Key Residues for Catalytic Function and Metal Coordination in a Carotenoid Cleavage Dioxygenase*

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Carotenoid cleavage dioxygenases (CCDs) are non-heme iron-containing enzymes found in all domains of life that generate biologically important apocarotenoids. Prior studies have revealed a critical role for a conserved 4-His motif in forming the CCD iron center. By contrast, the roles of other active site residues in catalytic function, including maintenance of the stringent regio- and stereo-selective cleavage activity, typically exhibited by these enzymes have not been thoroughly investigated. Here, we examined the functional and structural importance of active site residues in an apocarotenoid-cleaving oxygenase (ACO) from Synechocystis. Most active site substitutions variably lowered maximal catalytic activity without markedly affecting the $K_m$ value for the all-trans-8′-apocarotenol substrate. Native C15-C15′ cleavage activity was retained in all ACO variants examined suggesting that multiple active site residues contribute to the enzyme’s regioselectivity. Crystallographic analysis of a nearly inactive W149A-substituted ACO revealed marked disruption of the active site structure, including loss of iron coordination by His-238 apparently from an altered conformation of the conserved second sphere Glu-150 residue. Gln- and Asp-150-substituted versions of ACO further confirmed the structural/functional requirement for a Glu side chain at this position, which is homologous to Glu-148 in RPE65, a site in which substitution to Asp has been associated with loss of enzymatic function in Leber congenital amaurosis. The novel links shown here between ACO active site structure and catalytic activity could be broadly applicable to other CCD members and provide insights into the molecular pathogenesis of vision loss associated with an RPE65 point mutation.

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In metazoans, vitamin A (all-trans-retinol) and its metabolites, collectively referred to as retinoids, participate in many essential physiological processes, including embryonic development, growth, immune function, reproduction, and vision (1–4). Therefore, a sustainable supply of vitamin A is critical to fulfill the diverse biological functions associated with retinoids. However, the lack of a de novo vitamin A biosynthetic pathway in animals necessitates their dietary intake of provitamin A carotenoids or retinyl esters as vitamin A precursors (5). In mammals, β-carotene, the most abundant pro-vitamin A carotenoid in nature, is absorbed by intestinal mucosal cells and oxidatively metabolized into vitamin A-aldehyde (all-trans-retinal, RAL), which can be successively reduced to vitamin A and stored in the liver as fatty acid retinyl esters (e.g. retinyl palmitate) (6–8). Oxidative cleavage of the carotenoid polypeptide chain is catalyzed by a family of non-heme iron-dependent enzymes known as carotenoid cleavage dioxygenases (CCDs) that are found in all kingdoms of life (9–11). Besides their involvement in retinal formation, CCDs generate several other important apocarotenoids, including abscisic acid, strigolactones, pigments and volatiles in plants, and degrade stilbenoid compounds such as resveratrol and its derivatives as well as lignin catabolites in bacteria and fungi (11–16).

Although CCDs can often cleave multiple substrates, they generally display high regio- and stereo-selectivity with respect to the cleavage site within the carotenoid polypeptide chain as well as the polyene isomeric configuration (9, 11). For example, human BCO1, a retinal-forming CCD, catalyzes symmetrical cleavage of different all-trans-cyclic carotenoids at the central double bond position (C15-C15′ according to traditional carotenoid carbon numbering) of the polypeptide chain (17, 18), whereas BCO2, another human carotenoid-metabolizing CCD, is capable of cleaving not only cyclic but also acyclic carotenoids asymmetrically (C9-C10 position) (19–21). Not limited to mammals, these distinct enzymatic features (i.e. substrate promiscuity coupled with regio- and stereo-selectivity) are also commonly found in plant and bacteria CCDs (11, 22, 23). For example, a plant CCD called VP14 that plays a critical role in abscisic acid biosynthesis cleaves 9-cis-exoperoxycarotenoids selectively at the C11-C12 polypeptide double bond (24, 25). Interestingly, certain CCDs exhibit isomerase activity rather than or in addition to oxygenase activity. One notable example is RPE65,
which converts fatty acid all-trans-retinyl esters (predomi-
nantly palmitate) into 11-cis-retinol via a concurrent isomer-
ization and atypical hydrolysis reaction (26–28). This enzyme
plays an indispensable role in the regeneration pathway for
11-cis-retinal, the chromophore of retinal photoreceptor visual
pigments. RPE65 loss-of-function mutations cause severe reti-
nal dystrophies, such as retinitis pigmentosa and Leber congen-
ital amauros (LCA), leading to blindness (29–31).

The CCD structural fold consists of a seven-bladed β-propel-
ler capped on the top face by non-contiguous α-helices and
loops that together form the substrate binding tunnel. Residues
lining this tunnel are generally hydrophobic and thus provide
an ideal environment to accommodate apolar carotenoid sub-
strates (32–35). Despite this shared physicochemical property,
the tunnels observed in published CCD structures differ sub-
stantially in their shapes and amino acid compositions. The
geometric and steric restrictions imposed by the residues
within this region are believed to underlie the stereo and regi-
oselectivity of CCD enzymes (10). However, there are currently
no published structures of genuine CCD-substrate complexes
that validate this hypothesis, as the generally poor aqueous sol-
ubility of carotenoids in general poses a formidable challenge
to CCD structural studies (33, 34). Although a few prior studies
have examined the functional importance of active site residues
in CCD substrate interactions, metal binding, and catalytic activity
(34, 36–40), detailed structural and biochemical studies
concerning this issue are lacking.

The CCD iron catalytic center is positioned deep within the
substrate binding tunnel with the Fe(II) cofactor coordinated
by four strictly conserved His residues, leaving one or poten-
tially two open coordination sites, depending on the specific
CCD, accessible for ligand binding. In addition, three con-
served Glu residues indirectly contribute to iron coordination
through hydrogen bonding interactions with three of the direct
His ligands. This 4-His + 3-Glu dual-sphere metal-binding motif
distinguishes CCDs from other non-heme mononuclear
iron centers (41). Involvement of the Glu sphere is indispens-
able for CCD catalytic function as shown by previous studies
(28, 36, 42). Additionally, mutations in the RPE65 gene that
cause substitutions in second sphere Glu residues are associ-
ated with severe retinal dystrophy (43). Previous biochemical
studies imply that the negative-charged carboxylate groups of
these Glu residues are involved in iron charge neutralization
(36, 42), but their precise role in maintaining the structure of
the iron center and CCD catalytic activity remains elusive.

*Synechocystis* apocarotenoid oxygenase (ACO) is a prototyp-
ical CCD that is amenable to structure-function studies, includ-
ing crystallographic analysis of point mutants (32, 33, 44) in
contrast to mammalian CCDs for which crystallography of recombinantly generated protein is not yet feasible. Here, we
employed this enzyme as a model to probe the relationship
between CCD active site structure, catalytic activity, and regi-
oselectivity. Our results indicate that although many ACO
active site mutants have impaired catalytic activity, the regio-
selectivity of this enzyme is highly resistant to active site amino
acid substitutions. Structural analysis of a particularly detri-
mental point mutant (W149A) revealed unexpected disrup-
tions in regions of the active site distant from where the Trp
side chain normally resides. Two of these perturbed residues,
His-238 and Glu-150, are involved in iron coordination, the
latter residue being homologous to Glu-148 in RPE65, a posi-
tion at which Asp substitution has been associated with LCA
(42, 43). In both Asp-150 and Gln-150 ACO point mutants, the
non-native side chain fails to form a fully stable interaction with
His-238, which in turn disrupts the His-238–Fe(II) coordinate
bond. This destabilization of the iron center likely contributes
to the low catalytic activity of Asp/Gln-150 ACO mutants and
the retinal pathology associated with the E148D variant of
RPE65. Thus, in addition to providing key insights into the
active site determinants of CCD catalysis, these data provide
novel structural details regarding the role of second sphere Glu
residues in iron coordination by CCDs and help illuminate the
molecular pathology associated with an RPE65 point mutation.

### Results

**Identification of Potential Substrate-interacting Residues for Mutagenesis Studies**—ACO recognizes C25–35 β-apocarote-
noids/ols and specifically cleaves them at the C15-C15′ double bond to form RAL and a second linear apocarotenoid product
(Fig. 1A) (22). Among the different lengths of β-apocarote-
noids, ACO displays the highest activity toward all-trans-8′-
apocarotenol and its 3-hydroxy derivative. The former is com-
mercially available, making it convenient for activity studies
(Fig. 1A). To identify active site residues important for sub-
strate recognition and cleavage site selectivity, we carried out in
silico docking studies of all-trans-8′-apocarotenol in the ACO
active site (PDB accession code 4OU8). In multiple docking
runs, the top-binding pose, as assessed by the Autodock Vina

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**Figure 1.** ACO-catalyzed cleavage reaction and model of the ACO-sub-
strate complex. A, ACO accepts and specifically cleaves all-trans-8′-apocaro-
tenol at the C15-C15′ double bond (red wavy line) with incorporation of both
O2-originated oxygen atoms into the RAL and C10-aldehyde apocarotenoid
products. B, stick view of an in silico generated model of ACO (PDB code 4OU8)
complexed with all-trans-8′-apocarotenol. The most energetically favorable
binding pose is featured with residues located close to the docked apocaro-
tenoid molecule (in gray) shown as orange sticks. The iron cofactor, shown as
a brown sphere, is close to the scissile C15-C15′ double bond (labeled as C15).
scoring function (45), oriented the substrate with the β-ionone moiety interacting with residues at the entrance of the active site pocket and the C15-C15′ double bond in close proximity to the iron center consistent with the known biochemical properties of ACO (Fig. 1B) (22). This mode of binding resembles that of the apocarotenoid ligand modeled into the originally reported ACO crystal structure (PDB accession code 2BIW) (32). However, the apocarotenoid substrate in the previous structure was modeled in a di-cis configuration based on the bent electron density located in the active site cavity. This bent electron density feature was later shown to also be present in substrate-free crystals as well indicating that previously modeled substrate was spurious (33).

We used the in silico docked model to identify residues involved in substrate interactions. Three residues with bulky aromatic side chains (Trp-149, Phe-113, and Phe-236) near the entrance to the active site cavity appeared to be positioned appropriately to control the depth of substrate entry into the active site entry such that the 15–15′ double bond is positioned close to the iron center to facilitate efficient and regioselective cleavage, as suggested previously (Fig. 1B) (32). The scissile bond was surrounded by the phenyl side chains of Phe-69 and Phe-303 (Fig. 1B). Based on their proximity to the site of cleavage, we reasoned that they could reinforce cleavage selectivity as well as help stabilize reaction intermediates. In the innermost region of the cavity, three residues (Tyr-24, Leu-400, and Phe-371) appeared to help form a pocket that could accommodate the distal polar end of the substrate. As the only residue containing a polar hydroxyl moiety, Tyr-24 may form a water-mediated interaction with the substrate hydroxyl tail that is located 5.7 Å away in the docked model (Fig. 1B). Alternative conformations of the polar end of the substrate molecule could also enable a direct interaction between these two moieties.

We divided the above-mentioned residues into three groups based on their locations: the β-ionone ring binding region (Trp-149, Phe-113, and Phe-236), the region flanking the cleavage site (Phe-69 and Phe-303), and the distal end in the binding pocket (Tyr-24, Phe-371, and Leu-400) (Table 1).

### Table 1

<table>
<thead>
<tr>
<th>ACO mutations and their corresponding locations in the active site relative to the docked substrate ligand</th>
<th>Targeted residues</th>
<th>Mutants</th>
</tr>
</thead>
<tbody>
<tr>
<td>β-Ionone proximal region of the substrate</td>
<td>Phe-113, Trp-149, Phe-236</td>
<td>F113A, W149A, F236A</td>
</tr>
<tr>
<td>Flanking the substrate scissile double bond</td>
<td>Phe-69, Phe-303</td>
<td>F69Y, F303A, F69A/F303A</td>
</tr>
<tr>
<td>Distal end in the binding tunnel interior site</td>
<td>Tyr-24, Phe-371, Leu-400</td>
<td>Y24F, F371A, L400A</td>
</tr>
</tbody>
</table>

ACO Active Site Mutants Impair Catalytic Activity to Various Degrees without Altering Regioselectivity—Three residues located in the β-ionone binding region (Phe-113, Phe-236, and Trp-149) and two in the distal pocket (Phe-371 and Leu-400) were substituted with alanine to probe their contributions to substrate interactions. Tyr-24 also was mutated to a phenylalanine (Y24F) to examine its potential to hydrogen bond with the substrate’s hydroxyl group. Phe residues 69 and 303, which flank the scissile double bond, were replaced either independently with alanine (F69A and F303A) or together (F69A/F303A) to evaluate their involvement in regioselectivity and overall catalytic activity. The effect of introducing a polar functional group into the generally apolar environment that surrounds the catalytic center was examined with F69Y-substituted ACO. Mutants examined in this study are summarized in Table 1.

All ACO mutants were expressed and purified in the same manner as wild-type protein. All mutants featured detectable but variable levels of expression compared with wild-type ACO (Table 2). SDS-PAGE analysis of each purified ACO showed a similar purity, with the exception of F69Y, which was markedly diminished (Fig. 2A), consistent with its lower expression level compared with the other mutants (Table 2). Surprisingly, except for F69Y, oxidative cleavage activity, as assessed at a single substrate concentration by HPLC, was retained by all mutants (Fig. 2B). Among them, five ACO mutants (F69A, W149A, E150Q, F371A, and L400A) showed <5% activity, whereas the others displayed enzymatic activity ranging from 8 to 76% of the native protein (Fig. 2B).

To test possible changes in regioselectivity, we analyzed the reaction products by HPLC-MS. Whereas all-trans-retinal was the only β-apocarotenoid product of wild-type ACO detectable by UV-visible absorbance profiles, the atypical all-trans-12′-apocarotenal product of oxidative cleavage at the C11′-C12′ position was detected by HPLC-MS (Fig. 3, A and B). Each single mutant also produced RAL as the dominant product (Fig. 2B), indicating an unaltered selectivity for the cleavage site. Products derived from the double mutant F69A/F303A were also analyzed to assess whether the reduced steric hindrance in vicinity of the cleavage site in this mutant protein (Fig. 1A) could potentially render a more pronounced change in regioselectivity. Production of all-trans-12′-apocarotenal was slightly increased in this double mutant compared with wild-type protein, although the quantity produced was still below the limits of UV-visible detection (Fig. 3C). These results demonstrate that active site mutants differentially affected catalytic activity but that regioselectivity of ACO is preserved in the face of diverse active site perturbations.

Most ACO Active Site Mutants Have Reduced Maximal Enzymatic Activity with Minimally Altered Michaelis Constants—To further investigate the enzymology underlying the impaired activities, steady-state kinetic studies for each mutant were performed. Having established the cleavage selectivity of each ACO variant by HPLC, we employed a more rapid assay system for these steady-state kinetic studies in which substrate consumption is monitored spectrophotometrically in a plate reader. The steady-state kinetic parameters of native ACO determined by this method were in good agreement with those obtained previously by HPLC analysis (33), thus validating the assay methodology (Fig. 4). A majority of mutants showed only minimally perturbed \( K_m \) values (Y24F, F69Y, W149A, E150Q, F236A, F303A, F371A, and L400A), with four variants (F69A, F113A, E150D, and F69A/F303A) displaying ~2–3-fold increases as compared with native ACO (Table 2).
**Structure-Activity Relationships of Apocarotenoid Oxygenase**

**TABLE 2**

<table>
<thead>
<tr>
<th>ACO mutant</th>
<th>Expression level</th>
<th>$K_m^{a}$</th>
<th>$V_{max}$</th>
<th>$k_{cat}$</th>
<th>$k_{cat}/K_m$</th>
<th>RPE65 equivalent</th>
<th>Disease$^b$</th>
</tr>
</thead>
<tbody>
<tr>
<td>WT</td>
<td>+++++</td>
<td>24 ± 5</td>
<td>1.3 ± 0.1</td>
<td>17 ± 1</td>
<td>0.72 ± 0.16</td>
<td>Phe-16</td>
<td>Polymorphism</td>
</tr>
<tr>
<td>Y24F</td>
<td>++</td>
<td>27 ± 6</td>
<td>1.1 ± 0.1</td>
<td>15 ± 1</td>
<td>0.56 ± 0.13</td>
<td>Phe-61</td>
<td></td>
</tr>
<tr>
<td>F69A</td>
<td>+</td>
<td>67 ± 10</td>
<td>0.04 ± 0.01</td>
<td>0.67 ± 0.3</td>
<td>0.008 ± 0.002</td>
<td>Phe-61</td>
<td></td>
</tr>
<tr>
<td>F69Y</td>
<td>+</td>
<td>22 ± 4</td>
<td>0.012 ± 0.002</td>
<td>NA</td>
<td>NA</td>
<td>Phe-103</td>
<td></td>
</tr>
<tr>
<td>F113A</td>
<td>+</td>
<td>48 ± 20</td>
<td>0.23 ± 0.04</td>
<td>3.1 ± 0.5</td>
<td>0.066 ± 0.029</td>
<td>Phe-103</td>
<td></td>
</tr>
<tr>
<td>W149A</td>
<td>++</td>
<td>30 ± 5</td>
<td>0.018 ± 0.002</td>
<td>0.24 ± 0.02</td>
<td>0.008 ± 0.002</td>
<td>Thr-147</td>
<td></td>
</tr>
<tr>
<td>E150D</td>
<td>+++</td>
<td>80 ± 8</td>
<td>0.38 ± 0.02</td>
<td>0.5 ± 0.2</td>
<td>0.064 ± 0.007</td>
<td>Glu-148</td>
<td>LCA</td>
</tr>
<tr>
<td>E150Q</td>
<td>++</td>
<td>22 ± 4</td>
<td>0.014 ± 0.002</td>
<td>0.19 ± 0.03</td>
<td>0.009 ± 0.002</td>
<td>LCA</td>
<td></td>
</tr>
<tr>
<td>F236A</td>
<td>+++</td>
<td>21 ± 4</td>
<td>0.44 ± 0.03</td>
<td>0.80 ± 0.14</td>
<td>0.28 ± 0.04</td>
<td>Tyr-239</td>
<td>LCA</td>
</tr>
<tr>
<td>F303A</td>
<td>++</td>
<td>24 ± 4</td>
<td>0.13 ± 0.01</td>
<td>0.8 ± 0.5</td>
<td>0.076 ± 0.015</td>
<td>Phe-312</td>
<td></td>
</tr>
<tr>
<td>F303A</td>
<td>++</td>
<td>22 ± 4</td>
<td>0.014 ± 0.002</td>
<td>0.19 ± 0.27</td>
<td>0.009 ± 0.01</td>
<td>Phe-418</td>
<td></td>
</tr>
<tr>
<td>L400A</td>
<td>+</td>
<td>26 ± 2</td>
<td>0.046 ± 0.002</td>
<td>0.62 ± 0.03</td>
<td>0.024 ± 0.002</td>
<td>Pro-444</td>
<td></td>
</tr>
<tr>
<td>F69A/F303A</td>
<td>+</td>
<td>50 ± 9</td>
<td>0.12 ± 0.01</td>
<td>1.7 ± 0.1</td>
<td>0.033 ± 0.007</td>
<td>Phe-61/Phe-312</td>
<td></td>
</tr>
</tbody>
</table>

$^a$ Kinetic data were obtained with purified recombinant proteins.

$^b$ Disease was caused by mutation of the homologous position in RPE65.

$^c$ Protein purity was insufficient for accurate $k_{cat}$ calculation.

Note that most ACO mutants were purified to a degree comparable with that from duplicate results of a single experiment.

Crystal Structure of W149A ACO Reveals Major Disruptions in the Substrate-binding Cleft and Metal Coordination—The bulky hydrophobic side chain of Trp-149 and its proximity to the β-ionone ring of the substrate suggest an important function for this residue in substrate interactions (Fig. 1B). The near absence of activity in the W149A ACO variant (Fig. 2B) prompted us to investigate its underlying structural basis. To this end, W149A ACO was crystallized under the same conditions used for native ACO. The best crystals diffracted to ~2.8 Å resolution and were isomorphous to the previously reported orthorhombic ACO crystals (9). In this ACO mutant structure as well as those discussed below, we observed electron density in the solvent region consistent with the presence of a fifth ACO molecule in the asymmetric unit. The extra protomer was modeled in the final structure but was not used to draw structural conclusions due to its weak electron density relative to the other modeled chains. Superimposition of W149A and native ACO revealed an average root mean square deviation of ~0.425 Å between Co atoms. Absence of electron density for the Trp-149 side chain confirmed the Ala substitution (Fig. 5B).

Despite the overall structural similarity to native ACO, inspection of the W149A ACO catalytic center revealed an unexpected disruption in the iron coordination center (Fig. 5, A compared with B). Glu-150, a member of the conserved 3-Glu second sphere iron-binding motif, adjacent in sequence to the mutated site, adopted an alternative conformation that abolished its normal hydrogen bonding interaction with His-238. This change was accompanied by enhanced mobility of the His-238 side chain as evidenced by its weakened electron density as well as a loss of the coordinate bonding interaction with the iron cofactor (Fig. 5B), resulting in a 3-coordinate iron center. The perturbation in iron coordination resulted in a substantially elevated iron $B$-factor relative to the average $B$-factor for the protein in comparison with that of the wild-type protein at a comparable resolution (PDB accession code 4OU8 (33), which reflects increased iron mobility and/or reduced occupancy within the binding site. These structural perturbations could be directly attributed to alterations in the local structure surrounding position 149. Trp-149 and Glu-150 are located in

![Mass (Da)](image)

![Expression, purification, and activity of native and mutant ACOs](image)

and Fig. 4). By contrast, $k_{cat}$ values for most mutants were found to be substantially impaired. Y24F was the only variant with kinetic constants similar to the native enzyme, suggesting a less critical role for this residue in catalysis compared with the others examined (Table 2). Three variants (F113A, E150D, and F236A) had $k_{cat}$ values reduced by 70–80% with the remainder displaying maximal activities less than 10% of native protein (Fig. 4 and Table 2). The drastically decreased $k_{cat}$ values for many of the ACO variants examined imply important roles in maintenance of catalytic efficiency for residues in all regions of the active site pocket. The less pronounced effects of most substitutions on the stability of the enzyme-substrate complex at steady state (i.e. the $K_m$ value) together with the preserved regiospecificity of the variants indicate that residues lining the substrate binding tunnel work in concert to create a rigid platform for substrate binding and processing.

![Crystal Structure](image)

The perturbation in iron coordination resulted in a substantially elevated iron $B$-factor relative to the average $B$-factor for the protein in comparison with that of the wild-type protein at a comparable resolution (PDB accession code 4OU8 (33), which reflects increased iron mobility and/or reduced occupancy within the binding site. These structural perturbations could be directly attributed to alterations in the local structure surrounding position 149. Trp-149 and Glu-150 are located in
Structure-Activity Relationships of Apocarotenoid Oxygenase

FIGURE 3. Formation of all-trans-12′-apocarotenal by native and F69A/F303A ACO. A, detection of all-trans-12′-apocarotenal product formation by mass spectrometry. Extracted ion chromatogram of the reaction products generated by F69A/F303A ACO show the presence of an m/z 351.2 ion intensity peak at ~4.8 min. B, MS analysis of the product signal in A showing a dominant ion at m/z 351.19, consistent with the protonated form of all-trans-12′-apocarotenal generated by cleavage at the C11′-C12′ double bond of all-trans-8′-apocarotenol. Tandem MS analysis of the m/z 351.19 peak revealed a major fragmentation product at m/z 333.3 representing loss of water from the parent ion. C, semi-quantification of the atypical product formed by mutant and wild-type ACO according to the peak intensities in A. Formation of the atypical product was a linear function of the amount of enzyme in the reaction mixture up to a mass of 20 μg. Error bars represent standard deviations from experiments performed in duplicate.

the i and i + 1 positions of a type 1 β-turn connecting the middle strands of blade 1 of the β-propeller fold. Trp is infrequently found at the start of such β-turns (46), and the bulky indole moiety could help enforce the proper geometry of Glu-150. Notably, the ψ angle of ~9.3° for Glu-150 differs significantly from the ideal value of ~30° for an i + 1 residue in a type 1 β-turn, possibly as a consequence of steric effects resulting from the bulky Trp side chain (47). The Ala substitution at position 149 converts the β-turn into a type IV conformation with loss of a hydrogen bonding interaction between the i and i + 3 residues. These changes are accompanied by an ~2 Å shift in the Glu-150 Ca atom. Together, these structural alterations change the environment of the Glu-150 side chain such that its interaction with His-238 is no longer favored. In addition to alterations in the iron center, electron density for Phe-236, which is adjacent to His-238, was also substantially weakened indicating an enhanced mobility likely resulting from the His-238 structural perturbation (Fig. 5B). Interestingly, despite its extremely low activity, W149A displayed a $K_m$ value similar to that of native ACO, indicating a preserved substrate binding capacity. Because F236A exhibited an activity and enzymatic parameters comparable with native ACO, the activity loss observed in W149A ACO could be linked to disruption of the iron center and, to a lesser extent, a deficiency in substrate binding.

Glu-150 in the Second Sphere Is Critical for Metal Binding, Maintenance of Active Site Structure, and Catalytic Activity—Our results strongly suggested a critical role for Glu-150 in iron coordination and ACO catalytic function. To directly assess its structural and functional role in iron coordination, Glu-150 was substituted with Asp, which has a one-methylene shorter side chain and thus is likely incapable, within the confines of the β-turn structure, of hydrogen bonding with His-238. Because an equivalent substitution in RPE65 (E148D) was found in patients with LCA, characterization of this mutant would also improve our understanding of RPE65-associated retinal pathology. Whereas the expression and purification of E150D resembled that of native ACO, its maximal catalytic activity was reduced by ~80% with a 2–3-fold increased $K_m$ value (Fig. 2, A and B, and Table 2). To examine the structural consequences of this substitution, we determined the crystal structure of the mutant protein at a resolution of ~2.8 Å. Like the W149A mutant structure discussed above, the E150D crystals contained a fifth monomer in the asymmetric units albeit with weaker electron density support compared with the other chains. Electron density for Asp-150 was clearly evident but indicated that the side chain was flipped to a vacant site opposite the iron center (Fig. 6A). The hydrogen bonding interaction between Glu-150 and His-238 was eliminated, resulting in disruption of the coordination between His-238 and iron. Interestingly, His-238 is well resolved, but its side chain points away from the iron (Fig. 6A), clearly demonstrating an iron coordination defect in E150D ACO. Consequently, the altered three-coordinate metal center featured a reduced iron occupancy and/or high mobility as evidenced by an elevated iron $B$-factor. As in W149A ACO, Phe-236 exhibited a high mobility manifested by poorly resolved side chain electron density (Fig. 6A).

Previous mutagenesis studies of RPE65 suggest that the negative charge of carboxylate groups from the 3-Glu sphere contribute to iron charge neutralization (36, 42). It has also been suggested that the second sphere Glu residues help fix the first sphere His side chains in conformations capable of stably coordinating iron (48). To test these hypotheses, we replaced Glu-150 with Gln, which contains an uncharged side chain with a preserved capacity to hydrogen bond with His-238. This substitution yielded a nearly inactive enzyme (Table 2 and Fig. 4D). The crystal structure of E150Q ACO solved at 2.75 Å resolution revealed changes in the catalytic center largely resembling
those seen in the W149A and E150D ACO mutants, including loss of coordinate bonding between His-238 and the iron cofactor, an elevated iron $B$-factor, and increased mobility of Phe-236. However, unlike E150D ACO, the Gln residue maintained a hydrogen bonding interaction with His-238 (Fig. 6B). Indeed, the electron density for the His-238 imidazole ring is more discernable in E148Q ACO than that in the W149A and E150D mutants. However, coordination of His-238 to the iron was clearly disrupted, as demonstrated by the average $\sim 3.4$ Å distance separating the H238-N$^\bullet$ and iron atoms (Fig. 6B).

**Discussion**

The biological functions of CCDs have been described in numerous studies. However, uncertainties remain regarding...
how CCDs interact at the molecular level with carotenoids to determine their substrate specificity as well as their regio- and stereo-selectivity. With those questions in mind, we employed a cyanobacterial ACO as a model for the CCD family to systematically examine the functional and structural importance of its active site residues.

Based on our in silico ACO substrate model, the steric influence exerted by three hydrophobic residues (Trp-149, Phe-113, and Phe-236) could plausibly function as a “bottleneck” to limit passage of the substrate β-ionone moiety into the enzyme’s active site. This model agrees with a previous representation of the ACO-substrate complex proposed by Kloer and Schulz (9). Models in which the substrate orients with the β-ionone ring residing in the interior region of the tunnel are inconsistent with the lack of β-carotene cleavage by ACO (22). Rather the bottleneck hypothesis is consistent with experimental data showing that ACO can accept apocarotenoid substrates with polyene backbones of various lengths but only cleaves them at the C15-C15 position (Fig. 1) (22). The unperturbed regioselectivity in bottleneck residue point mutants shown here suggests a degree of functional redundancy exerted by these and potentially other active site residues with single mutations insufficient to alter the preferred site of cleavage.

Among the residues examined in this study, the position homologous to Phe-236 in ACO has also been studied in mouse BCO1 (Tyr-235) and maize CCD1 (Met-345) (34, 49). Interestingly, Tyr-235 in BCO1 was hypothesized to stabilize a proposed cationic reaction intermediate (49) that may be generated during CCD-catalyzed oxidative reactions (50). This was supported by the fact that substitutions of Tyr-235 with aromatic residues (Y235F/Y235W) minimally perturbed BCO1 activity, whereas non-aromatic amino acid replacements such as Y235L led to ~50% activity reduction (49). Indeed, a similar carbocation stabilization mechanism has been proposed for RPE65 (37) and some isoprenoid metabolizing enzymes (51, 52). In contrast to BCO1, the F236A substitution in ACO and the homologous mutant M276A in CCD1 failed to exhibit significant activity loss (34), which argues against a role for Phe-236 in carbocation stabilization, at least in ACO. CCD sequence alignments and structural superpositioning suggest that a consensus Phe residue may fulfill this function. This conserved Phe residue (Phe-69 in ACO, Phe-171 in VP14, and Phe-61 in RPE65) is located across from the iron center near the predicted position of the scissile double bond (9, 34, 35) where it could play a key role in stabilizing reaction intermediates.

Changes in substrate regio- and stereo-selectivity in response to active site alterations have been demonstrated in several dioxygenases, e.g. naphthalene and toluene dioxygenases (53–55). Inspired by these studies, we first analyzed the reaction products of native ACO and identified an atypical apocarotenoid product formed by cleavage at the C11′-C12′ site.
Structure-Activity Relationships of Apocarotenoid Oxygenase

However, the trace amount of this atypical product, detectable only by mass spectrometry, indicated that ACO-catalyzed polyene cleavage is tightly regulated with a high degree of precision (Fig. 3). We hypothesized that mutation of a substrate-interacting residue could enhance formation of the atypical product. Surprisingly, all tested ACO mutants faithfully generated RAL as the dominant product (Fig. 2B), with only a minor increase in formation of the atypical product observed for the F69A/F303A dual mutant (Fig. 3). These data suggest that residues in the substrate-binding cleft act together in substrate binding and processing, such that regioselective cleavage is maintained in the presence of subtle active site changes. However, formation of the atypical product could be under-represented in our analyses, as it could be further processed by the enzyme with RAL as the final product.

Conserved second sphere Glu and Asp residues are frequently found in non-heme iron enzymes. In 15-lipoxygenase, for instance, Glu-357 in the outer iron sphere is critical for enzyme activity as well as reaction specificity (56, 57). However, among this group of enzymes, CCDs appear to employ the most extensive use of a negatively charged second ligand sphere for maintenance of the iron center structure and function. The strict conservation of Glu, or in rare cases Asp, at the second sphere positions in this ancient family indicates a profound selective pressure maintaining the anionicity that envelopes the 4-His-Fe(II) center (41, 58). The neutral 4-His inner sphere of CCDs contrasts with metal-binding ligand sets of most other non-heme iron proteins, which usually contain at least one negatively charged residue (59). The non-heme iron of photosystem II constitutes the only other known example of a 4-His-Fe(II) center lacking protein-associated negatively charged iron ligands, which instead possesses a bound (bi)carbonate ligand (60). Therefore, the neutral CCD 4-His combination alone could be insufficient to cage the iron thus requiring outer sphere anionic ligand(s) to achieve a stable coordination complex. The ability of Dke1 (61) and cysteine dioxygenase (62) to complex Fe(II) with a 3-His triad motif somewhat argues against this hypothesis, although these enzymes also contain iron-bound solvent molecules that may contribute to charge stabilization and may adopt conformations with anionic side chains residing in first coordination sphere (63). By contrast, most CCD iron centers are likely capable of binding only a single solvent molecule and have no additional residues with coordination potential in their vicinity (10, 44). The presence of anionic Glu side chains in close proximity to the His ligands is expected to elevate the pKₐ of the inner sphere imidazole rings (64), which in turn would stabilize the His-N₈–iron coordinate bond and tune the reactivity of Fe(II) toward dioxygen. The disrupted interaction between Glu-150 and His-238 in W149A ACO coupled with the severe loss of activity in this mutant emphasize the importance of the 3-Glu sphere.

To further probe the role of Glu-150 in ACO function, two point mutants, E150D and E150Q, were structurally and kinetically characterized. In E150Q ACO, the loss of iron coordination by His-238 together with an ~3.3-fold reduced kₐ value unambiguously demonstrated a critical role of Glu in iron coordination and catalytic function. The structure of E150Q ACO revealed that this non-native residue, like Glu, forms a hydrogen bonding interaction with His-238, similar to that of the native Glu side chain. However, this interaction did not promote formation of a stable coordinate bond between His-238 and the iron cofactor. Interestingly, E150Q ACO was essentially inactive despite being expressed at a level comparable with the wild-type ACO suggesting that the Gln side chain, per se, exerts an additional negative effect on the catalytic properties of this enzyme. These results are consistent with previous studies showing that Gln substitution for one of the conserved outer sphere Glu residues in RPE65 was substantially more detrimental for catalytic activity than an Asp substitution (36, 42). It is conceivable that the Asp–150 ACO variant could partially stabilize His-238 in a catalytically competent orientation, particularly in the presence of substrate, to enable the observed residual activity. Indeed, in the crystal structure of E150D, weak electron density was consistently observed between the iron cofactor and His-238, which suggests that the ensemble of His conformations may include the native rotamer and that this subpopulation may be responsible for the residual activity. Cumulatively, our results reinforce the notion that the negativity of the 3-Glu motif is indispensable for CCD catalysis.

In addition to a role in iron coordination, a recent crystal structure of RPE65 complexed with emixustat, a retinoid-mimetic inhibitory amine, disclosed a role for Glu-148 in retinyl cation stabilization (65). In this high resolution crystal structure, the Glu-148-O⁺ atom is located 2.9 Å from the amine group of the compound, which suggests high tendency to form an electrostatic interaction with the C15 retinyl carbocation during catalysis (65). In the calculated ACO substrate model, the distance between the C15 site in the substrate and oxygen atom of Glu-150 is ~6 Å. Electrostatic contacts could be formed between Glu-150 and cationic apocarotenoid intermediates, but further structural and biochemical studies are needed to determine whether such a role is fulfilled by this residue in ACO.

In summary, our study utilized a systematic mutagenesis approach targeting active site residues that are potentially involved in carotenoid interactions. Detailed biochemical and kinetic data obtained with purified enzymes allowed us to test the functions of these residues in substrate interactions. In addition, our crystallographic and kinetic studies on selected ACO mutants provided insights needed to understand the unusual 4-His 3-Glu iron coordination system in CCDs, particularly the 3-Glu outer sphere. As a consequence of our studies with ACO, we also provide a plausible molecular mechanism underlying the functional impairment of an RPE65 mutant associated with human retinal dystrophy.

Materials and Methods

In Silico Ligand Docking—The substrate coordinate and stereochemical restraint files for all-trans-8'-apocarotenol were generated with the PRODRG server (66). Adjustments to the polypeptide configuration and regularization of the substrate bond lengths and angles were performed using COOT (67). The substrate coordinate file was then processed with Autodock Tools to generate a pdbqt file with polar hydrogens added to the ligand. The polypeptide single bonds were allowed to freely rotate. Docking was accomplished with a 2.8 Å resolution ACO crystal...
Structure (PDB code 4OU8) (33) in which water molecules in the model were removed. Autodock Tools were used to convert the model to a pdbqt format with polar hydrogens added in the protein model. Docking experiments of ACO with substrate were then carried out with Autodock Vina 1.1.2 (45). Multiple hits were identified during the docking trials that were carried out using a search area that included the entire ACO active site. The top binding pose, in which the C15-C15' double bond was placed in close proximity to the iron center with the β-ionone ring positioned near the membrane binding surface of the protein was used to guide the mutagenesis study.

Molecular Biology, Protein Expression, and Purification—All Synechocystis ACO point mutants were generated from a previously described pET3a-ACO expression plasmid (33) with a QuickChange site-directed mutagenesis kit (Stratagene, Santa Clara, CA) and confirmed by DNA sequencing. ACO was expressed as described previously with minor changes (33, 44). The LB culture supplemented with ampicillin (100 μg/ml) and ferrous iron (ammonium ferrous iron sulfate or Mohr's salt, 50 μg/ml) was grown at 37 °C, with 230 rpm shaking in an A<sub>600</sub> of ~0.6, and induced by adding isopropyl 1-thio-β-D-galactopyranoside to a final concentration of 100 μM. At the same time, additional ampicillin (100 μg/ml) and iron salt (50 μg/ml) were added into the culture. After an overnight incubation at 28 °C, cells were harvested by centrifugation and suspended in 20 mM HEPES-NaOH, pH 7.0. Cells were either flash-frozen and stored at −80 °C or used immediately. ACO purification was performed as described previously (33). All ACO mutants were expressed and purified identically to the wild-type protein. Concentrations of purified ACO samples were determined using an A<sub>280</sub>nm extinction coefficient of 75,249 M<sup>−1</sup>·cm<sup>−1</sup> as determined by amino acid analysis of purified wild-type ACO (Protein Chemistry Laboratory, Texas A&M University).

Enzymatic Assays, High Performance Liquid Chromatography (HPLC), and Mass Spectrometry (MS) Analyses—Activity studies of ACO and mutants were performed by previously established methods (33). Briefly, 2 μg of purified ACO was added to 200 μl of reaction buffer consisting of 20 mM HEPES-NaOH, pH 7.0, 0.05% (w/v) Triton X-100, and 1 mM tris(2-carboxyethyl)phosphine, pH 7.0. The all-trans-8'-apocarotenol substrate in DMSO was added to initiate the reaction. The reaction proceeded for 20 min at 28 °C with 500 rpm shaking in the dark and then was quenched with 200 μl of methanol. Products were extracted with 500 μl of hexane, and the analysis was performed by HPLC on a ZORBAX SIL (5 μm, 4.6 × 250 mm) normal phase column (Agilent, Santa Clara, CA). One hundred μl of extract were injected into the column, and elution with hexane/ethyl acetate (4:1, v/v) was carried out at 1.4 ml/min. RAL was quantified by plotting peak areas of known quantities of an authentic standard (TRC, Toronto, Canada). Enzymatic studies of all ACO mutants were conducted identically to those carried out with wild-type protein. For mass spectrometry analysis of products formed by wild-type and mutant forms of ACO, hexane extracts (100 μl) from the enzyme reaction mixtures were injected into the HPLC system as described above, and the eluates were directed into the atmospheric pressure chemical ionization probe source of an LXQ linear ion trap mass spectrometer (Thermo Scientific, Waltham, WA). Mass spectra were analyzed with the Xcalibur 2.0.7 software package.

Determination of Kinetic Parameters—For analysis of steady-state kinetics, ACO-catalyzed reactions were performed in a 96-well plate (Thermo Fisher, Waltham, MA) with different all-trans-8'-apocarotenol concentrations ranging from 0 to 150 μM. Inhibition of ACO activity was observed at substrate concentrations above 150 μM. One hundred μl of enzyme solution containing 4 μg of purified protein in reaction buffer and substrate at various concentrations in the same buffer were individually added to the plate wells. The plate was incubated at 28 °C for 10 min before mixing the enzyme solution with substrate. Reaction progression was monitored with a Flexstation3 microplate reader (Molecular Devices, Sunnyvale, CA) by measuring the change in substrate absorbance at 424 nm over time. Reactions at each substrate concentration were performed in triplicate. Care was taken to include only the linear portion of the progress curves for initial velocity measurements. A standard curve made with known substrate concentrations was used to quantify substrate depletion. Analysis of the kinetic data was performed with SigmaPlot (Systat Software, Inc., San Jose, CA).

Protein Crystallization, Structural Determination, and Analysis—Crystallization of ACO mutant enzymes was performed as described previously for the wild-type protein (33, 44). Briefly, purified protein samples were loaded onto a 25-ml Superdex 200 gel filtration column equilibrated with 20 mM HEPES-NaOH, pH 7.0, containing 0.02% (w/v) Triton X-100. A single symmetrical peak that eluted at ~13 ml was collected and concentrated to 8–10 mg/ml. For crystallization, 1.5–2 μl of purified enzyme at 10 mg/ml in gel filtration buffer was mixed with reservoir solution containing 0.1 M BisTris propane–HCl, pH 6.0, 21–23% (w/v) sodium polyacrylate 2100, and 0.2 M NaCl in a 1:1 ratio. Numerous trials yielded crystals with suboptimal morphology and poor diffraction quality. To improve crystal quality, a micro-seeding method was applied to obtain crystals of each mutant ACO. Specifically, after mixing protein samples with reservoir solution, a small quantity of crushed wild-type ACO microcrystals was applied to each drop. Crystallization was carried out by the hanging-drop vapor-diffusion method at 8 °C. Rod-shaped crystals typically appeared within 2–3 weeks. Mature crystals were directly harvested and flash-cooled in liquid nitrogen before x-ray exposure. Diffraction data were collected at the NE-CAT 24-ID-E beamline of the Advanced Photon Source and indexed, integrated, and scaled with the XDS package (68). Mutant ACO crystals were isomorphous to previously reported orthorhombic wild-type ACO crystals, and their structures were determined by rigid body refinement in REFMAC5 with PDB code 4OU9 used as the starting model (69). Manual adjustments to the structure were made with COOT (67), and restrained refinement was carried out in REFMAC5 (69). Structures were validated with MOLPROBITY (70) and the PDB structure validation server (71). In crystals of each mutant ACO, a fifth, less well defined ACO molecule was identified within the asymmetric unit. The x-ray data and refinement statistics are summarized in Table 3. All structural figures were prepared with PyMOL (Schrödinger, New York).
Structure-Activity Relationships of Apocarotenoid Oxygenase

TABLE 3
X-ray crystallographic data collection and refinement statistics for ACO mutants

<table>
<thead>
<tr>
<th>Data collection*</th>
<th>W149A ACO</th>
<th>E150D ACO</th>
<th>E150Q ACO</th>
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<tbody>
<tr>
<td>Crystal name</td>
<td>NECAT 24-ID-E</td>
<td>0.97910</td>
<td>0.97919</td>
</tr>
<tr>
<td>Beamline</td>
<td>0.97921</td>
<td>97.921</td>
<td>97.921</td>
</tr>
<tr>
<td>Wavelength (Å)</td>
<td>0.97921</td>
<td>97.921</td>
<td>97.921</td>
</tr>
<tr>
<td>Space group</td>
<td>P2₁,2₁,2₁</td>
<td>a = 118.42 b = 124.95 c = 203.97</td>
<td></td>
</tr>
<tr>
<td>a = 118.14 b = 125.26 c = 203.57</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Unit cell parameters (Å)</td>
<td>97.921</td>
<td>97.921</td>
<td>97.921</td>
</tr>
<tr>
<td>Resolution (Å)</td>
<td>48.62–2.80 (2.87–2.80)*</td>
<td>48.58–2.81 (2.88–2.81)*</td>
<td>47.42–2.75 (2.82–2.75)*</td>
</tr>
<tr>
<td>Unique reflections</td>
<td>75,013 (5,505)</td>
<td>74,574 (5,384)</td>
<td>75,683 (3,786)</td>
</tr>
<tr>
<td>Completeness (%)</td>
<td>100.0 (100.0)</td>
<td>99.9 (98.6)</td>
<td>95.9 (65.1)</td>
</tr>
<tr>
<td>Multiplicity</td>
<td>8.2 (8.1)</td>
<td>8.1 (7.9)</td>
<td>3.8 (3.1)</td>
</tr>
<tr>
<td>(I/σI)</td>
<td>11.0 (1.1)</td>
<td>7.9 (1.1)</td>
<td>7.6 (1.4)</td>
</tr>
<tr>
<td>Rmerge (%)</td>
<td>20.3 (217.9)</td>
<td>23.4 (172.5)</td>
<td>20.0 (124.4)</td>
</tr>
<tr>
<td>CC1/2 (%)</td>
<td>99.5 (30.5)</td>
<td>99.1 (49.6)</td>
<td>98.5 (33.7)</td>
</tr>
<tr>
<td>Wilson B-factor (Å²)</td>
<td>61</td>
<td>57</td>
<td>57</td>
</tr>
</tbody>
</table>

Refinement

| Resolution (Å)   | 48.62–2.80 | 48.58–2.81 | 47.42–2.75 |
| No. of observations | 71,417 | 70,990 | 72,122 |
| No atoms          | 18,975 | 18,839 | 18,840 |
| Protein           | 18,841 | 18,835 | 18,840 |
| Water             | 154 | 69 | 107 |
| Metal/Ion         | 5 Fe, 2 Cl | 5 Fe | 5 Fe |
| B-factors (Å²)    | 75 [61] | 73 [59] | 70 [56] |
| Protein           | 76 [62] | 73 [59] | 71 [56] |
| Water             | 42 [42] | 39 [38] | 35 [35] |
| Metal/Ion         | 107 [93] (Fe), 59 [59] (Cl) | 99 [87] (Fe) | 89 [72] (Fe) |
| Root mean square deviations | 0.01 | 0.01 | 0.01 |
| Bond angles (%)   | 1.38 | 1.33 | 1.27 |
| Ramachandran plot* | 97/0 | 97/0 | 96/0 |
| PDB accession code | 5KJA | 5KJB | 5KJD |

* Values in parentheses are those for the highest resolution shell of data.
* Data are as calculated in XD (68).
* Values in brackets are (B-factors) for the four well resolved chains (A–D).
* Data are as calculated in MOLPROBITY (70).

Author Contributions—X. S., K. P., and P. D. K. conceived and designed the study. X. S., J. Z., M. G., K. P., and P. D. K. performed experiments and analyzed the data. X. S. and P. D. K. wrote the manuscript. All authors reviewed the results and approved the final version of the manuscript.

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References

**Structure-Activity Relationships of Apocarotenoid Oxygenase**


Structure-Activity Relationships of Apocarotenoid Oxygenase
