used (Fig. 7b). These results suggest that the higher carbon flux from GGPP to carotenoid synthesis was promoted by a combination of the expression of CzPSY gene product in C. reinhardtii as well as the expression of the endogenous copy of the gene. The CzPSY transcript level reached 140-fold the level of the endogenous PSY of the wild type in the transformants T1 and T10, however the violaxanthin and lutein contents in these transformants were only 2.0- and 2.2-fold higher than the parental strain, respectively, in T10 transformant, and 7% and 21% higher, respectively, in T1. Although transcriptional control has been shown to be the most important regulation factor for carotenogenic genes, a possible explanation for these results is that post-transcriptional and translational controls play also important roles, as well as the stability of RNA, since silencing of the PSY endogenous gene by artificial miRNAs has been described in Chlamydomonas (Molnar et al. 2009). Moreover, an increase in the levels of PSY cannot be directly correlated with an increase in carotenoids content, since the protein could not be correctly targeted, processed or assembled into a fully functional complex. In transgenic tomato and potato plants expressing a bacterial phytoene synthase (CRTB), despite CRTB enzyme activity being substantially elevated, there was only a moderate increase in total carotenoid content and no linear correlation between the levels of transcript, protein, enzyme activity, and total carotenoids (Fraser et al. 2002; Ducreux et al. 2005).

Overexpression of an exogenous PSY in tomato resulted in an accumulation of phytoene, and in a decrease in the flux control coefficient for PSY. Then, the increase in PSY activity shifts probably the regulatory step from PSY to a later enzyme (Shewmaker et al. 1999; Fraser et al. 2002). Accumulation of β-carotene in higher plants overexpressing exogenous PSY, probably indicates that β-carotene hydroxylases become rate-limiting as the carotenogenic flux increases (Ducreux et al. 2005). In our study no increase in the content of phytoene or other carotenoids intermediates were detected, showing that other enzymes in the pathway are not limited in Chlamydomonas, and that phytoene, β-carotene or other intermediates are not accessible to the downstream enzymes of the pathway in the single cell of Chlamydomonas. Phytoene was consecutively converted to the downstream metabolites α-carotene, lutein, β-carotene and violaxanthin, catalyzed by endogenous carotenoid biosynthetic enzymes such as PDS, ZDS, LCYb, LCYe, CHYb, and ZEP.

In plants and microalgae, carotenoid biosynthesis is part of the plastidic terpenoid metabolism. GGPP is a common intermediate to the different terpenoid biosynthetic pathway such as carotenoids, chlorophylls, gibberellins and quinones. The engineering of any terpenoid pathway may have a direct effect on other branches of the pathway. In the case of carotenoid overproduction, the increase of carbon flux into the carotenoid pathway can produce limitations especially in the synthesis of gibberellins, chlorophylls,
and quinones with negative effects on growth and photosynthesis, respectively. In some higher plants overexpressing PSY genes undesired collateral effects or low carotenoids increase were observed (Farré et al. 2010). The constitutive expression of phytene synthase genes in tomato (Lycopersicum esculentum) and tobacco (Nicotiana tabacum) resulted in dwarfism and chlorosis in the plants (Fray et al. 1995; Busch et al. 2002) correlated with a decrease of giberellins. But so far, in our case, no atypical phenotype was observed in the Chlamydomonas transformants, showing even a similar growth rate than the wild type cultures (data not shown).

In conclusion, C. zofingiensis is a model green microalga to study the regulation of the carotenoid biosynthetic pathway, since it accumulates both lutein and astaxanthin. In this study the phytene synthase gene from C. zofingiensis has been isolated and functionally characterized. The overexpression of CzPSY in Chlamydomonas cells under the control of strong constitutive promoters has resulted in a significant enhancement in the content in violaxanthin and lutein, reaching 2.0- and 2.2-fold the values of control untransformed cells, respectively. This is a successful case of manipulation of the carotenogenic pathway in eukaryotic microalgae, which opens up the possibility of enhancing the productivity of microalgal-based systems to produce carotenoids and offers an excellent tool to gain basic knowledge about an important pathway that at present is not yet completely characterized and a promising alga for industrial applications.

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