Substrate Specificity of Purified Recombinant Chicken \(\beta\)-Carotene 9',10'-Oxygenase (BCO2)*

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Carlo dela Seña 1,5, Jian Sun 1, Sureshbabu Narayanasamy 1, Kenneth M. Riedel 1, Yan Yuan 1, Robert W. Curley, Jr. 1, Steven J. Schwartz 1, and Earl H. Harrison 1,5

From the 1 Department of Human Nutrition, 5 Ohio State Biochemistry Program, 6 Department of Food Science and Technology, and 5 College of Pharmacy, The Ohio State University, Columbus, Ohio 43210

Provitamin A carotenoids are oxidatively cleaved by \(\beta\)-carotene 15,15'-dioxygenase (BCO1) at the central 15-15' double bond to form retinal (vitamin A aldehyde). Another carotenoid oxygenase, \(\beta\)-carotene 9',10'-oxygenase (BCO2) catalyzes the oxidative cleavage of carotenoids at the 9'-10' bond to yield an ionone and an apo-10'-carotenoid. Previously published substrate specificity studies of BCO2 were conducted using crude lysates from bacteria or insect cells expressing recombinant BCO2. Our attempts to obtain active recombinant human BCO2 expressed in vertebrates were unsuccessful. We have expressed recombinant chicken BCO2 in the strain \(\varepsilon\) coli BL21-Gold (DE3) and purified the enzyme by cobalt ion affinity chromatography. Like BCO1, purified recombinant chicken BCO2 catalyzes the oxidative cleavage of the provitamin A carotenoids \(\beta\)-carotene, \(\alpha\)-carotene, and \(\beta\)-cryptoxanthin. Its catalytic activity with \(\beta\)-carotene as substrate is at least 10-fold lower than that of BCO1. In further contrast to BCO1, purified recombinant chicken BCO2 also catalyzes the oxidative cleavage of 9-cis-\(\beta\)-carotene and the non-provitamin A carotenoids zeaxanthin and lutein, and is inactive with all-trans-lycopene and \(\beta\)-apo-carotenoids. Apo-10'-carotenoids were detected as enzymatic products by HPLC, and the identities were confirmed by LC-MS. Small amounts of 3-hydroxy-\(\beta\)-apo-8'-carotenal were also consistently detected in BCO2-\(\beta\)-cryptoxanthin reaction mixtures. With the exception of this activity with \(\beta\)-cryptoxanthin, BCO2 cleaves specifically at the 9'-10' bond to produce apo-10'-carotenoids. BCO2 has been shown to function in preventing the excessive accumulation of carotenoids, and its broad substrate specificity is consistent with this.

There are three members of the carotenoid oxygenase family expressed in vertebrates. \(\beta\)-Carotene 15,15'-dioxygenase (BCO1)2 catalyzes the oxidative cleavage of provitamin A carotenoids to form retinal (vitamin A aldehyde). RPE65 is an isomerohydrolase that catalyzes the conversion of all-trans-retinyl esters to 11-cis-retinol and is an important component of the visual cycle (1–5). \(\beta\)-Carotene 9',10'-oxygenase (BCO2) catalyzes the oxidative cleavage of carotenoids at the 9'-10' or 9'-10 bond to yield an ionone and an apo-10'-carotenoid. Studies in mice suggest that BCO2 cleavage prevents oxidative stress from carotenoid accumulation, especially in the mitochondria (6). The provitamin A carotenoids \(\beta\)-carotene and \(\beta\)-cryptoxanthin are also cleaved by BCO2 to produce \(\beta\)-apo-10'-carotenal (6–8), which can then be oxidatively cleaved by BCO1 to produce retinal (8, 9).

Unlike BCO1, BCO2 has also been shown to cleave non-provitamin A carotenoids such as zeaxanthin and lutein to produce 3-hydroxy-apo-10'-carotenals (6, 7). The significance of these compounds in mammals is unknown. However, (3R)-hydroxy-\(\beta\)-apo-10'-carotenol, also known as galloxanthin, has been identified as one of the dominant carotenoids in quail retina (10). This is formed from the reduction of (3R)-hydroxy-\(\beta\)-apo-10'-carotenol, which can be formed from BCO2 cleavage of \(\beta\)-cryptoxanthin, zeaxanthin, or lutein. Galloxanthin is found in the lipid droplets of avian cones that is necessary for the filtering of the short wavelength UV light in the avian eye. This process is evolutionarily important for the enhanced color vision of birds as compared with other taxa. Ferret hepatic homogenates have been shown to reduce (3R)-hydroxy-\(\beta\)-apo-10'-carotenal to the alcohol (7). The production of \(\beta\)-apo-10'-carotenol has been shown in vivo. BCO1 KO mice, which have elevated expression of BCO2, accumulate \(\beta\)-apo-10'-carotenol on a diet with only \(\beta\)-carotene as the sole source of apocarotenoids (11). In humans, the dominant carotenoids in the retina are lutein, 3'-epilutein, and zeaxanthin (10). Given that BCO2 is also highly expressed in the human retinal epithelium (12), it is interesting that BCO2 cleavage products of xanthophylls are not the dominant carotenoids. It is possible that human BCO2 and avian BCO2 have different substrate specificities, or another regulatory mechanism(s) exists to favor the BCO2 cleavage of lutein and zeaxanthin in avian retina but not in humans (13, 14).

The published BCO2 substrate specificity studies as of the time of writing were done with crude lysates of bacteria and insects cells expressing recombinant BCO2 (6, 7, 14). In this work, we describe the substrate specificity and kinetics of \(\beta\)-carotene cleavage of purified recombinant chicken BCO2. We show that the enzyme reacts with \(\beta\)-carotene (albeit with
Expression and Purification of Recombinant BCO2 cDNA of chicken BCO2 with an N-terminal hexahistidine tag formed into isoforms

Plasmids

Plasmids for bacterial expression containing human BCO2 isoforms a and d with a C-terminal His tag were purchased from GeneCopoeia. A pET-28a plasmid vector containing the cDNA of chicken BCO2 with an N-terminal hexahistidine tag was a gift from Drs. Matthew Toomey and Joseph Corbo of Washington University at St. Louis.

Expression and Purification of Recombinant BCO2

For human BCO2 isoforms a and d, the plasmid was transformed into E. coli BL21-Gold (DE3) (Stratagene) or T7 Express (New England Biolabs) according to the manufacturer’s instructions. For chicken BCO2, the plasmid was transformed into E. coli BL21-Gold (DE3). The transformed bacterial cells were grown in LB broth (Sigma-Aldrich) at 30 °C to an optical density (OD) of 0.5–0.7. The incubator temperature was then lowered to 16 °C, and the cells were harvested by centrifugation. The cell pastes were frozen at −80 °C and thawed prior to lysis. The protein was purified using the same protocol for recombinant human BCO1 using cobalt ion affinity chromatography as in our previous publication (9).

In Vitro BCO2 Assay

The in vitro BCO2 enzyme assay was based on the method of Amengual et al. (6). All substrates were tested at a concentration of 20 μM at 37 °C with the indicated incubation time and protein concentration. The working enzyme solution for a single reaction was prepared by combining the required amount of enzyme and storage buffer to a total volume of 100 μL, 40 μL of reaction buffer (250 mM Tricine-KOH, pH 8.0 at 37 °C, 625 mM NaCl), 4 μL of 500 μM FeSO₄, 2 μL of 0.5 M triis(2-carboxyethyl)phosphine, and 14 μL of deionized water. The substrate solution was prepared by first mixing the required amount of carotenoid in hexanes or ethanol, 9 μL of 10 mM α-tocopherol, 225 μL of 4% octyl thioglycoside in ethanol, and 400 μL of acetone. The mixture was dried under nitrogen and stored at −20 °C. The next day, immediately before use, the residue was brought up to ice bath temperature, redissolved in 200 μL of acetone, dried down under nitrogen, and redissolved in 180 μL of deionized water. The substrate and enzyme solutions were incubated at 37 °C for 5 min, and the reaction was initiated by adding 40 μL of the substrate to the enzyme for a total reaction volume of 200 μL. The reaction was quenched with 50 μL of 37% formaldehyde and incubated for a further 10 min at 37 °C. Then 500 μL of 1:1 (v/v) acetonitrile-isopropyl alcohol was added. The resulting mixture was filtered through a 0.22-μm syringe filter, and 100 μL was subjected to HPLC. Time 0, no enzyme, and heat-denatured enzyme controls were used.

Analytical HPLC Methods

Method A—This is identical to Method A as described in our previous publication (9) using a Zorbax Eclipse XDB-C$_8$ LC column (4.6 × 50 mm, 1.8 μm; Agilent) with or without a Zorbax Eclipse XDB-C$_{18}$ guard column (4.6 × 12.5 mm, 5 μm; Agilent).

Method B—For lutein, the chromatography described above cannot separate the 3-hydroxy-β-apo-10’-carotenal and 3-hydroxy-α-apo-10’-carotenal produced from its reaction with BCO2. However, these were separated by the following chromatography system: column, YMC Carotenoid (YMC Co.), 4.6 × 150 mm, 5-μm particle size; flow rate, 0.9 ml/min; column temperature, 35 °C; solvent, 95% methanol: 5% water; elution time, 12 min. Elution was monitored at 380, 414, and 453 nm.

Method C—The reaction products and isomers of the parent carotenoids were analyzed using a YMC Carotenoid S-3 column (4.6 × 250 mm, 3-μm particle size), a flow rate of 1.4 ml/min, and a column temperature of 35 °C. The following elution profile was used with methanol (solvent A) and methyl tert-butyl ether (solvent B): gradient from 5 to 25% B over 30 min, gradient from 25 to 5% solvent B over 30 s, and 5% solvent B for 2.5 min. Elution was monitored at 450 and 360 nm.

LC-MS Analysis

HPLC was performed with Method A described above with the following modifications. First, formic acid was used instead of ammonium acetate. Second, the column used was also Zorbax XDB-C$_{18}$ but with dimensions of 4.6 × 150 mm and 5-μm particle size.

The HPLC was interfaced with a quadrupole time-of-flight (Q-TOF) mass spectrometer (Q-TOF Premier, Micromass, Manchester, UK) via an atmospheric pressure chemical ionization probe. β-Carotene, β-apo-8’-carotenal, β-apo-10’-carotenal, β-apo-12’-carotenal, β-apo-14’-carotenal, and retinal were ionized in atmospheric pressure chemical ionization negative mode as their respective radical anions, 536.438, 416.310, 390.310, 350.260, 310.230, and 284.210 m/z. The Q-TOF system allowed for quantitative detection with the confidence of accurate mass typically being 1 ppm. Mass spectra were acquired in V-mode (~8000 resolution) from 50 to 1000 m/z with a scan time of 1 s, peak centroiding, and enhanced duty cycle enabled for the parent m/z. At intervals of 30 s, a 0.1-s lock spray scan was acquired with leucine enkephalin as the lock spray compound (554.2615 m/z) to correct for minor deviations in calibration due to temperature fluctuations. Prior to...
analysis, the Q-TOF was fully calibrated from 114 to 1473 m/z using a solution of sodium formate. The resultant MS spectra were acquired and integrated with MassLynx software, V4.1 (Micromass). Source parameters were as follows: 30-μA corona current; 550 °C probe; 110 °C source block; 35-V cone; 100 liter/h cone gas (N2); 400 liter/h desolvation gas (N2); and collision energy of 8 eV (non-fragmenting) with argon as the collision-induced dissociation gas ($4.2 \times 10^{-3}$ millibar).

**Results**

**Recombinant Human BCO2 Expressed in E. coli Is Inactive**—The longest isoform of human BCO2 is isoform a (accession number NP_114144.4), corresponding to a 579-amino acid peptide. Isoform d (accession number NP_001243327.1) corresponds to a 506-amino acid peptide and is the same sequence used by Kim et al. (16) in their study claiming to produce active recombinant human BCO2 in E. coli strain ER2566 (equivalent to T7 Express). Interestingly, this isoform is missing one of the conserved histidines that coordinate the catalytic ferrous ion (Fig. 1) (17).

Both isoforms have a behavior similar to that of recombinant human BCO1 expressed in E. coli (9): high rates of expression favor formation of inclusion bodies, but the small amount of His-tagged protein can be recovered from the supernatant by cobalt ion affinity chromatography. However, there is no detectable enzymatic cleavage activity from the crude superna-

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**FIGURE 1.** Sequence alignment of BCO2, BCO1, and RPE65. The GKAA insertion that differentiates human BCO2 isoforms from other BCO2 orthologs is highlighted in green. The conserved residues (four histidines and one glutamate) that were identified to be crucial for activity in mouse BCO1 (17) are highlighted in violet. The sequence alignment was generated by ClustalW (37), and the image was generated by ESPript (38) (green and violet highlights were added separately). The NCBI accession numbers are as follows: human BCO2 isoforms a–e, NP_114144.4, NP_001032367.2, NP_001243326.1, NP_001243327.1, and NP_001243329.1, respectively; macaque BCO2, AFE80390.1; mouse BCO2, NP_573480.1; chicken BCO2 isoforms x1–x3, XP_004948199.1, XP_417929.2, and XP_015153786.1, respectively; human BCO1, NP_059125.2; mouse BCO1, NP_067461.2; human RPE65, NP_000320.1; bovine RPE65, NP_776878.1. The dots are space markers for 10 amino acids of the topmost sequence.
Substrate Specificity of Chicken BCO2

![Image](http://www.jbc.org/)

**FIGURE 2. Purification of recombinant chicken BCO2 using cobalt affinity column chromatography.** The SDS-polyacrylamide gel on the left shows the elution of recombinant His-tagged human BCO2 (theoretical mass, 64,202 Da) from the cobalt column by increasing concentrations of imidazole. Only the fractions that show a single band (lanes 5–7) were combined and desalted into the final storage buffer. The SDS-polyacrylamide gel on the right shows the combined fractions and the protein obtained after desalting.

-tant and the purified protein from either strain of *E. coli* when tested with β-carotene, β-cryptoxanthin, and lutein. Furthermore, we did not detect any activity when the *in vitro* assay was conducted in the same manner as BCO1 (9) or by the method of Amengual et al. (6). The inactivity of recombinant human BCO2 expressed in *E. coli* has been reported by others (13, 14).

**Expression and Purification of Recombinant Chicken BCO2—**

The pET28a-chicken BCO2 clone is similar to isoform x2 of *Gallus gallus* BCO2 mRNA in PubMed (accession number XP_417929.2; corresponding to a 579-amino acid peptide), but the first 31 amino acids of the latter (MPSSQERMFSKIAL-SAVSILLANLRLHLLSSL) are replaced with 16 amino acids (MGSSSHHHHHSSGGVLPRGS) corresponding to the His tag. The theoretical molecular mass of the protein is 64,202 Da.

As with recombinant human BCO2 and BCO1, recombinant chicken BCO2 is not very soluble under the expression conditions in BL21-Gold (DE3), but the small amount of soluble protein was easily recovered to a high degree of purity by cobalt ion affinity chromatography as shown in Fig. 2.

**Kinetics of Cleavage of β-Carotene by Chicken BCO2—**

Kinetic data for the cleavage of all-trans-β-carotene by purified recombinant chicken BCO2 is shown in Fig. 3. The production of β-apo-10′-carotene is linear for up to 40 min (Fig. 3A) and up to 5 μg of BCO2/200-μl reaction (Fig. 3B). The substrate-velocity curve (Fig. 3C) fits the Michaelis-Menten equation with a *V*~max~ of 0.93 nmol of β-apo-10′-carotene/mg of BCO2/h (standard error of regression, 0.05; two independent experiments) and *K*~m~ of 2.03 μM (standard error of regression, 0.27; two independent experiments) obtained from nonlinear regression using GraphPad Prism 4. From these values, the calculated *k*~cat~ is 9.9 × 10^−4~ min~−1~ and *k*~cat~/*K*~m~ (catalytic efficiency) is 487 m~−1~ min~−1~. This catalytic efficiency is at least 10 times lower than what we have previously measured for purified recombinant human BCO1 (6098 m~−1~ min~−1~) (9).

**Chicken BCO2 Cleaves Full-length β-Ring Carotenoids at the 9′-10′ Bond—**
The activity of purified recombinant chicken BCO2 was tested with the major dietary carotenoids (all-trans-β-carotene, 9-cis-β-carotene, α-carotene, β-cryptoxanthin, zeaxanthin, lutein, and lycopene) and β-apocarotenals (β-apo-8′-carotene, β-apo-10′-carotene, β-apo-12′-carotene, and β-apo-14′-carotene) using a 20 μM concentration of each substrate and 10 μg of BCO2/200-μl reaction with a 2-h reaction time. Time 0, no enzyme, and heat-denatured enzyme controls were conducted for all reactions.

Because 9-cis-β-carotene is an asymmetrical molecule, we expected to see two apocarotenoid cleavage products, all-trans- and 9-cis-β-apo-10′-carotene (Fig. 4). In our original chromatography (HPLC Method A), there was only one enzymatic product peak observed from all-trans- and 9-cis-β-carotene, and these have the same retention time as standard all-trans-β-apo-10′-carotene (Fig. 5, A–C). However, when using a different chromatographic system (HPLC Method C), the cleavage product of 9-cis-β-carotene elutes earlier (about 0.1 min) than the cleavage product of all-trans-β-carotene and standard all-trans-β-apo-10′-carotene (Fig. 5D). The peak of the cleavage product of 9-cis-β-carotene shows a very small shoulder to the right. However, the heat-denatured control shows that there is a distorted peak at that elution time. Thus, we cannot rule out the possibility that some all-trans-β-apo-10′-carotene is also produced by enzymatic cleavage. Nevertheless, it is quite clear that the cleavage product of 9-cis-β-carotene is not solely all-trans-β-apo-10′-carotene. It is also worth noting that we cannot detect any conversion of the parent 9-cis-β-carotene to the all-trans form (and vice versa) during the reaction (Fig. 5E). Thus, the major (if not the only) product of BCO2 cleavage of 9-cis-β-carotene under the test conditions is 9-cis-β-apo-10′-carotene.

α-Carotene is an asymmetrical molecule, and BCO2 cleavage can produce two products, β-apo-10′-carotene and α-apo-10′-carotene (Fig. 6). Consequently, we observed two product peaks (Fig. 7A), one with the same retention time as β-apo-8′-carotene (peak b) and another earlier eluting peak that corresponds to α-apo-8′-carotene (peak a).

β-Cryptoxanthin is also an asymmetrical molecule, and BCO2 cleavage can produce two products, β-apo-10′-carotene and 3-hydroxy-β-apo-10′-carotene (Fig. 6). We observed three product peaks (Fig. 7B). β-Apo-10′-carotene (peak c) was identified by its identical retention time to the standard compound and LC-MS, and 3-hydroxy-β-apo-10′-carotene (peak c) was identified by LC-MS. 3-Hydroxy-β-apo-8′-carotene, a product of 7-8 double bond cleavage, was identified by LC-MS (peak d). Note that because of the response range used in the figure, this peak appears small, but its height is comparable with that of the product peak for all-trans β-carotene (Fig. 5A). The UV-visible spectra of peak d is also consistent with 3-hydroxy-β-apo-8′-carotenal. The chromophore of carotenoids and apocarotenoids is the conjugated double bond system, and hydroxylation that does not disrupt the latter does not cause large shifts in *λ*~max~. Thus, the UV spectra of 3-hydroxy-β-apo-10′-carotenal and β-apo-10′-carotenol produced from the reaction and that of standard β-apo-10′-carotenol show the same *λ*~max~ (Fig. 8, left column). However, the UV spectrum of peak d is more similar to that of standard β-apo-8′-carotenol rather than that of β-apo-10′-carotenol (Fig. 8, right column).

A substrate-velocity curve was done for β-cryptoxanthin (Fig. 9). However, apart from β-apo-10′-carotenol and β-apo-8′-carotenol, we do not have authentic standards for other apocarotenoid cleavage products necessary for their quantification.
Thus, the 3-hydroxy-β-apocarotenal product peaks were estimated by using the analogous β-apocarotenals. Even though the hydroxyl group is not expected to cause large changes in \( \lambda_{\text{max}} \) and extinction coefficients for as long as the conjugated system is not disrupted (18), we cannot guarantee that the instrument response is virtually the same. Thus, kinetic parameters cannot be reliably calculated. Even if the standards were available, the data cannot be fit to the classic Michaelis-Menten model because the enzyme cleaves the substrate at three positions, and each position is presumably a separate binding mode.

Zeaxanthin is a symmetrical molecule, and BCO2 cleavage will yield 3-hydroxy-α-apo-10′-carotenal (Fig. 6). The original chromatography used (HPLC Method A) shows only one product peak (Fig. 7D, peak g). However, HPLC Method B was able to separate the two products (Fig. 10, top panel). 3-Hydroxy-β-apo-10′-carotenal was distinguished from the α isomer by its identical retention time to the 3-hydroxy-β-apo-10′-carotenal produced from BCO2 cleavage of zeaxanthin (Fig. 10, bottom panel).

It has been reported that murine BCO2 can cleave the two ionone rings from the carotenoid to yield rosafluene dialdehyde (6). However, by identification of the masses of the cleavage products by LC-MS, we did not find evidence of its formation even with a 2-h reaction time and 25 \( \mu \text{g} \) of protein/200- \( \mu \text{l} \) reaction. Also, incubation of β-apo-10′-carotenal with chicken BCO2 did not show any evidence of a reaction. We also did not detect any reaction between chicken BCO2 and β-apo-8′-carotenal, β-apo-12′-carotenal, β-apo-14′-carotenal, and lycopene (chemical structures shown in Fig. 11).

Discussion

Products of excentric cleavage of carotenoids are important in the production of plant hormones such as abscisic acid and strigolactone (19). For vertebrates, the central cleavage product of provitamin A carotenoids, retinal, is still the most important in terms of function. Our finding that the central bond cleavage enzyme BCO1 cleaves only provitamin A carotenoids (except lycopene) specifically at the 15-15′ bond to yield retinal (9) is only consistent with the latter’s biological importance. Thus, the existence of a carotenoid 9′,10′-oxygenase in vertebrates is interesting. Current literature suggests that the primary function of BCO2 in mice is to prevent oxidative stress from accumulation of carotenoids in the mitochondria (6), and so far, a
specific biological activity for β-apo-10'-carotenoids in vertebrates has not been identified (20) apart from their provitamin A activity (9, 21, 22).

The inactivity of human BCO2 in the in vitro assay systems tested can be explained by two possibilities. One is that it really is inactive or has very low activity, which can explain why full-length xanthophylls (lutein, 3'-epilutein, and zeaxanthin) are found in the human macula (10) despite BCO2 being highly expressed in the human retinal epithelium (12). Li et al. (13) noted four residues in human BCO2 (GKAA, positions 169–172; Fig. 1) that are not present in murine BCO2. However, expression of human BCO2 with the GKAA deletion in E. coli engineered to synthesize zeaxanthin failed to decrease the yellow color of the cells (an indication of carotenoid cleavage) unlike murine BCO2. In contrast, insertion of the GKAA into murine BCO2 resulted in retention of the yellow color. The GKAA sequence is also absent in all isoforms of chicken BCO2 (Fig. 1). However, the GKAA sequence is present in macaque BCO2, and activity was observed for heterologously expressed macaque BCO2 with an N terminus truncation (14).

The other possibility is that because it is associated with the inner mitochondrial membrane, as demonstrated by Palczewski et al. (23), it becomes misfolded during lysis or is missing the interactions with the membrane or membrane proteins necessary for activity. A similar phenomenon is observed for the related enzyme RPE65, which is also membrane-associated (24) and inactive when purified, but the activity is restored when it is reassociated with liposomes (25). In the study by Palczewski et al. (23), β-apo-10'-carotenol is produced when human BCO2 is expressed in E. coli engineered to synthesize β-carotene, possibly because BCO2 is still properly folded within the cell and/or has the necessary interactions with membranes and/or other proteins. Thus, the possibility that human BCO2 is active in vivo still cannot be completely ruled out.

As mentioned in our results, the recombinant chicken BCO2 we used differs from the chicken BCO2 isoform x2 in the N terminus, which is where the mitochondrial targeting sequence, if any, would be located (26). This may have been a happy accident that made the protein more amenable to purification. However, Li et al. (13) replaced the N-terminal 197 amino acids of human BCO2 with the N-terminal 150 amino acids of murine BCO2, but this did not result in an observable cleavage of zeaxanthin in their cell-based assay. They also substituted a range of amino acid residues of human BCO2 with the corresponding residues of mouse BCO2, but this failed to confer detectable activity. Palczewski et al. (23) expressed human BCO2 with the N-terminal mitochondrial targeting sequence truncated at different lengths in β-carotene-producing E. coli and observed β-apo-10'-carotenol production with all variants of the enzyme. However, they also did not observe activity in cell-free systems (14).

Our results show that, with the exception of β-cryptoxanthin, chicken BCO2 cleaves carotenoids specifically at the 9-10 or 9'-10' position to yield an apo-10'-carotene. The ability of chicken BCO2 to cleave β-cryptoxanthin at the 7'-8' double bond to produce 3-hydroxy-β-apo-8'-carotenol is possibly due
to the asymmetry of the molecule, causing a minor, alternative binding mode with BCO2.

As of the time of writing, there are no available crystal structures of BCO1 and BCO2. The only carotenoid oxygenase crystal structures available are those for an apocarotenoid 15,15'-oxygenase (27) and a 9-cis-epoxycarotenoid 11,12-oxygenase (28). Based on these crystal structures, the carotenoid oxygenases discriminate between substrates by the size of the hydrophobic substrate tunnel and the interactions of certain hydrophobic and aromatic residues (for a review, see Sui et al. (29)). Our experiments with BCO1 show that hydroxylation of the ring in β-cryptoxanthin results in a lower $K_m$ compared with β-carotene and total loss of activity when the two rings are hydroxylated as in zeaxanthin (9). This suggests that a hydroxyl group may form unfavorable interactions with the hydrophobic substrate tunnel of BCO1 or that it is simply an issue of size; i.e., the 3-hydroxy-β-ionone ring is too large to pass through the substrate tunnel. The ability of BCO2 to cleave both carotenoids with and without 3-hydroxyl groups on the ionone rings, as shown by this study and others (6, 7), suggests that the substrate tunnel of BCO2 is larger than that of BCO1 and/or that the 3-hydroxyl groups do not form unfavorable interactions with the hydrophobic substrate tunnel.

BCO2 clearly catalyzes the oxidative cleavage of 9-cis-β-carotene (Fig. 5) and produces mostly 9-cis-β-apo-10'-carotenal (as discussed under “Results,” the production of the all-trans product cannot be completely ruled out). This is not surprising because if the enzyme were to produce all-trans-β-apo-10'-carotenal, the kinked cis 9-10 end of 9-cis-β-carotene would have to enter the substrate tunnel. The size of the substrate tunnel may not allow for this, and thus BCO2 is regioselective (or even regiospecific) for the trans 9'-10' end of 9-cis-β-carotene.

The large difference in the product peak areas obtained from the hydrocarbon carotenoids (β-carotene and α-carotene) compared with the hydroxylated carotenoids (β-cryptoxanthin, lutein, and zeaxanthin) (Figs. 5 and 7) suggests that hydroxylated carotenoids are the preferred substrates of the enzyme. It is worth noting that, at the lowest concentration tested for the α-cryptoxanthin substrate-velocity plot (7.5 μM; Fig. 9), the sum of the specific activities for β-apo-10'-carotenal and 3-hydroxy-β-apo-10'-carotenal production is 12× greater than that at the highest concentration tested for β-carotene (5 μM; Fig. 3C). Furthermore, at saturating β-cryptoxanthin, the sum of the specific activities for β-apo-10'-carotenal and 3-hydroxy-β-apo-10'-carotenal production is 33× greater than the calculated $V_{max}$ for β-carotene. Kinetic data obtained using lysates from ferret BCO2-expressing Sf9 cells also suggest that lutein and zeaxanthin, which both contain 3- and 3'-hydroxyl groups, are better substrates than β-cryptoxanthin, which contains only one 3-hydroxyl group (7). Our semiquantitative results, at this point, also support this, although determination of kinetic parameters is necessary to make this conclusive. Similar to the issue with β-cryptoxanthin noted under “Results,”
asymmetrical full-length carotenoids that are cleaved at two sites presumably have two binding modes and thus two \( K_m \) values. This should be taken into consideration when comparing \( K_m \) values with symmetrical substrates such as \( \beta \)-carotene.

The differences in peak areas of the product peaks from the asymmetrical carotenoid substrates (\( \alpha \)-carotene, \( \beta \)-cryptoxanthin, lutein) (Figs. 7 and 10) suggest that the enzyme displays an apparent regioselectivity. As mentioned previously, we do not have the standards for quantification of cleavage products except for \( \beta \)-apo-10'-carotenal, and thus we cannot conclude with absolute certainty that the small differences in peak areas of the product peaks of \( \alpha \)-carotene (Fig. 7) and lutein (Fig. 10) are due to true regioselectivity or the differences in extinction coefficients of the compounds as well as other factors that contribute to instrument response. With \( \beta \)-cryptoxanthin, the difference between the \( \beta \)-apo-10'-carotenol and 3-hydroxy-\( \beta \)-apo-10'-carotenal product peaks (Fig. 7B) is larger. The difference in the extinction coefficients of these two compounds is not expected to be very large (18), and thus it is likely (and not surprising) that the enzyme really does display regioselectivity with \( \beta \)-cryptoxanthin.

Unlike murine BCO2 (6), chicken BCO2 cannot perform a second cleavage on \( \beta \)-apo-10'-carotenal to yield rosafuene dialdehyde. If galloxanthin (3-hydroxy-\( \beta \)-apo-10'-carotenol) is an important carotenoid in avian vision and is indeed generated via BCO2 cleavage of zeaxanthin or lutein (Fig. 6), then it makes functional sense that chicken BCO2 does not cleave apo-10'-carotenals. If the function of BCO2 in mice is to degrade excess carotenoids, then it also makes functional sense that it can cleave full-length carotenoids twice, producing smaller molecules that are easier to clear from the circulation.

Chicken BCO2 has no activity with all-trans-lycopene, which is in agreement with the report on ferret BCO2 (30). This is also consistent with the preference of BCO2 for substrates that contain a 3-hydroxy-\( \beta \)-ionone ring. Ferret BCO2, however, is able to cleave cis isomers of lycopene (30). Ferrets accumulate cis-...
lycopene isomers when supplemented with all-trans-lycopene (30). Thus, the preference of ferret BCO2 for the latter is consistent with its proposed role of preventing excess accumulation of carotenoids. Similarly, the majority of lycopene in human fluids and tissues are cis isomers (31, 32), and it will not be surprising if human BCO2 can also catalyze their oxidative cleavage. The acid, alcohol, and aldehyde forms of the 9'/H11032-10'/H11032-cleavage product of lycopene have shown biological activity in cell culture, inducing retinoic acid receptor β mRNA levels and phase II enzymes at the mRNA and protein levels (33).

Chicken BCO2 also does not have activity with the other β-apocarotenals in the series (β-apo-8'-carotenal, β-apo-12'-carotenal, and β-apo-14'-carotenal). Again, this is consistent with the importance of the 3-hydroxy-β-ionone ring for BCO2 cleavage.  β-carotene, with two β-ionone rings, is a relatively poor substrate, and it is not surprising that the β-apocarotenals, with only one β-ionone ring per molecule, are altogether inactive.

It is interesting that the substrates that are inactive with BCO2 (all-trans-lycopene and β-apocarotenals) are substrates for BCO1 (Fig. 11). With the β-apocarotenals, this
Substrate Specificity of Chicken BCO2

| Substrate         | Chemical structure |
|-------------------|--------------------|
| β-carotene        | ![β-carotene Structure](image) |
| α-carotene        | ![α-carotene Structure](image) |
| β-cryptoxanthin   | ![β-cryptoxanthin Structure](image) |
| Lycopene          | ![Lycopene Structure](image) |
| β-apo-8'-carotenal| ![β-apo-8'-carotenal Structure](image) |
| β-apo-10'-carotenal| ![β-apo-10'-carotenal Structure](image) |
| β-apo-12'-carotenal| ![β-apo-12'-carotenal Structure](image) |
| β-apo-14'-carotenal| ![β-apo-14'-carotenal Structure](image) |
| Zeaxanthin        | ![Zeaxanthin Structure](image) |
| Lutein            | ![Lutein Structure](image) |
| 9-cis β-carotene  | ![9-cis β-carotene Structure](image) |

**FIGURE 11. Structures of the substrates tested with purified recombinant chicken BCO2 and human BCO1 (9).** The red bars represent the cleavages observed with BCO2, and the blue bars, BCO1.

In summary, our results show that chicken BCO2 has broader substrate specificity than BCO1, consistent with its proposed function of preventing oxidative stress brought about by carotenoid accumulation in the mitochondria. It cleaves only full-length carotenoids with ionone rings, and the hydroxylated carotenoids are cleaved to a greater extent than their hydrocarbon counterparts under the conditions tested. Furthermore, the enzyme displays regioselectivity with asymmetrical substrates.

**Author Contributions**—E. H. H. conceived and coordinated the study and cowrote the paper with C. d. S. C. d. S., J. S., and Y. Y. prepared the enzyme and designed, conducted, and analyzed the experiments. S. N. and R. W. C. synthesized compounds. K. M. R., S. J. S., and C. d. S. designed and conducted the LC-MS experiments. All authors reviewed the results and approved the final version of the manuscript.

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Carlo dela Seña, Jian Sun, Sureshbabu Narayanasamy, Kenneth M. Riedl, Yan Yuan, Robert W. Curley, Jr., Steven J. Schwartz and Earl H. Harrison

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