

Complete Biosynthetic Pathway of the C₅₀ Carotenoid Bacterioruberin from Lycopene in the Extremely Halophilic Archaeon *Haloarcula japonica*

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ABSTRACT

Haloarcula japonica, an extremely halophilic archaeon that requires high concentrations of NaCl for growth, accumulates the C₅₀ carotenoid bacterioruberin (BR). By homology analysis, a gene cluster, including *c0507*, *c0506*, and *c0505*, was found and predicted to be involved in the synthesis of bacterioruberin. To elucidate the function of the encoded enzymes, we constructed *Ha. japonica* mutants of these genes and analyzed carotenoids produced by the mutants. Our research showed that *c0507*, *c0506*, and *c0505* encoded a carotenoid 3,4-desaturase (CrtD), a bifunctional lycopene elongase and 1,2-hydratase (LyeJ), and a C₅₀ carotenoid 2',3'-hydratase (CruF), respectively. The above three carotenoid biosynthetic enzymes catalyze the reactions that convert lycopene to bacterioruberin in *Ha. japonica*. This is the first identification of functional CrtD and CruF in archaea and elucidation of the complete biosynthetic pathway of bacterioruberin from lycopene.

IMPORTANCE

Haloarcula japonica, an extremely halophilic archaeon, accumulates the C₅₀ carotenoid bacterioruberin (BR). In this study, we have identified three BR biosynthetic enzymes and have elucidated their functions. Among them, two enzymes were found in an archaeon for the first time. Our results revealed the biosynthetic pathway responsible for production of BR in *Ha. japonica* and provide a basis for investigating carotenoid biosynthetic pathways in other extremely halophilic archaea. Elucidation of the carotenoid biosynthetic pathway in *Ha. japonica* may also prove useful for producing the C₅₀ carotenoid BR efficiently by employing genetically modified haloarchaeal strains.

Carotenoids are natural pigments synthesized by bacteria, archaea, algae, fungi, and plants (1). They are involved in photosynthesis as accessory pigments (2), functioning as antioxidants (3, 4), light protection pigments (5, 6), and membrane stabilizers (7). Thus far, more than 750 carotenoids have been identified and classified as C₃₀, C₄₀, and C₅₀ carotenoids, depending on the number of carbons in their carotene backbones (1). Most of these compounds are based on the symmetric C₄₀ backbone, phytoene, which is formed by condensation of two molecules of geranylgeranyl pyrophosphate (GGPP; C₂₀PP) (8). Only a small number of C₃₀ carotenoids, which arise from the fusion of two molecules of farnesyl pyrophosphate (FPP; C₁₅PP), and even fewer C₅₀ carotenoids, which are derived from the C₄₀ structure by the addition of two 5-carbon (C₅) isoprene units, have been discovered to date (1).

In nature, several restricted groups of bacteria, including species belonging to the bacterial genera *Corynebacterium*, *Dietzia*, and *Micrococcus*, and extremely halophilic archaea, such as *Haloarcula salinarum*, show accumulation of C₅₀ carotenoids (9, 10). To date, only three biosynthetic pathways of cyclic C₅₀ carotenoids, the ε-cyclic C₅₀ carotenoid decaprenoxanthin in *Corynebacterium glutamicum* (11–13), the β-cyclic C₅₀ carotenoid 2,2'-bis-(4-hydroxy-3-methylbut-2-enyl)-β,β-carotene in *Dietzia* sp. strain CQ4 (14), and the γ-cyclic C₅₀ carotenoid sarcinaxanthin in *Micrococcus luteus* NCTC2665 (15), have been described, based on their structures. The acyclic C₅₀ carotenoid bacterioruberin (BR) is known to be produced in some extremely halophilic archaea. BR of *Haloferax volcanii* has been isolated and identified by chromatographic (thin-layer chromatography [TLC] and high-performance liquid chromatography [HPLC]), spectroscopic (mass

spectrometry [MS], nuclear magnetic resonance [NMR] analysis, and circular dichroism [CD] spectroscopy), and chemical (silylation and methylation) methods (16). *Haloferax mediterranei* can also accumulate BR, and a two-stage culture method has been applied to increase production of C₅₀ carotenoids in the laboratory (17). In addition to studies on the antioxidant properties of BR (18), the regulating mechanism of BR biosynthesis in *Hb. salinarum* has been investigated (19). *Hb. salinarum* is also known to produce the bacteriorhodopsin light-induced proton pump, comprising retinal and bacterioopsin (20). The carotenoids and retinal biosynthetic pathways of *Hb. salinarum* have been briefly described previously (10). That report proposed that retinal and BR are synthesized from a common precursor, lycopene, which is generated from phytoene, as occurs in C₄₀ carotenoid biosynthetic pathways. Although the enzyme Lye, which catalyzes the commit-

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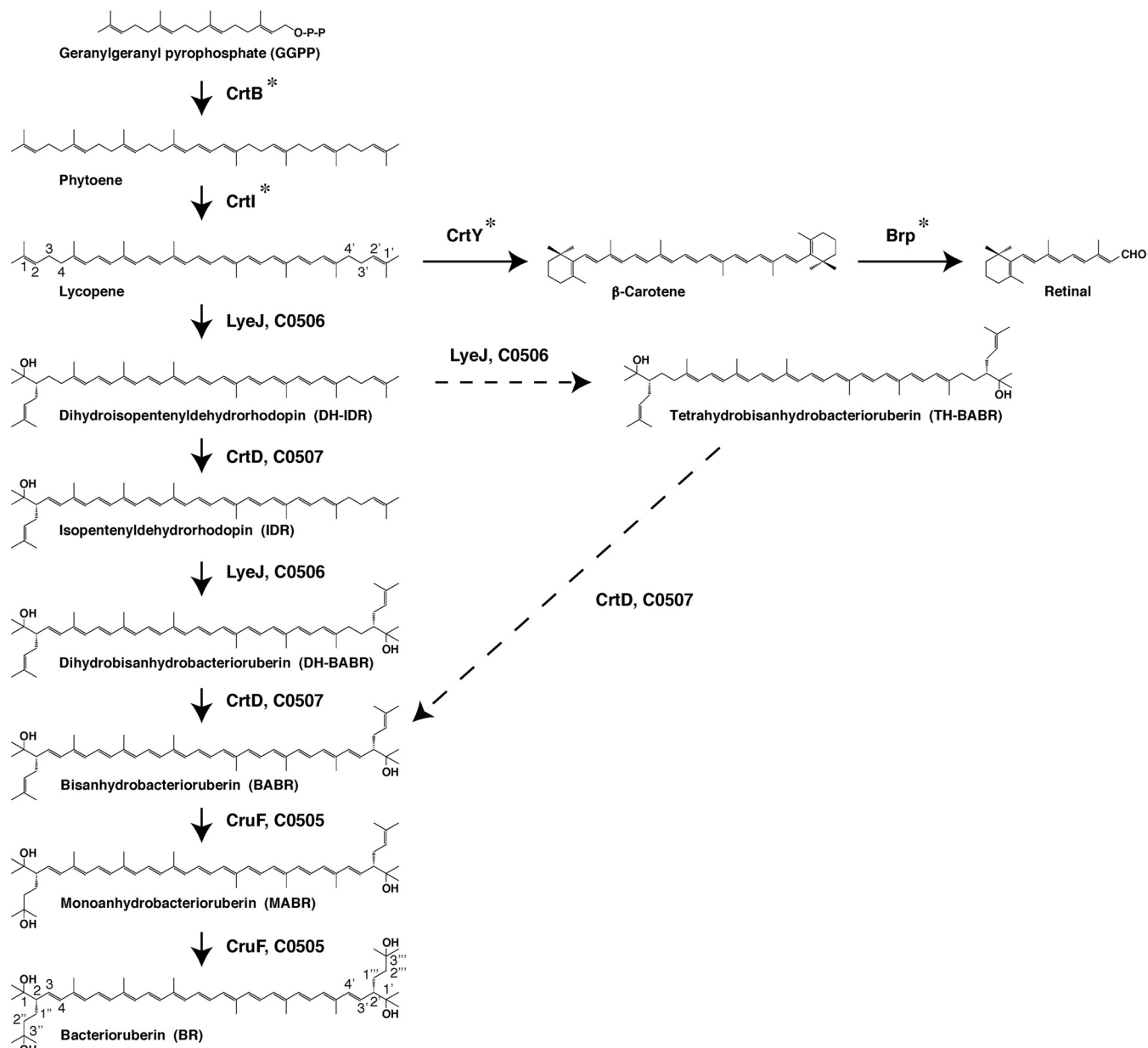


FIG 1 Proposed main steps in the carotenoid and retinal biosynthetic pathways of *Haloarcula japonica*. Solid arrows signify the main carotenoid biosynthetic steps. The dashed arrows indicate other possible carotenoid biosynthetic steps. Asterisks indicate the unidentified carotenoid biosynthetic enzymes.

ted step in BR biosynthesis, has been described in *Hb. salinarum*, the details of the BR biosynthetic pathway remain unclear.

Haloarcula japonica is a predominantly triangular, disc-shaped, extremely halophilic archaeon that requires high concentrations of NaCl for growth (21). In our previous study, we found that *Ha. japonica* can produce isopentenyldehydrorhodopin (IDR; C₄₅), bisanhydrobacterioruberin (BABR; C₅₀), and monoanhydrobacterioruberin (MABR; C₅₀) as intermediates and that BR is produced as the final product (22). In addition, it has been suggested that *Ha. japonica* also produces the retinal proteins cruxrhodopsin (23) and cruxhalorhodopsin (24). Thus, retinal could also be synthesized from carotenoids in *Ha. japonica*. Based on the results of our previous study and the proposed biosynthetic pathways of

carotenoids in *Hb. salinarum* (10), the carotenoid pathways and retinal biosynthetic pathways in *Ha. japonica* are suggested to be as shown in Fig. 1. Phytoene synthase (CrtB) condenses two GGPP molecules to yield the colorless carotenoid phytoene. Lycopene is generated from phytoene via a series of desaturation reactions that are catalyzed by phytoene desaturase (CrtI). Subsequently, the steps are divided into the retinal biosynthetic pathway and the BR biosynthetic pathway. In retinal synthesis, the cyclization of lycopene to β-carotene is catalyzed by lycopene β-cyclase (CrtY), and β-carotene is then cleaved to form a retinal (C₂₀) by β-carotene dioxygenase (Brp) cleavage. In the BR biosynthetic pathway, lycopene is used as a precursor and is converted to BR by introduction of two C₅ isoprene units, two double bonds, and four hydroxyl

TABLE 1 Bacterial strains and plasmids used in this study

Strain or plasmid	Relevant property or properties ^a	Source or reference
Strains		
<i>E. coli</i> JM109	Host for cloning vectors	Laboratory stock
<i>E. coli</i> JM110	Host used for preparing plasmid that is free of Dam or Dcm methylation	Laboratory stock
<i>Ha. japonica</i>	Wild type (JCM 7785 ^T)	21
<i>Ha. japonica</i> Δ <i>c0507</i>	<i>c0507</i> gene mutant of <i>Ha. japonica</i>	This work
<i>Ha. japonica</i> Δ <i>c0506</i>	<i>c0506</i> gene mutant of <i>Ha. japonica</i>	This work
<i>Ha. japonica</i> Δ <i>c0505</i>	<i>c0505</i> gene mutant of <i>Ha. japonica</i>	This work
<i>Ha. japonica</i> Δ <i>c0507</i> (pJ <i>c0507</i>)	Δ <i>c0507</i> complement with pJ <i>c0507</i>	This work
<i>Ha. japonica</i> Δ <i>c0506</i> (pJ <i>c0506</i>)	Δ <i>c0506</i> complement with pJ <i>c0506</i>	This work
<i>Ha. japonica</i> Δ <i>c0505</i> (pJ <i>c0505</i>)	Δ <i>c0505</i> complement with pJ <i>c0505</i>	This work
Plasmids		
pUC119	<i>E. coli</i> cloning vector; Ap ^r	Laboratory stock
pWL102	<i>E. coli</i> -haloarchaea shuttle vector; Ap ^r Mev ^r	29
pDrHj2 ^b	<i>E. coli</i> -haloarchaea shuttle vector; Ap ^r Mev ^r	Laboratory stock
pWL102-Δ <i>c0507</i>	pWL102 derivative containing disrupted fragment of <i>c0507</i> gene	This work
pDrHj2-Δ <i>c0506</i>	pDrHj2 derivative containing disrupted fragment of <i>c0506</i> gene	This work
pDrHj2-Δ <i>c0505</i>	pDrHj2 derivative containing disrupted fragment of <i>c0505</i> gene	This work
pJFZ33	pWL102 derivative carrying <i>Ha. japonica</i> <i>csg</i> promoter and <i>ftsZ2</i> gene; Ap ^r Mev ^r	27
pJ <i>c0507</i>	pJFZ33 derivative in which <i>ftsZ2</i> is replaced by the <i>c0507</i> gene	This work
pJ <i>c0506</i>	pJFZ33 derivative in which <i>ftsZ2</i> is replaced by the <i>c0506</i> gene	This work
pJ <i>c0505</i>	pJFZ33 derivative in which <i>ftsZ2</i> is replaced by the <i>c0505</i> gene	This work

^a Ap^r, ampicillin resistance; Mev^r, pravastatin resistance.

^b The pDrHj2 plasmid was constructed as follows. The sequence between *Sac*I and *Kpn*I in pUC119 was replaced with the mevinoline resistance gene, and the sequence between *Hind*III and *Xba*I was replaced with the gene of a halophilic β-galactosidase (*bgaH*) under the control of *csg* promoter. The *bgaH* gene was obtained from *Haloferax alicantei* and was oriented in the direction opposite that in which the mevinoline resistance gene was oriented. The *csg* promoter is from *Haloarcula japonica*. *Ha. japonica* has a large amount of a glycoprotein (cell surface glycoprotein [CSG]) on the cell surface, suggesting that the *csg* promoter is powerful.

groups into the lycopene. All of the related enzymes and the complete pathway for biosynthesis of BR from lycopene remain unknown.

In this study, we identified and characterized three main BR biosynthetic enzymes in *Ha. japonica*, namely, C0506, C0507, and C0505. They function as a bifunctional lycopene elongase and 1,2-hydratase (LyeJ), a carotenoid 3,4-desaturase (CrtD), and a C₅₀ carotenoid 2',3''-hydratase (CruF), respectively, to generate BR from lycopene. These findings contribute to elucidation of the complete carotenoid biosynthetic pathway in extremely halophilic archaea.

MATERIALS AND METHODS

Microbial strains and growth conditions. All strains and plasmids used in this study are listed in Table 1. The wild-type and mutant strains of *Haloarcula japonica* (JCM 7785^T) were grown at 37°C in the dark with a complex medium, as described previously (25). The medium was supplemented with 8 μg/ml pravastatin (a gift from Daiichi Sankyo, Tokyo, Japan) instead of mevinoline, when required. *Escherichia coli* was cultured in LB medium at 37°C (26). Ampicillin (50 μg/ml) was added when required.

Isolation of genomic DNA and total RNA from *Ha. japonica*. *Ha. japonica* genomic DNA was isolated as described previously (27). Total RNA of *Ha. japonica* was prepared using Sepasol RNA I (Nacalai Tesque, Kyoto, Japan) according to the manufacturer's instructions. The resulting total RNA was further treated with DNase I (GE Healthcare, Bucks, United Kingdom) to remove trace amounts of contaminating genomic DNA.

RT-PCR and PCR. Oligonucleotide primers for reverse transcription-PCR (RT-PCR) and PCR were purchased from Operon Biotechnologies (Tokyo, Japan). The oligonucleotide primers used in this study are listed in Table 2. RT-PCR was performed under the following conditions. Total RNA (1.0 μg) was reverse transcribed at 45°C for 60 min in 20 μl of the reaction buffer, which contained 20 pmol of each primer (*c0507*-A,

c0506-A, or *c0505*-A), 0.3 mM (each) deoxynucleotide triphosphate, 2.5 mM manganese (II) acetate, 20 U RNase inhibitor (Toyobo, Osaka, Japan), and 5 U *rTth* DNA polymerase (Toyobo). The cDNA generated was then amplified by PCR. PCR was carried out using KOD-Plus-Dash or KOD-Dash (Toyobo) DNA polymerase, according to the manufacturer's instructions. Primer set *c0507*-S/*c0507*-A, primer set *c0506*-S/*c0506*-A, or primer set *c0505*-S/*c0505*-A was used to confirm the transcription of *c0507*, *c0506*, or *c0505* in the wild-type and mutant strains of *Ha. japonica*.

Construction of *c0507*, *c0506*, and *c0505* mutant strains. Mutants with deletions and frameshift mutations were constructed by homologous recombination (28). The plasmid pWL102-Δ*c0507*, containing a disrupted *c0507* gene, was constructed as follows. The targeted gene, *c0507*, was amplified from *Ha. japonica* using the primer set *c0507*-S1/*c0507*-A1. The amplified PCR fragment was ligated into the *Sma*I site of pUC119, yielding plasmid pUC119-*c0507*. A 652-bp DNA fragment, ranging from base 378 to base 1029 of *c0507*, was excised from pUC119-*c0507* by *Csp*45I and *Eco*RV digestion, followed by blunting and self-ligation, to yield plasmid pUC119-Δ*c0507*. Plasmid pUC119-Δ*c0507* was digested with *Eco*RI and *Hind*III, releasing a disrupted *c0507* gene. Plasmid pWL102 (29) was also digested with *Eco*RI and *Hind*III to eliminate the *Hf. volcanii* ori and was then ligated to the disrupted *c0507* gene, yielding plasmid pWL102-Δ*c0507*.

Plasmids pDrHj2-Δ*c0506*, containing a disrupted *c0506* gene, and pDrHj2-Δ*c0505*, containing a disrupted *c0505* gene, were constructed as follows. The targeted genes, *c0506* and *c0505*, were amplified from *Ha. japonica* using primer set *c0506*-S1/*c0506*-A1 and primer set *c0505*-S1/*c0505*-A1, respectively. Amplified PCR fragments were ligated into the *Sma*I site of pUC119, yielding plasmid pUC119-*c0506* and pUC119-*c0505*, respectively. A 292-bp DNA fragment, ranging from base 271 to base 562 of *c0506*, and a 221-bp DNA fragment, ranging from base 318 to base 538 of *c0505*, were removed from the pUC119-*c0506* and pUC119-*c0505* plasmids by *Xba*I PCR, using primer set *c0506*-S3/*c0506*-A3 and primer set *c0505*-S3/*c0505*-A3, respectively. This was followed by self-ligation, yielding plasmids pUC119-Δ*c0506* and pUC119-Δ*c0505*, respec-

TABLE 2 Sequences of the primers used in this study

Primer	Sequence ^a	Usage	
c0507-S	5'-GGTTGGCCTCCAGCTCATTG-3'	To confirm the transcription of target genes by RT-PCR	
c0507-A	5'-CTCGTGGTCCGGCAGGAGTTC-3'		
c0506-S	5'-TCGCCCTGTTTCTGTACTTCAC-3'		
c0506-A	5'-GATATCTGGAATCGCCGAGAAGG-3'		
c0505-S	5'-CACCGAGAACCGGTTTACCATC-3'		
c0505-A	5'-ACTGCGGCATCTCGTAGATCC-3'		
c0507-S1	5'-ATGTGGCCACGAATGCCATAC-3'		To amplify the target genes by PCR for the construction of mutants
c0507-A1	5'-GGTCTGTCGGGAGTTCTGGC-3'		
c0506-S1	5'-GGACATCGCCTGAGTAATGCC-3'		
c0506-A1	5'-GGTCTTTGCCGCTACCCATAC-3'		
c0506-S3	5'-TCTGGGCGATGGGGATGCAC-3'		
c0506-A3	5'-TTTCGGATGTGCTCGTCGATATCGG-3'		
c0505-S1	5'-GCTGTATGGGTAGCGGCAAAG-3'		
c0505-A1	5'-GAGCTGGCTGATCCAAACGG-3'		
c0505-S3	5'-TGGATCCTGAGCCGTAGCTATC-3'		
c0505-A3	5'-AGCTCGATGCCGTAGGAGTACAG-3'		
c0507-S4	5'-GAATTCATATGAGTGACTTGTCCGGTG-3'	To amplify the intact target genes by PCR for the complementation study	
c0507-A4	5'-GGGGATCCTCATTAGTGGTGGTGGTGGTGGGCGATGCTCTCGATGAGC-3'		
c0506-S4	5'-GAATTCATATGCCAACTCCCAGACAG-3'		
c0506-A4	5'-GGGGATCCTCATTAGTGGTGGTGGTGGTGGTGGTGGCCATACAGCATCACCCAC-3'		
c0505-S4	5'-GAATTCATATGGGTAGCGGCAAAGACC-3'		
c0505-A4	5'-GGCCATGGTCATTAGTGGTGGTGGTGGTGGTGGCCAGACGGCCCGACCG-3'		

^a The introduced NdeI, BamHI, and NcoI restriction sites are underlined.

tively. Plasmid pUC119- Δ c0506 was digested with EcoRI. After blunting, it was further digested with BamHI, and the fragment containing the disrupted c0506 gene was ligated to pDrHj2 (containing only the ColE1 ori), which had been digested with SmaI and BamHI, to yield plasmid pDrHj2- Δ c0506. A fragment containing the disrupted c0505 gene was amplified from pUC119- Δ c0505 using the c0505-S1/c0505-A1 primer set, and the fragment was ligated to pDrHj2, which had been digested with SmaI, to yield plasmid pDrHj2- Δ c0505.

Plasmids pWL102- Δ c0507, pDrHj2- Δ c0506, and pDrHj2- Δ c0505 were subjected to passage through *E. coli* JM110 to avoid the restriction barrier formation of extremely halophilic archaea (30). Transformation of *Ha. japonica* was performed using the polyethylene glycol method (27). Transformants were plated onto agar plates containing pravastatin. Pravastatin-resistant colonies were cultured in liquid medium without pravastatin for 96 h and plated onto agar plates without pravastatin in order to isolate recombinants in which the targeted gene on the genome was replaced with the corresponding disrupted gene. Gene disruption was confirmed by PCR analysis.

Complementation of mutant strains. Three plasmids, pJc0507, pJc0506, and pJc0505, were constructed as follows. The targeted genes, c0507, c0506, and c0505, were amplified from *Ha. japonica* using primer set c0507-S4/c0507-A4, primer set c0506-S4/c0506-A4, and primer set c0505-S4/c0505-A4, respectively. Amplified PCR fragments were ligated into the SmaI site of pUC119, yielding plasmids pUc0507, pUc0506, and pUc0505, respectively. Plasmids pUc0507 and pUc0506 were digested with NdeI and BamHI. The sequences containing targeted genes were ligated to pJFZ33 (a recombinant plasmid in which the *Ha. japonica* csg promoter sequence, the NdeI restriction site, the *Ha. japonica* ftsZ2 gene sequence, and the BamHI and NcoI restriction sites were inserted into the *E. coli*-haloarchaea pWL102 shuttle vector), which had also been digested with NdeI and BamHI, to yield plasmids pJc0507 and pJc0506, respectively. Plasmid pUc0505 was digested with NdeI and NcoI. The sequence containing c0505 was ligated to pJFZ33, which had also been digested with NdeI and NcoI, to yield a pJc0505 plasmid.

Plasmids pJc0507, pJc0506, and pJc0505 were subjected to passage

through *E. coli* JM110 and then transformed into the corresponding mutant strains as described above.

Total carotenoid extraction from *Ha. japonica*. The wild-type and mutant strains of *Ha. japonica* were precultured at 37°C. Four milliliters of preinoculum was transferred to a 2-liter Erlenmeyer flask containing 400 ml of liquid medium and cultured for 240 h at 37°C in the dark. Cells were harvested by centrifugation (4°C, 4,400 × g, 20 min). Carotenoids were extracted from *Ha. japonica* essentially as described previously (22) with the following modifications. Cells were suspended in acetone/methanol (7:2 [vol/vol]) which contained 0.1% (wt/vol) 2,6-di-*t*-butyl-*p*-cresol antioxidant to avoid the destruction of carotenoids. After cell disruption by sonication, samples were centrifuged (4°C, 840 × g, 2 min) to obtain the supernatant. The pellets were reextracted until all visible pigments were retained in the liquid phase. Finally, the supernatants were collected and evaporated to dryness in a vacuum. The dried carotenoid extract was used for monitoring the HPLC analysis of total carotenoids and mass spectrometric analysis of the purified carotenoids.

HPLC analysis of total carotenoids. The dried carotenoid extract was dissolved in a small volume of chloroform-methanol (3:1) and analyzed using an HPLC system (SCL-10A; Shimadzu, Kyoto, Japan) equipped with a μ Bondapak C₁₈ column (Waters, Milford, MA) (3.9 by 300 mm). The eluent was methanol-water (9:1) for the first 10 min, followed by 100% methanol, at a flow rate of 1.5 ml/min (31). Absorption spectra of the carotenoids were recorded with a photodiode-array detector (SPD-M20A; Shimadzu) attached to the HPLC apparatus.

Mass spectrometric analysis of purified carotenoid. The dried carotenoid extract was dissolved in 2 ml of acetone-hexane (1:1) and was then applied onto a column of DEAE-Toyopearl 650M (Tosoh, Tokyo, Japan) to separate the polar lipids from the carotenoids (32). The column was washed with acetone-hexane (1:1), and the colored fractions were collected. After evaporation to dryness, the collected fractions were dissolved in 1 ml chloroform-methanol (3:1) and spotted onto high-performance thin-layer chromatography (HPTLC) plates (HPTLC silica gel 60 with concentrating zone; Merck Millipore, Darmstadt, Germany). The HPTLC plates were developed with petroleum ether-acetone (7:1) in the dark, and

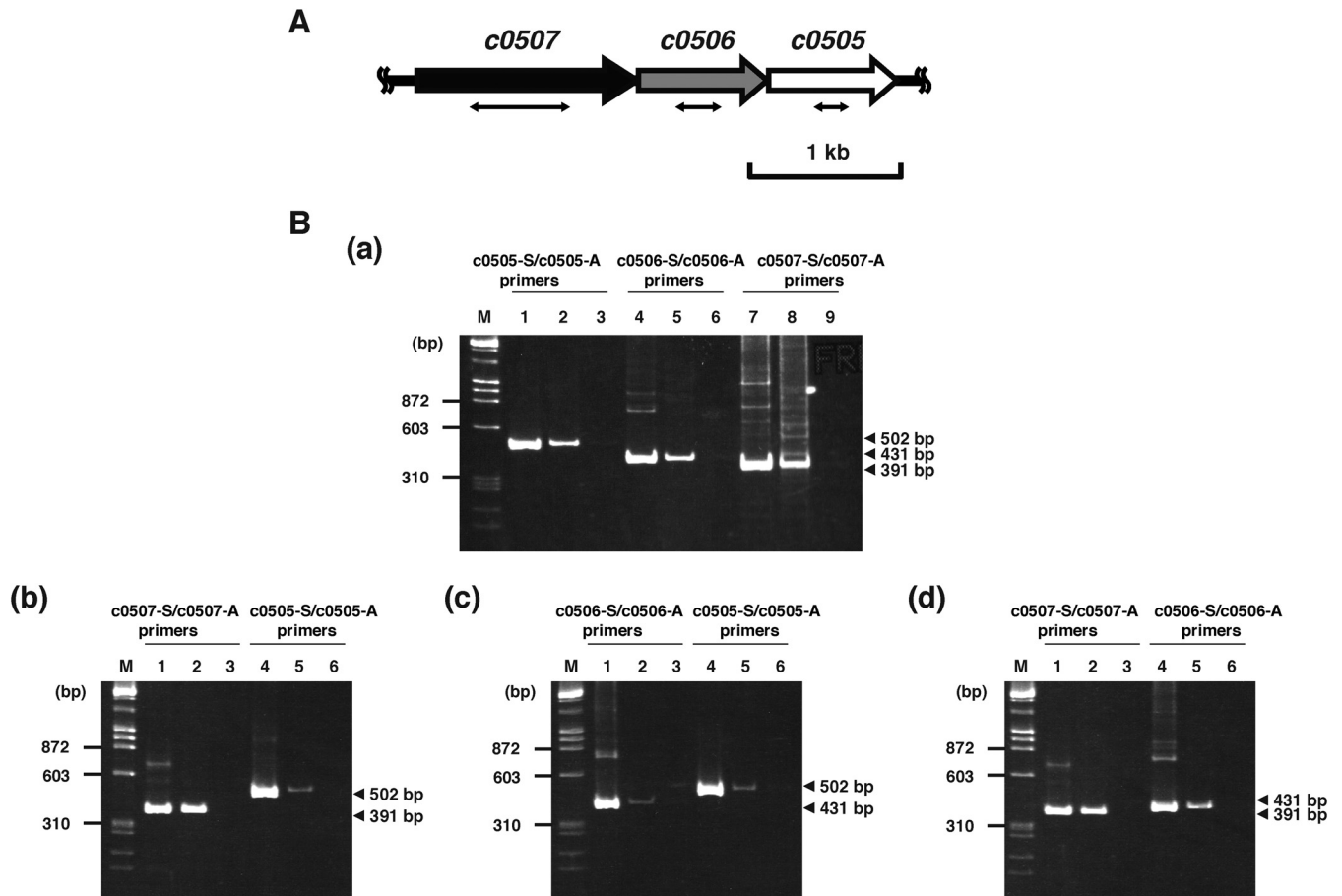


FIG 2 The *c0507-c0506-c0505* gene cluster and agarose gel electrophoresis of the RT-PCR products. (A) The *c0507-c0506-c0505* gene cluster. The double-headed arrows show the deleted parts in the corresponding mutants: a 652-bp fragment, ranging from base 378 to base 1029 of *c0507*, was removed in the $\Delta c0507$ strain; a 292-bp fragment, ranging from base 271 to base 562 of *c0506*, was removed in the $\Delta c0506$ strain; and a 221-bp fragment, ranging from base 318 to base 538 of *c0505*, was removed in the $\Delta c0505$ strain. (B) Agarose gel electrophoresis of the RT-PCR product. (Panel a) The genomic DNA and total RNA were isolated from wild-type *Ha. japonica*. (Panel b) The genomic DNA and total RNA were isolated from the $\Delta c0506$ strain. (Panel c) The genomic DNA and total RNA were isolated from the $\Delta c0507$ strain. (Panel d) The genomic DNA and total RNA were isolated from the $\Delta c0505$ strain. Lanes 1, 4, and 7, the positive-control reaction product using genomic DNA as the template; lanes 2, 5, and 8, the RT-PCR product; lanes 3, 6, and 9, the negative-control reaction product with total RNA not subjected to reverse transcription.

each separated fraction of carotenoids was collected. The collected silica powder was suspended with 1 ml chloroform-methanol (3:1) and then filtered through a polytetrafluoroethylene (PTFE) 0.2- μ m-pore-size filter (Lab Lab, Tokyo, Japan). Carotenoids in each fraction were further separated using the HPLC system described above, with the following modification. Elution was performed with 100% methanol at a flow rate of 1.5 ml/min. Carotenoids in peaks were collected, and their relative molecular masses were measured using an MStation JMS-700 mass spectrometry system (Jeol, Tokyo, Japan) in the fast-atom-bombardment (FAB) mode, with *m*-nitrobenzyl alcohol as a matrix.

Nucleotide sequence accession numbers. The DNA sequence data for *c0507*, *c0506*, and *c0505* were deposited in the DNA Data Bank of Japan (DDBJ), the European Molecular Biology Laboratory (EMBL), and the GenBank nucleotide sequence database. Their GenBank accession numbers are [LC008542](#), [LC008543](#), and [LC008544](#), respectively.

RESULTS

Identification of candidate genes. Recently, the draft genome sequence of *Ha. japonica* was determined (33). By homology analysis, some open reading frames (ORFs) (*c0507*, *c0506*, and *c0505*) were predicted to be candidate genes encoding carotenoid biosyn-

thetic enzymes. C0506 had 31% amino acid sequence identity to lycopene elongase (CrtEb) from *Corynebacterium glutamicum* and had 60% amino acid sequence identity to a lycopene elongase homolog (Lye) from *Hb. salinarum* (10). C0507 had 29% amino acid sequence identity to CrtI from *Pantoea ananatis* (34). Furthermore, C0507 also had 26% and 24% amino acid sequence identity to CrtD from *Rhodobacter capsulatus* (35) and *Deinococcus radiodurans* R1 (36), respectively. C0505, the homolog of CruF, had 30% and 31% amino acid identity to those of *Synechococcus* sp. strain PCC 7002 (37) and *Deinococcus radiodurans* R1 (38), respectively. In addition, *c0507*, *c0506*, and *c0505*, in that order, cluster on the genome of *Ha. japonica* (Fig. 2A), and RT-PCR analysis revealed that those genes were cotranscribed. Since carotenoid synthesis-related genes in some bacteria are assembled in clusters or in neighborhoods (34, 39), these genes were predicted to be involved in the carotenoid synthesis of *Ha. japonica*.

Analysis of *c0506* and *c0507*. To characterize *c0506*, a mutant of this gene was constructed by homologous recombination and designated strain $\Delta c0506$. The $\Delta c0506$ strain showed the same

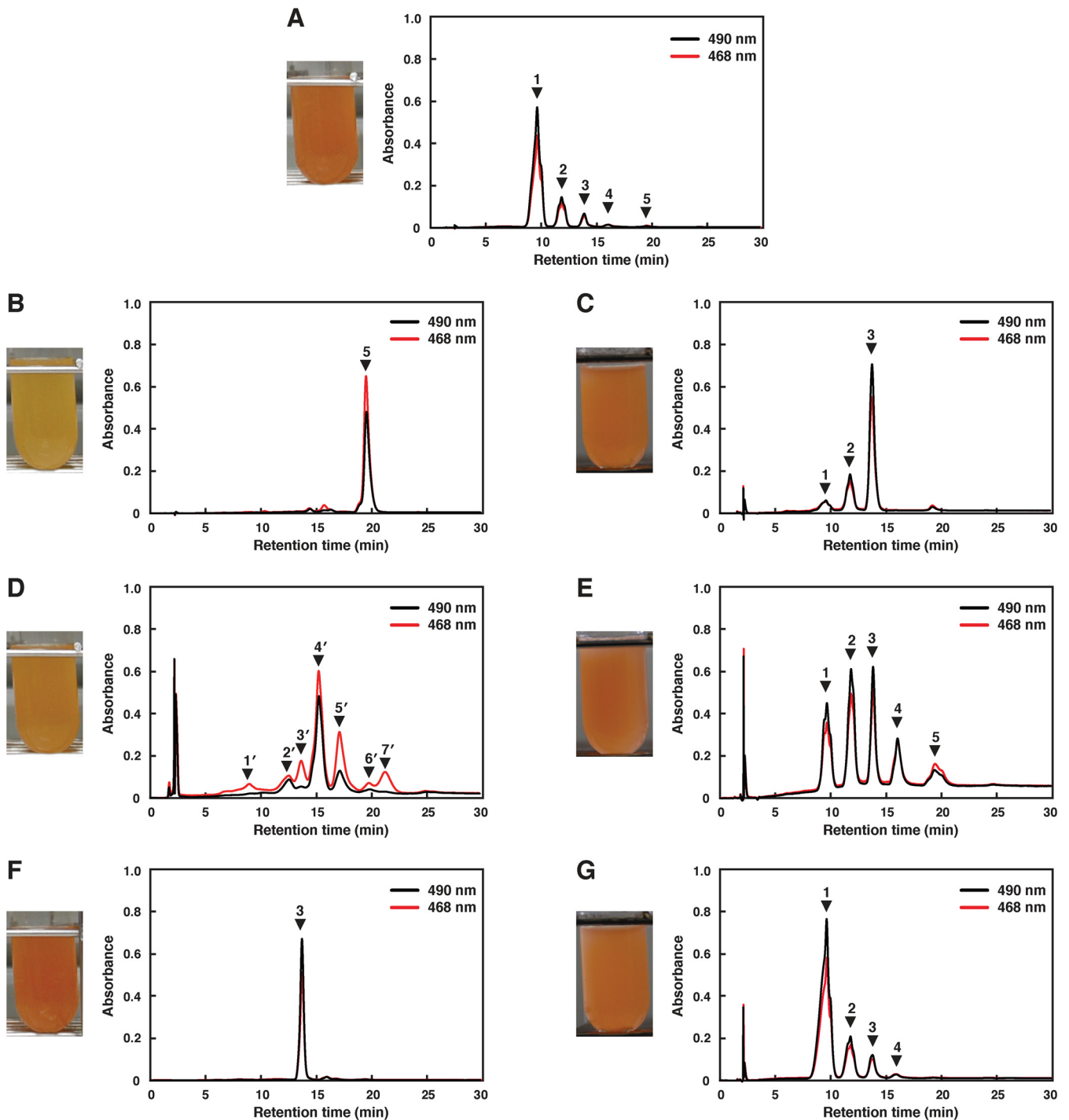


FIG 3 Cell suspension color of the wild type and of the mutants and their genetically modified *Ha. japonica* strains, and HPLC elution profiles of carotenoids extracted from them. (A) Wild type. (B) $\Delta c0506$ strain. (C) $\Delta c0506(pJc0506)$ strain. (D) $\Delta c0507$ strain. (E) $\Delta c0507(pJc0507)$ strain. (F) $\Delta c0505$ strain. (G) $\Delta c0505(pJc0505)$ strain. Peak 1, bacterioruberin (BR); peak 2, monoanhydrobacterioruberin (MABR); peak 3, bisanhydrobacterioruberin (BABR); peak 4, isopentenyldehydrorhodopin (IDR); peak 5, lycopene; peak 4', tetrahydrobisanhydrobacterioruberin (TH-BABR); peak 5', dihydroisopentenyldehydrorhodopin (DH-IDR). Peaks 1', 2', 3', 6', and 7' might be derivatives of lycopene precursors. The eluent was methanol-water (9:1 [vol/vol]) for the first 10 min and then 100% methanol (1.5 ml/min).

growth rate as wild-type *Ha. japonica*. The transcription of *c0507* and *c0505* in the $\Delta c0506$ strain were confirmed by RT-PCR (Fig. 2B, panel b). The disruption of *c0506* did not affect the expression of the upstream *c0507* gene, although the transcription level of the

downstream *c0505* gene was reduced to less than half that in wild-type *Ha. japonica*. The cell suspension of the $\Delta c0506$ strain was yellow in color, while the cell suspension of wild-type *Ha. japonica* was red in color (Fig. 3). Thus, the carotenoids produced were

TABLE 3 Characteristics of carotenoids extracted from the wild type and the mutants and the corresponding genetically modified *Ha. japonica* strains

Strain	Peak no.	Retention time (min)	λ_{\max} (nm) in HPLC eluent	Conjugated double bonds (<i>n</i>)	Molecular mass (<i>m/z</i>)		Carotenoid ^a
					Measured	Calculated	
<i>Ha. japonica</i> WT	1	9.6	469, 492, 525	13	740	740	BR
	2	11.8	469, 491, 522	13	722	722	MABR
	3	13.8	467, 490, 521	13	704	704	BABR
	4	16.0	453, 479, 509	12	620	620	IDR
	5	19.4	442, 468, 499	11	536	536	Lycopene
<i>Ha. japonica</i> $\Delta c0506$	5	19.4	443, 469, 500	11	536	536	Lycopene
<i>Ha. japonica</i> $\Delta c0507$	4'	14.8	441, 466, 498	11	708	708	TH-BABR
	5'	16.7	440, 465, 495	11	622	622	DH-IDR
<i>Ha. japonica</i> $\Delta c0505$	3	13.7	468, 491, 523	13	704	704	BABR
<i>Ha. japonica</i> $\Delta c0506(pJc0506)$	1	9.6	469, 491, 523	13	ND	740	BR
	2	11.8	468, 491, 523	13	ND	722	MABR
	3	13.8	466, 490, 522	13	ND	704	BABR
<i>Ha. japonica</i> $\Delta c0507(pJc0507)$	1	9.6	468, 492, 524	13	ND	740	BR
	2	11.8	469, 491, 522	13	ND	722	MABR
	3	13.8	466, 490, 521	13	ND	704	BABR
	4	16.0	454, 480, 510	12	ND	620	IDR
	5	19.4	443, 468, 499	11	ND	536	Lycopene
<i>Ha. japonica</i> $\Delta c0505(pJc0505)$	1	9.6	468, 492, 525	13	ND	740	BR
	2	11.8	467, 492, 522	13	ND	722	MABR
	3	13.8	466, 490, 520	13	ND	704	BABR
	4	16.0	454, 479, 509	12	ND	620	IDR

^a BR, bacterioruberin; MABR, monoanhydrobacterioruberin; BABR, bisanhydrobacterioruberin; IDR, isopentenyldehydrorhodopin; TH-BABR, tetrahydrobis-anhydrobacterioruberin; DH-IDR, dihydroisopentenyldehydrorhodopin. Carotenoids were analyzed using an HPLC system equipped with a μ Bondapak C₁₈ column (Waters) (3.9 by 300 mm) and were eluted with methanol-water (9:1 [vol/vol]) for the first 10 min and then with 100% methanol (1.5 ml/min). Absorption spectra of the carotenoids were recorded with a photodiode array detector attached to the HPLC apparatus (Fig. 3). The relative molecular masses of the purified carotenoids were measured using an MStation JMS-700 mass spectrometry system (Jeol) in the FAB mode with *m*-nitrobenzyl alcohol as a matrix. ND, not determined.

likely to be different. The carotenoids produced by the $\Delta c0506$ strain were analyzed by HPLC and compared with those of the wild-type *Ha. japonica*. Five peaks were detected in wild-type *Ha. japonica* and had been identified as BR (Fig. 3A, peak 1), MABR (peak 2), BABR (peak 3), IDR (peak 4), and lycopene (peak 5), respectively, in our previous study (Fig. 3 and Table 3) (22). In contrast, only one carotenoid (Fig. 3B, peak 5) was detected in the $\Delta c0506$ strain. This carotenoid exhibited a retention time of 19.4 min and had λ_{\max} values of 443, 469, and 500 nm, which corresponded to an acyclic carotenoid having 11 conjugated double bonds ($n = 11$). This was similar to the results seen with peak 5 of wild-type *Ha. japonica* (representing lycopene, with a retention time of 19.4 min and λ_{\max} values of 442, 468, and 499 nm [$n = 11$]). Mass spectrometric analysis showed that peak 5 of the $\Delta c0506$ strain had an *m/z* value of 536. Based on the retention time, absorbance spectrum, and mass spectrometric results, peak 5 of the $\Delta c0506$ strain was identified as representing lycopene.

In the $\Delta c0506$ strain, no C₄₅ and C₅₀ carotenoids were detected, suggesting that the biosynthetic pathway for carotenoids was discontinued at lycopene. Therefore, we deduced that *c0506* of *Ha. japonica* encodes a functional enzyme that catalyzes the conversion of lycopene to a BR precursor.

There seem to be two types of desaturation reactions in the carotenoid biosynthetic pathway of *Ha. japonica*. In the first reaction, CrtI converts phytoene into lycopene by the introduction of

four double bonds. In the second reaction, CrtD forms double bonds at C-3,4 and C-3',4' of the lycopene derivatives. According to homology analysis results, C0507 shows amino acid sequence homology to both CrtI and CrtD. To investigate the function of C0507, a mutant of *c0507*, designated strain $\Delta c0507$, was constructed by homologous recombination. The $\Delta c0507$ strain showed the same growth rate as wild-type *Ha. japonica*. The transcripts of *c0506* and *c0505* in the $\Delta c0507$ strain were confirmed by RT-PCR, although their transcription levels were reduced to less than half those seen in wild-type *Ha. japonica* (Fig. 2B, panel c). The cell suspension of the $\Delta c0507$ strain was orange in color (Fig. 3D).

The HPLC elution profile of carotenoids produced by the $\Delta c0507$ strain is shown in Fig. 3D. The absorption spectrum indicated that peak 1', peak 2', peak 3', peak 4', peak 5', peak 6', and peak 7' had 9, 10, 9, 11, 11, 7, and 9 conjugated double bonds, respectively. Among these, peak 4' and peak 5', representing the main carotenoids in the $\Delta c0507$ strain, possessed the most abundant conjugated double bonds, suggesting that peak 4' and peak 5' represent the final products or their derivatives. The absorption spectra of peak 4' and peak 5' are shown in Fig. 4. Mass spectrometric analysis also showed that peak 4' and peak 5' had *m/z* values of 708 (4 mass units more than BABR) and 622 (2 mass units more than IDR), respectively (Table 3). Based on these data, peak 4' and peak 5' were identified as representing TH-BABR and

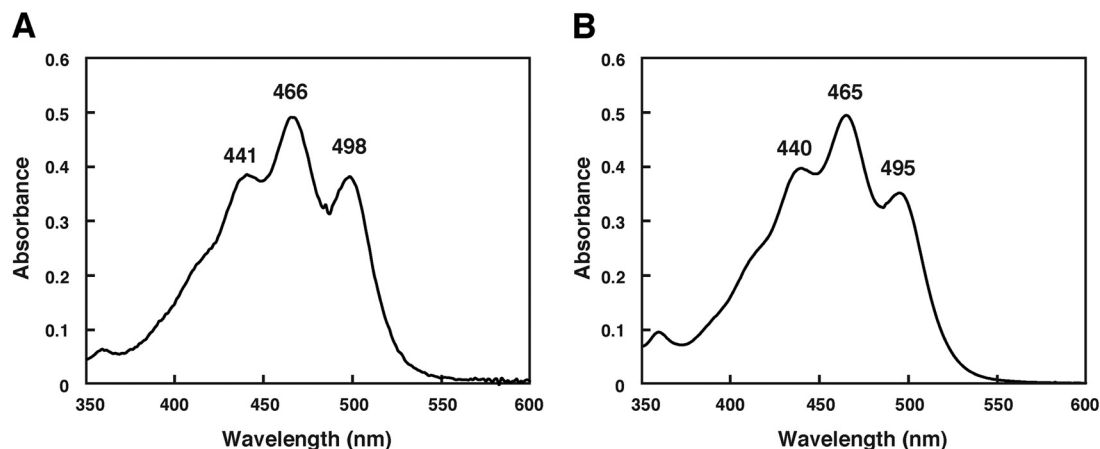


FIG 4 Absorption spectra of the carotenoids extracted from the $\Delta c0507$ strain. (A) Peak 4' (tetrahydrobisnhydrobacterioruberin [TH-BABR]; λ_{\max} = 441, 466, and 498 nm). (B) Peak 5' (dihydroisopentenyldehydrorhodopin [DH-IDR]; λ_{\max} = 440, 465, and 495 nm). Spectra were measured using a photodiode-array detector attached to the HPLC apparatus, with a mobile phase of 100% methanol.

dihydroisopentenyldehydrorhodopin (DH-IDR; C₄₅), respectively. Since phytoene did not accumulate in the $\Delta c0507$ strain, the desaturation reactions that convert phytoene into lycopene were not catalyzed by C0507, or the desaturation reactions were catalyzed cooperatively by C0507 and other phytoene desaturases. We reached that conclusion because, in a *Bradyrhizobium* strain, two distinct *crt* gene clusters are involved in the synthesis of its carotenoids (spirilloxanthin and canthaxanthin). Each cluster contains the genes *crtE* (GGPP synthase), *crtB*, and *crtI*, leading to the common precursor lycopene (39). In addition, the main products of the $\Delta c0507$ strain, TH-BABR and DH-IDR, lacked the double bonds at C-3,4 and C-3',4'. Therefore, we deduced that C0507 functions as a CrtD, involved in the desaturation reactions that form double bonds at C-3,4 of DH-IDR and C-3',4' of DH-BABR. This is the first time that a CrtD has been identified functionally in archaea. Peak 1', peak 2', peak 3', peak 6', and peak 7' remain unexplained. Because the number of conjugated double bonds in these compounds was less than 11 and because the number of conjugated double bonds in lycopene was 11, these peaks might represent derivatives of lycopene precursors.

The biosynthetic pathway for carotenoids in the $\Delta c0506$ strain was discontinued at lycopene, and the $\Delta c0507$ strain accumulated DH-IDR and TH-BABR. C0506 was suggested to catalyze the conversion of lycopene to DH-IDR. The conversion contains two reactions, *viz.*, introduction of a C₅ isoprene unit at C-2 and hydration at C-1,2 of the terminal of lycopene (Fig. 1). Therefore, C0506 in *Ha. japonica* is likely to be a bifunctional lycopene elongase and 1,2-hydratase. The *lye* gene of *Hb. salinarum*, a homolog of *c0506*, was expressed in *E. coli*, and the gene product catalyzes the conversion of lycopene to TH-BABR (10). In this conversion, introduction of a C₅ isoprene unit at C-2 and hydration at C-1,2 of the terminal of lycopene were both catalyzed by *Lye*. This result supports our prediction that C0506 would be a bifunctional enzyme. So we redefined *Lye* as a bifunctional enzyme and designated C0506 from *Ha. japonica* [*Lye*].

In order to confirm the function of C0506 and C0507, an *in vivo* complementation study was performed. Plasmid pJc0506, containing the intact *c0506* gene, was introduced into the $\Delta c0506$ strain, and the transformant was named the $\Delta c0506$ (pJc0506) strain. HPLC analysis revealed that the biosynthesis of the final BR

product was restored (Fig. 3C and Table 3). However, the main products were BABR and MABR. That was probably a result of the fact that the transcription level of the downstream *c0505* gene is still low in the complemented strain. Plasmid pJc0507, containing the intact *c0507* gene, was introduced into the $\Delta c0507$ strain. The transformant $\Delta c0507$ (pJc0507) restored the production of BR (Fig. 3E and Table 3). Since the transcription levels of the downstream genes, *c0506* and *c0505*, are probably still low in the complemented strain, the intermediates (MABR, BABR, IDR, and lycopene) accumulated and the proportion of final product diminished in the $\Delta c0507$ (pJc0507) transformant.

Analysis of *c0505*. The hydroxyl groups at C1 and C1' of lycopene were introduced by C0506. However, the enzyme that introduces hydroxyl groups to the C3'' and C3''' of BABR had not been identified. C0505, the homolog of CruF, was selected as a candidate enzyme for this reaction. Its mutant was constructed by homologous recombination and designated strain $\Delta c0505$. The $\Delta c0505$ strain also showed the same growth rate as wild-type *Ha. japonica*. The transcriptions of *c0507* and *c0506* in the $\Delta c0505$ strain were confirmed by RT-PCR, and the disruption of *c0505* did not affect the expression of the upstream genes in the same cluster (Fig. 2B, panel d).

The cell suspension of the $\Delta c0505$ strain was red in color. In HPLC, only one peak (Fig. 3F, peak 3) was detected in the $\Delta c0505$ strain, and the peak exhibited a retention time of 13.7 min and λ_{\max} values of 468, 491, and 523 nm ($n = 13$) (Table 3). Mass spectrometric analysis showed that peak 3 of the $\Delta c0505$ strain had an m/z value of 704, compatible with that of BABR. Based on these data, peak 3 of the $\Delta c0505$ strain was identified as representing BABR. These data demonstrated that the carotenoid biosynthetic pathway of the $\Delta c0505$ strain was discontinued at BABR and indicated that C0505 catalyzes the reaction that introduces hydroxyl groups to C3'' and C3''' of BABR to generate BR (Fig. 1). Therefore, C0505 in *Ha. japonica* is a C₅₀ carotenoid 2'',3''-hydratase.

The *in vivo* complementation study was also assessed using the intact *c0505* gene-introduced transformant $\Delta c0505$ (pJc0505). HPLC analysis revealed that the carotenoid composition of the $\Delta c0505$ (pJc0505) transformant was restored and was similar to that of wild-type *Ha. japonica* (Fig. 3G and Table 3).

DISCUSSION

The C₅₀ carotenoid BR is known to accumulate in several extremely halophilic archaea (15–19), but the identity of the responsible biosynthetic pathway remains unclear. In this study, we have identified three main genes encoding enzymes responsible for the biosynthesis of BR from lycopene for the first time.

The biosynthetic pathway for carotenoids in the $\Delta c0506$ strain was discontinued at lycopene, and the $\Delta c0507$ strain accumulated DH-IDR and TH-BABR; hence, C0506 was suggested to be a bifunctional lycopene elongase and 1,2-hydratase, catalyzing the reaction that introduces a C₅ isoprene unit and a hydroxyl group to the terminal of lycopene. In strain $\Delta c0507$, TH-BABR and DH-IDR were detected. Therefore, C0507 was proposed to be a carotenoid 3,4-desaturase and to be involved in the desaturation reactions that form double bonds at C-3,4 of DH-IDR and C-3',4' of DH-BABR. It seems that two types of possible biosynthetic pathways from lycopene to BABR exist in wild-type *Ha. japonica*. The first type of pathway involves immediate conversion of DH-IDR, generated from lycopene by C0506 to IDR by C0507. Subsequently, C0506 further introduces a C₅ isoprene unit and a hydroxyl group to the other terminal of IDR to generate the C₅₀ carotenoid DH-BABR, which is then desaturated by C0507 to form BABR. The second type involves C0506-mediated catalysis of reactions that introduce two C₅ isoprene units and two hydroxyl groups to each end of the C₄₀ lycopene to generate the C₅₀ TH-BABR, which is then converted to BABR by C0507. IDR was detected as an intermediate, but TH-BABR was not detected in the wild-type *Ha. japonica* strain. Thus, it can be suggested that the former is the main pathway in wild-type *Ha. japonica*. On the other hand, it has been proposed that the biosynthetic pathway of BABR from lycopene in *Hb. salinarum* involves the latter pathway (10); therefore, the carotenoid biosynthetic pathway in *Ha. japonica* is possibly different from that in *Hb. salinarum*.

Based on the proposed carotenoid and retinal biosynthetic pathways in *Ha. japonica* (Fig. 1), the $\Delta c0506$ strain was expected to accumulate β -carotene or retinal in addition to lycopene. However, only lycopene was detected when the $\Delta c0506$ strain was grown in the dark. Our previous study showed that the transcription of the cruxrhodopsin (a retinal protein) gene is regulated by high light intensity (23), and β -carotene was detected when *Ha. japonica* was grown in light (unpublished data). So it is possible that β -carotene cannot be accumulated when the $\Delta c0506$ strain is grown in the dark.

The carotenoid 1,2-hydratase catalyzes the synthesis of some carotenoids that contain hydroxyl groups by hydration at the C-1,2 or C-1',2' double bond. Two types of carotenoid 1,2-hydratases (CrtC and CruF) have been discovered to date. The first type is the CrtC-type carotenoid 1,2-hydratase that has been primarily found in purple bacteria, such as *Rhodobacter capsulatus*, which produces spheroidene (35, 40). The second type is the CruF-type carotenoid 1,2-hydratase that has been found in bacteria such as *Synechococcus* sp. strain PCC 7002, which produces myxol-2' fucoside (41), and two species of *Deinococcus* (38). C0505, the C₅₀ carotenoid 2'',3''-hydratase from *Ha. japonica*, is a homolog of the CruF-type carotenoid 1,2-hydratase. This is the first time that an example of CruF has been functionally identified in archaea, although C0505 catalyzes hydration at the C-2'',3'' and C-2''',3''' double bond of BABR instead of at the C-1,2 or C-1',2' double bond of lycopene. The carotenoid hydratases of the same CruF

type from different bacteria would have their own substrate specificities.

In a previous genomic sequence analysis of *Ha. japonica*, *c0507* revealed its sequence homology to *crtI* of *P. ananatis* (34). However, our results demonstrated that C0507, the CrtD equivalent, is involved in desaturation reactions that form double bonds at C-3,4 of DH-IDR and C-3',4' of DH-BABR. Thus, the CrtI that converts phytoene to lycopene has not been identified. In addition, C0184, C1220, C1219, and C1158 from *Ha. japonica* also show amino acid sequence homology to GGPP synthase (CrtE), phytoene synthase (CrtB), lycopene β -cyclase (CrtY), and β -carotene cleavage dioxygenase (Brp), respectively. Additionally, further studies are necessary to investigate the remaining unidentified carotenoid biosynthetic enzymes in order to elucidate the complete carotenoid biosynthetic pathway in *Ha. japonica*.

It has been suggested that the antioxidant capacity of carotenoids is linked to the number of conjugated double bonds and hydroxyl groups (3, 4, 42). BR, which contains 13 conjugated double bonds and four hydroxyl groups, is regarded to be an effective free-radical scavenger. Our previous study has shown that the free-radical scavenging capacity of BR is much higher than that of β -carotene, which contains 11 conjugated double bonds (22). Moreover, BR is a dipolar C₅₀ carotenoid, and it was suggested to act as a “rivet” in the cell membrane, to increase its rigidity and mechanical strength (7).

In summary, we have identified three carotenoid biosynthetic enzymes and have elucidated their functions. Among them, CrtD and CruF were found in an archaeon for the first time. Our results revealed the biosynthetic pathway responsible for production of BR from lycopene in *Ha. japonica* and provide a basis for investigating carotenoid biosynthetic pathways in other extremely halophilic archaea. Elucidation of the carotenoid biosynthetic pathway in *Ha. japonica* may also prove useful for producing the C₅₀ carotenoid BR efficiently by employing genetically modified haloarchaeal strains.

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