Aromatic Residues in the Substrate Cleft of RPE65 Protein Govern Retinol Isomerization and Modulate Its Progression*5
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Background: RPE65 retinol isomerohydrolase is essential for vision, but its catalytic mechanism is poorly understood.

Results: Mutating aromatic residues in the substrate cleft abolishes or modifies RPE65 activity to make 13-cis- instead of 11-cis-retinol.

Conclusion: The RPE65 substrate-binding cleft is molded by aromatic residues to promote specific isomerization of retinol.

Significance: This further defines the vitamin A isomerization step central to the vertebrate visual cycle.

Previously, we showed that mutating RPE65 residue Phe-103 preferentially produces 13-cis-retinol instead of 11-cis-retinol, supporting a carbocation/radical cation mechanism of retinol isomerization. We asked whether this modulation of specificity can occur with residues other than Phe-103 and what role it plays in substrate binding and isomerization. We modeled the substrate-binding cleft of RPE65 to identify residues lining its surface. Many are phenylalanines and tyrosines, including three Phe residues (Phe-61, Phe-312, and Phe-526) forming an arch-like arrangement astride the cleft and Tyr-338. Also, Phe-418 sits at the neck of the cleft, lending a bend to the volume enclosed by the cleft. All mutations of Phe-61, Phe-312, and Phe-418 result in severely impaired or inactive enzyme. However, mutation of Phe-526 and Tyr-338, like Phe-103, decreases 11-cis-retinol formation, whereas increasing the 13-cis isomer. Significantly, 2 of these 3 residues, Phe-103 and Tyr-338, are located on putatively mobile interstrand loops.

The process of vision is initiated by photoisomerization of 11-cis-retinylidene bound to photoreceptor visual pigments, such as rhodopsin and cone opsins, to the all-trans conformation. Rhodopsin is thereby activated to metarhodopsin, initiating the phototransduction cascade. All-trans-retinal is ultimately released following the decay of metarhodopsin. To keep the visual pigments in a state capable of responding to light, they must be regenerated with 11-cis-retinal. This is achieved in the vertebrate retina by a process termed the visual cycle (1, 2). The visual cycle is localized in the retinal pigment epithelium (RPE) and retina and involves an assemblage of enzymes and isomer-specific binding proteins, some specific to the retina/RPE and some not. In the retinal visual cycle, RPE65 (3, 4) is the enzyme that isomerizes and cleaves all-trans-retinyl esters (all-trans-RE) into 11-cis-retinol (11-cis-ROL) (5–7), which is then converted to the 11-cis-retinol chromophore of opsins. Recently, our laboratory has shown that RPE65 acts as a leaky isomerase that can isomerize all-trans-RE to both 11-cis-ROL and 13-cis-ROL (8). Although there is no known physiological role for 13-cis-ROL, and 13-cis-retinol cannot regenerate opsin, it has been shown to accumulate as esters in Rdh5 knock-out mouse RPE (9). RDH5 is the primary short-chain retinol dehydrogenase in RPE that oxidizes 11-cis-ROL to 11-cis-retinol (10). Its loss may disrupt the process of mass action crucial for 11-cis-retinoid production in the visual cycle, thereby allowing accumulation of 13-cis-retinoids (9). In our previous studies, we could change the ratio of 11-cis- to 13-cis-ROL with specific conservative mutations of RPE65; for example, F103L increased the proportion of 13-cis-ROL, whereas T147S increased the proportion of 11-cis-ROL (8). We concluded that the mechanism underlying the leaky isomerization implicates a carbocation or a radical cation intermediate with loss of polyene bond order allowing for production of either 11-cis-ROL or 13-cis-ROL by RPE65 and that ultimate specificity for 11-cis isomers occurs downstream by a process of mass action applied by 11-cis-specific binding proteins such as CRALBP and apoprotein opsins (8), as suggested previously (9, 11). We hypothesize that the aromatic side chain-rich environment of the RPE65 substrate-binding cleft is primarily responsible for favoring an 11-cis-specific outcome.

At present, there are crystal structures for three members of the carotenoid oxygenase family: a cyanobacterial apocarote-
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Site-directed Mutagenesis of RPE65—QuikChange XL site-directed mutagenesis kit (Agilent, La Jolla, CA) was used for mutagenesis of the RPE65 open reading frame cloned in pVitro2 (InvivoGen, San Diego, CA). Primer sequences for mutagenesis are given in supplemental Table S1. Mutants were verified by sequence analysis of DNA minipreps (Northwoods DNA, Solway, MN). Validated mutant and wild-type plasmids were purified by Qiagen purification kits (Qiagen, Valencia, CA).

Transient Transfection and Cell Culture—Cell culture methods and transient transfection protocols were previously published (7, 8). In a typical experiment, 3 × 10^7 HEK293-F (Invitrogen) cells were transfected with 30 μg of pVitro2 plasmid (containing RPE65 wild type or mutant) and CRALBP open reading frames and 30 μg of pVitro3 (InvivoGen) plasmid (containing lecithin–retinol acyltransferase (LRAT)) and retinol dehydrogenase 5 (RDH5) open reading frames in the presence of 40 μl of 293Fectin transfection reagent (Invitrogen), all in a total volume of 30 ml. 24 h after transfection, all-trans-ROL was added to a final concentration of 2.5 μM, and the cells were cultured for a further 7 h and then harvested for analysis. Immunoblot analysis was carried out as described previously (8).

Retinoid Extractions and HPLC—Culture fractions of 29-ml volumes of transfected HEK293-F cells were centrifuged, cells were harvested, and retinoids were extracted and saponified as described previously (7). Isomeric retinols were analyzed on 5-μm particle LiChrospher (Alltech, Deerfield, IL) normal phase columns (2 × 250 mm) on an isocratic HPLC system equipped with a diode array UV-visible detector (Agilent 1100/1200 series, Agilent Technologies, New Castle, DE), following Landers and Olson (16) as modified by us (7). Data were analyzed using ChemStation32 software (Agilent).

Construction of Models and Ligand Docking Simulations—A structural alignment template was made from the crystal structures of ACO (Protein Data Bank (PDB) ID: 2BIW), VP14 (PDB ID: 3NPE), and bovine RPE65 (PDB ID: 3KV C). The murine RPE65 sequence was then aligned to this template and submitted to the Swiss Model server (17, 18) to obtain a generalized model that covered loops missing in the crystal structures (residues 109–126 are missing in the bovine RPE65 structure). The substrate-binding cleft was visualized using the MSMS reduced surface package in Chimera (19). It was noted that the three crystal structures appear to represent differing degrees of openness at the presumed substrate-binding cleft, VP14 being the most open and the bovine RPE65 being the most closed. Therefore, an additional open murine RPE65 model based on the VP14 structure was also constructed with the water and dioxygen coordinated to the iron, as seen in VP14 structure. Three-dimensional models of ligands were either obtained from the PubChem database when available or made using the PRODRG program (20, 21). Ligand dockings of substrate (all-trans-retinyl palmitate), intermediates (11-cis- and 13-cis-retinyl palmitate), and products (11-cis-ROL and palmitate) were carried out using Autodock Vina (22). In general, the ligand constraints chosen allowed for all possible torsions. Docking of all-trans, 11-cis, and 13-cis isomers of both esters and retinols were carried out on both the bovine RPE65 structure and the murine models. The top hits from the docking simulations were chosen based on two criteria. The first was that they were in the correct orientation, and the second was that the retinyl O-alkyl bond...
was positioned over the iron center. Except where noted, models were visualized using the PyMOL Molecular Graphics System (Version 1.4, Schrödinger, LLC). The σ-A-weighted difference density for 3KVC, calculated using structure factors deposited by Golczak et al. (31) on the PDB, was obtained from the EDS Uppsala Electron Density Server (EDS) (23).

RESULTS

Structural Modeling—Examination of the three crystal structures of the carotenoid oxygenase family showed that they share a common seven-bladed β-propeller fold, although the three proteins are less than 30% identical in sequence. For the propeller region, the root mean square deviation between RPE65 and ACO is 1.1 Å (for 209 Cα atoms) and between RPE65 and VP14 is 1.4 Å (for 206 Cα atoms). A cleft was observed above the propeller region, over the iron center that was assumed to be the substrate-binding site. Comparison of these clefts showed the RPE65 cleft to be more compact than the ACO cleft. VP14, which contains a dioxygen on the sixth iron coordination position, showed the most extended conformation (Fig. 2). The differences in cleft volumes can be explained by contributions from the dynamic loops in the upper half of the proteins that likely undergo order-disorder transition. In RPE65, these include residues 108–127 (missing in both crystal structures), 258–281 (missing and high B-factor in 3KVC and 3FSN, respectively), and 332–361. This would be modified by the state of the protein during crystallization, i.e. with or without substrate or product. The residues comprising the surface of the cavity in RPE65 were identified using the MarkUs Cavity Server (24). The cavity overlaps with the residual density, as seen in the RPE65 crystal structure, and the surrounding accessible area (supplemental Fig. S1). A total of 62 residues were identified as contributing to this surface (Fig. 3A; supplemental Table S2), with 6 of these contributing to the iron-coordinating complex (3 histidine (out of 4 coordinating) and 3 glutamic acid residues). Of the remaining, 10 were phenylalanine, 5 were tyrosine, 2 were tryptophan, 9 were leucine, and 3 were isoleucine residues. Thus, aromatic residues contribute about 30% of the cavity internal surface with the majority being due to phenylalanines or tyrosines. In the three structures, an arch of phenylalanines (with one leucine replacement in ACO) surmounts the iron center: in RPE65, Phe-61, Phe-312, and Phe-526; in ACO, Phe-69, Phe-311, and Leu-483; and in VP14, Phe-171, Phe-411, and Phe-589. The RPE65 arch residues include Phe-61 that is part of the FDG conserved motif and 2 residues immediately adjacent to two of the iron-coordinating histidines (Phe-312 and Phe-526). Therefore, these 3 residues were selected for mutational studies. (Phe-61 was addressed in our previous study (8), but we provide additional insight into its role herein.) In addition, comparisons of the mouse RPE65 models based on ACO and VP14 indicated a change in the positions of both Phe-103 (mutation of which was previously shown to favor 13-cis-ROL production, (8)) and Tyr-338 in the different conformations. This suggests that the loops on which these residues are located, amino acids 84–133 for Phe-103 and amino acids 333–361 for Tyr-338, are relatively dynamic or mobile (Fig. 3B). This is known to be the case for the amino acids 82–133 loop as the center residues of this loop (amino acids 109–126) were poorly ordered in the RPE65 crystal (15). Loop residues Thr-101, Glu-102, Phe-103, Glu-127, Val-128, and Thr-129 contribute to the internal surface of the cleft (supplemental Table S2). Secondary structure topology of the amino acids 333–361 loop reveals three predicted α helices (15), and residues in two of these helices (Val-337, Tyr-338, Leu-341, and Leu-346) contribute to the internal surface of the cleft (supple-
Inactivating Mutations—Mutations of the “Phe arch” residues Phe-61 or Phe-312 to other conservative substitutions severely diminished or eliminated all catalytic activity (Table 1). Phe-61, part of the FDG motif seen in all carotenoid oxygenases, is located on the top of the Phe arch. It is positioned right above the iron, on the other side of the substrate in the cleft. Phe-61 displays negligible activity upon any substitution, suggesting a conserved role for this residue. Tyrosine, due to its hydrophilic nature, may destroy the putative carbocation/cat-ion radical intermediate, and tryptophan may be too bulky a substituent for that location, sterically hindering the substrate. Phe-312 is also irreplaceable, suggesting a high invariance at this position. Due to its location and proximity to the substrate, Phe-312 possibly plays a role in steric shielding to allow proper substrate-binding and/or variable loops. In the inactivating mutations, 13-cis-retinol production was comparable with thermal levels, as described previously (8).

Modulating Mutations—In contrast to the inactivating effects of mutations at Phe-61 and Phe-312, substitution at the 3rd Phe arch residue Phe-526 altered but did not abolish activity. Mutation with any residue other than alanine decreased 11-cis-ROL (but not 13-cis-ROL) production in excess of what would be expected by the lower expression levels of these mutants (Table 2 and Fig. 4A). However, the F526A mutation promoted leaky isomerase activity, selectively increasing 13-cis-ROL production with a concomitant reduction in 11-cis-ROL production. Tyr-338, one of the dynamic residues seen in the crystal structures, selectively reduced 11-cis-ROL production in all its mutants. Y338L, Y338F, and Y338W produced the 13-cis-ROL product in similar amounts to wild type when corrected for expression (Table 2 and Fig. 4B), and Y338A displayed increased 13-cis-ROL (1.63-fold increase) similar to the F103L or F103I mutations previously reported, although somewhat more severe in the reduction of 11-cis production (8). This result suggests that Tyr-338 substitutions also reduce 11-cis-ROL isomerization specifically, but that 13-cis isomerization is essentially unaffected by most substitutions or increased by the Y338A mutation. We observed this increased 13-cis-retinol production by the Y338A mutant to be elevated when compared with thermal production of 13-cis-retinol either in the presence or in the absence of wild-type RPE65 at all time points analyzed (supplemental Fig. S2). We suggest that the major effect of Tyr-338, just as in Phe-103 and Phe-526, is to stabilize the cis isomers as they are being positioned in the cleft prior to cleavage; 13-cis-RE, being lower in its energy of formation, may be more favored to bind in the cleft of these mutants as they are more permissive.

Aromatic Residues Govern RPE65 Isomerization—The bovine RPE65 crystal structure, derived from native material, contains residual densities in cleft pockets. When we docked the products, 11-cis-ROL and palmitate, they traced the density areas (Fig. 5), suggesting that these residual densities may represent the presence of the products in the crystal. There is a bend in the residual electron density going around Phe-418 (highlighted in Fig. 5), indicating that this residue helps shape the cavity. To evaluate the docking simulations, two assumptions were made: first, that the retinyl ester had to be positioned in the phenylalanine arch prior to cleavage, and second, that the retinyl ester O-alkyl bond should be positioned over the iron catalytic center in a specific orientation. Docking of all-trans-retinyl palmitate, however, was unable to meet the basic criteria as it was never docked appropriately over the iron center in any simulation. However, when all-trans-retinyl palmitate was docked on the mouse RPE65 model based on the open VP14 crystal, the highest affinity docking placed it in a cleft between the two loops carrying Phe-103 and Tyr-338.

### Table 1

Expression and cis-retinol production of HEK293-F cells with mutants inactivating RPE65 activity

| Expressiona | ± S.D. | 11-cis-ROLa | ± S.D. | 13-cis-ROLa | ± S.D. |
|-------------|--------|-------------|--------|-------------|--------|
| WT          | 1      | 1           | 1      | 1           | 1      |
| F61A        | 0.11   | 0.04        | 0      | 0.25        | NA     |
| F61Y        | 0.56   | 0.08        | 0.08   | 0.42        | 0.07   |
| F61L        | 0.39   | 0.15        | 0.01   | 0.37        | 0.05   |
| F61W        | 0.22   | 0.05        | 0.02   | 0.38        | 0.06   |
| F312L       | 0.21   | 0.11        | 0.03   | 0.32        | 0.04   |
| F312Y       | 0.2    | 0.08        | 0.02   | 0.39        | 0.03   |
| F312W       | 0.3    | 0.1         | 0      | 0.29        | 0.03   |
| F418A       | 0.13   | 0           | 0      | 0.51        | 0.11   |
| F418L       | 0.18   | 0           | 0.04   | 0.42        | 0.1    |
| F418Y       | 0.12   | 0.08        | 0      | 0.4         | 0.08   |
| F418W       | 0.15   | 0.04        | 0      | 0.39        | 0.11   |

a Data are given as -fold change from wild-type expression ± S.D.; n ≥ 3.

b Data are given as -fold wild-type RPE65 isomerase activity ± S.D.; n ≥ 3.

NA, not applicable.
Use of the mouse model was required to model the loop (carrying Phe-103), which was unresolved in the bovine RPE65 crystal structure, as indicated previously (15). (This loop is almost perfectly conserved between bovine and mouse RPE65s with only one conservative substitution at amino acid 124 (Arg in bovine replaced by Lys in mouse)). In contrast, both 11-cis-retinyl palmitate and 13-cis-retinyl palmitate docked in the bovine RPE65 crystal structure in the presence of water modeled into coordination position 5 of the iron (Fig. 6, B and C). However, if dioxygen was modeled into position 6 (as in the VP14 crystal), no retinyl palmitate docking within the phenylalanine arch was obtained (not shown). These results suggest that the early steps of isomerization of retinyl esters occur in the capture and binding of the substrate, thus allowing it to fit appropriately over the iron center for cleavage.

### Modeling of Substrate and Intermediate Docking in Mutant RPE65

Docking simulations were also carried out on mutants of the various residues studied. Both F61L and F312L showed docked substrates with the ester bond significantly off position when compared with wild type (Fig. 7, A and C). Leucines occur naturally in paralogous positions to Phe-61 and Phe-312 in some carotenoid oxygenases of plants and in the carotenoid isomero-oxygenases of insects (25), respectively. However, in RPE65, mutations to these residues render RPE65 inactive, which suggests that the role of these residues is to orient the substrate in such a way that the ester cleavage activity of the enzyme can occur. F418L showed docked substrate with distortions around the 11-cis-retinol (blue), palmitate (light gray), and iron (orange) are depicted. 

### Table 2

| Expression | ± S.D. | 11-cis-ROL | ± S.D. | 13-cis-ROL | ± S.D. | 11-cis expression | 13-cis expression |
|------------|--------|------------|--------|------------|--------|------------------|------------------|
| WT         | 1      | 1          |        | 1          |        |                  |                  |
| F526A      | 0.88   | 0.12       | 0.68   | 0.08       | 1.30   | 0.24             | 0.77             | 1.48             |
| F526Y      | 0.54   | 0.09       | 0.10   | 0.02       | 0.56   | 0.13             | 0.18             | 1.04             |
| F526L      | 0.46   | 0.10       | 0.00   | 0.02       | 0.54   | 0.14             | 0.00             | 1.11             |
| F526W      | 0.38   | 0.05       | 0.03   | 0.02       | 0.54   | 0.14             | 0.07             | 1.42             |
| Y338A      | 0.31   | 0.11       | 0.10   | 0.03       | 1.63   | 0.22             | 0.31             | 5.25             |
| Y338L      | 0.29   | 0.10       | 0.03   | 0.03       | 0.62   | 0.10             | 0.09             | 2.14             |
| Y338W      | 0.24   | 0.11       | 0.01   | 0.02       | 0.74   | 0.16             | 0.05             | 3.08             |
| Y338F      | 0.08   | 0.06       | 0.15   | 0.04       | 0.71   | 0.15             | 1.85             | 8.83             |
| F103L      | 0.61   | 0.12       | 0.28   | 0.01       | 1.22   | 0.11             | 0.46             | 2.00             |
| F103W      | 0.57   | 0.11       | 0.24   | 0.01       | 0.61   | 0.04             | 0.42             | 1.07             |
| F103Y      | 0.33   | 0.07       | 0.21   | 0.03       | 0.36   | 0.06             | 0.64             | 1.09             |
| F103I      | 0.27   | 0.06       | 0.03   | 0.01       | 0.44   | 0.04             | 0.11             | 1.63             |

* Data are given as -fold change from wild-type expression ± S.D.; n ≥ 3.
* Data are given as -fold wild-type RPE65 isomerase activity ± S.D.; n ≥ 3.

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![Figure 4. Mutations of Phe-526 and Tyr-338 modulate product isomer specificity of RPE65 isomerase activity.](image)

![Figure 5. Product docking in RPE65. Docking of 11-cis-ROL (purple) and palmitate (light gray) in the cleft of the bovine RPE65 crystal structure is shown. The darker green mesh represents α-A-weighted difference density (contoured at 3σ) as observed in the 3KVC crystal structure.](image)
to be positioned in the effective orientation, thus allowing 13-cis product formation (Fig. 7, B and D). The reduction of 11-cis isomerization activity in this mutant could reflect the requirement of an aromatic residue to stabilize the 11-cis isomer as the docking simulation appears to be feasible. Docking of both 11-cis-retinyl palmitate and 13-cis-retinyl palmitate in F526L is also essentially the same as in the wild type. However, given the complete loss of activity with respect to 11-cis-ROL production, it is probable that formation and/or stabilization of the 11-cis isomer is compromised by a leucine mutation in this member of the phenylalanine arch. Likewise, the F103L mutation and the Y338A mutation showed apparently normal docking for both isomers (Fig. 7, B and D), suggesting a failure of isomerization to 11-cis-ROL in these mutants. Overall, the dockings show a clustering of the substrate conformations in inactivating mutants and modulating mutants into two respective groups.

**DISCUSSION**

In this study, we demonstrate, using site-directed mutagenesis and structure-based modeling, that an assemblage of aromatic residues in the substrate-binding cleft of RPE65 governs the progression and specificity of retinoid isomerization by this enzyme. These data extend our previous conclusions (8) that RPE65 utilizes a cationic intermediate (carbocation or radical cation) in its mechanism by showing that multiple aromatic residues contribute to the retinyl moiety-binding cavity within which isomerization occurs. This environment is a result of contributions from the various aromatic side chains we have studied and appears to be crucial to isomerization. Furthermore, we interpret the residual unresolved density in the RPE65-binding cleft as being due to the products of the reaction, not only fatty acid, as indicated previously (15), but also 11-cis-retinol. Finally, based on docking of various modeled substrates, we can discern a temporal sequence for the progression of retinoid isomerization by RPE65.

Based on our previous findings that we could modulate cis isomer specificity by mutation of single residues (8), we determined that isomerization of all-trans-ROL to 11-cis-ROL occurs via a cationic intermediate with loss of polyene bond order as this alternative was the only one that could accommodate possible formation of isomers besides 11-cis. Although a carbocation intermediate was considered originally based on
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Prior studies (9, 11, 26), we concluded that the data could just as easily support a radical cation intermediate. Support for the latter possibility was provided by our finding that aromatic lipophilic spin traps such as N-tert-butyl-α-phenylnitrone (PBN) inhibit RPE65 activity (26), and this was corroborated by others (27).

Phe-103 was one of the original residues that we showed to modulate isomer specificity (8). Although it is itself resolved, Phe-103 is located on a flexible loop that is not completely modeled in the RPE65 crystal structure (15). Paralogous to Phe-113 in ACO (12, 13), Phe-103 is one of the aromatic residues of the substrate-binding tunnel of RPE65. It is part of the conserved EFG triplet motif in metazoan carotenoid oxygenases, of which the glycine is invariably conserved throughout the family. Herein we show that two other tunnel aromatic residues, Tyr-338 and Phe-526, are also required for 11-cis product specificity. Of these, Tyr-338 is also located on a loop, what we term the metazoan loop because it is conserved only among animal carotenoid oxygenases. This loop is observed in the RPE65 crystal structure, and the aromatic side chain of Tyr-338 inserts into the tunnel. Phe-526 is part of what we have termed the phenylalanine arch. A similar arrangement of phenylalanines also occurs in VP14, where it is proposed to be important for “caging” the substrate over the iron (14), a role we predict to be repressed in RPE65 by the paralogous residues Phe-61, Phe-312, and Phe-526. The opening up of space around residues 526 and 338, in their respective Ala mutants, probably allows for the more energetically favorable 13-cis product to be formed. This is similar to the F103L mutant, which also affects product specificity. The varying effects of the Phe arch residues, located at the core of RPE65, point to their critical role in the RPE65 retinol isomerase mechanism. Along with Phe-103, we conclude that Tyr-338 and Phe-526 also constrain cleft shape and volume to favor production of 11-cis-ROL. Such a key role for phenylalanine in binding polyene chains has previously been shown for carotenoid binding in light-harvesting complex where Garcia-Martín et al. (28) showed that about 25% of total protein-carotenoid contacts are with phenylalanine residues. Conversely, phenylalanines, unlike tyrosines, are rarely involved in catalysis (29).

An important finding of the cavity modeling of RPE65, ACO, and VP14 is the observation of the relative constriction of the RPE65 cavity when compared with those of ACO and VP14. We propose that this may partly reflect the closed state of the RPE65 crystal (15) and the mechanistic requirements for the assured conformational molding of the delocalized retinyl moiety to the 11-cis conformation. Selective mutational alteration of the cavity, such as we have performed here with residues Tyr-338 and Phe-526, extends our findings on Phe-103 (and Thr-147 (8)). These alterations disturb the predicted molecular molding, allowing isomerization to the more thermodynamically favorable 13-cis isomer. Therefore, in all these mutants, of Phe-103, Phe-526, and Tyr-338, we conclude that the dominant effect is reduced stringency disfavoring 11-cis isomerization rather than preferential selection for 13-cis-RE binding.

Consideration of these data allows us to put forward a temporal sequence for isomerization by RPE65. We propose that: 1) all-trans-RE enters the open ready state conformation of RPE65, perhaps via initial recognition of the β-ionone ring; 2) the bound all-trans-RE undergoes a single electron oxidation by an as yet unidentified means, resulting in the delocalization of the retinyl bond order; 3) the delocalized retinyl moiety in the substrate-binding cleft is constrained by the geometry of the closing pocket to adopt the 11-cis conformation as RPE65 transitions to the state captured by the RPE65 crystal structure (15); 4) the O-alkyl bond is positioned over and interacts with the iron, which catalyzes cleavage; 5) attack by water completes hydrolysis and restores retinyl bond order; 6) 11-cis-retinol is released; and 7) RPE65 returns to the open ready state. This chronology is different from the mechanism proposed by Kiser et al. (15) in a number of respects. These authors proposed that initially, the oxygen of the all-trans-RE acyl bond interacts with the iron, resulting in cleavage of the O-alkyl bond. They proposed that the leaving of the palmitate anion results in formation of a retinyl carbocation, which can then be constrained to the 11-cis conformation. However, our docking studies indicate that this is too late for delocalization to occur as the predicted binding energy of all-trans-RE is quite unfavorable in the closed state modeled in the RPE65 crystal structure. Instead, we postulate that the retinyl moiety must be in a delocalized state prior to approach of the O-alkyl bond to the iron center and its ultimate cleavage. We propose that single-electron oxidation generating a radical cation intermediate is the source of the delocalized state. An earlier loss of bond order allows the retinyl moiety to be constrained to the 11-cis conformation, which has a favorable binding energy in the closed state, whereas the O-alkyl bond is brought into register for cleavage. This point is supported by the inhibitory effect of aromatic lipophilic spin traps (such as PBN) on RPE65 activity, an effect that would be less likely if delocalization via radical formation occurs later as the putative PBN-retinyl ester adduct is not predicted to be able to access the binding cleft in its later closed state (26). Consequently, we propose that O-alkyl cleavage occurs later. However, we do concur with the proposal of Kiser et al. (15, 30) that the bond-cleaving activity of RPE65 has become that of O-alkyl cleavage, with retinol isomerization an acquired function. Thus, by strategic placement of aromatic residues to modulate the geometry and chemistry of its catalytic cleft, RPE65 has evolved from a carotenoid oxygenase to an isomerohydrolase role to drive the vertebrate retinal visual cycle.

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REFERENCES
1. Saari, J. C. (2000) Biochemistry of visual pigment regeneration: the Friedenwald lecture. Invest. Ophthalmol. Vis. Sci. 41, 337–348
2. Lamb, T. D., and Pugh, E. N., Jr. (2004) Dark adaptation and the retinoid cycle of vision. Prog. Retin. Eye Res. 23, 307–380
3. Hamel, C. P., Tsilou, E., Pfeiffer, B. A., Hooks, J. I., Detrick, B., and Redmond, T. M. (1993) Molecular cloning and expression of RPE65, a novel retinal pigment epithelium-specific microsomal protein that is post-transcriptionally regulated in vitro. J. Biol. Chem. 268, 15751–15757
4. Redmond, T. M., Yu, S., Lee, E., Bok, D., Hamasaki, D., Chen, N., Goletz, P., Ma, J. X., Crouch, R. K., and Pleifer, K. (1998) Rpe65 is necessary for production of 11-cis-vitamin A in the retinal visual cycle. Nat. Genet. 20,
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344–351
5. Jin, M., Li, S., Moghrabi, W. N., Sun, H., and Travis, G. H. (2005) Rpe65 is the retinoid isomerase in bovine retinal pigment epithelium. Cell **122**, 449–459

6. Moiseyev, G., Chen, Y., Takahashi, Y., Wu, B. X., and Ma, J. X. (2005) RPE65 is the isomerase domain in the retinoid visual cycle. Proc. Natl. Acad. Sci. U.S.A. **102**, 12413–12418

7. Redmond, T. M., Poliakov, E., Yu, S., Tsai, J. Y., Lu, Z., and Gentleman, S. (2005) Mutation of key residues of RPE65 abolishes its enzymatic role as isomerohydrolase in the visual cycle. Proc. Natl. Acad. Sci. U.S.A. **102**, 13658–13663

8. Redmond, T. M., Palczewski, K., Kuo, S., Chander, P., and Gentleman, S. (2010) RPE65, visual cycle retinol isomerase, is not inherently cis-specific: support for a carbocation mechanism of retinol isomerization. J. Biol. Chem. **285**, 1919–1927

9. Maeda, A., Maeda, T., Imanishi, Y., Golczak, M., Moiseyev, G. P., and Palczewski, K. (2006) Achromatophores in mouse models of congenital blindness: formation and storage of retinyl esters. Biochemistry **45**, 4210–4219

10. Kuo, S., and Schulz, G. E. (2010) Retinol dehydrogenases (RDHs) in the visual cycle. Exp. Eye. Res. **91**, 788–792

11. McBee, J. K., Kuksa, V., Alvarez, R., de Lera, A. R., Prezhdo, O., Haeseleer, M., Kiser, P. D., Golczak, M., Lodowski, D. T., Chance, M. R., and Palczewski, K. (2006) Structural and biological aspects of RPE65 isomerohydrolase activity. J. Biol. Chem. **281**, 267–269

12. Kloer, D. P., Ruch, S., Al-Babili, S., Beyer, P., and Schulz, G. E. (2005) Isomerization of all-trans-retinol to cis-retinol in bovine retinal pigment epithelial cells: dependence on the specificity of retinoid-binding proteins. Biochemistry **39**, 11370–11380

13. Kuo, S., Ruch, S., Al-Babili, S., Beyer, P., and Schulz, G. E. (2005) The structure of a retinal-forming carotenoid oxygenase. Science **308**, 267–269

14. Mandal, M. N., Moiseyev, G. P., Elliott, M. H., Kasus-Jacobi, A., Li, X., Chen, H., Zheng, L., Nikolaeva, O., Floyd, R. A., Ma, J. X., and Anderson, R. E. (2011) NinaB combines carotenoid oxygenase and retinoid isomerase activity in a single polypeptide. Proc. Natl. Acad. Sci. U.S.A. **105**, 19000–19005

15. Kiser, P. D., Golczak, M., Lodowski, D. T., Chance, M. R., and Palczewski, K. (2009) The crystal structure of RPE65, the retinoid isomerase of the visual cycle. Proc. Natl. Acad. Sci. U.S.A. **106**, 17325–17330

16. Oberhauser, V., Voolstra, O., Bangert, A., von Lintig, J., and Vogt, K. (2008) NINA combines carotenoid oxygenase and retinoid isomerase activity in a single polypeptide. Proc. Natl. Acad. Sci. U.S.A. **105**, 19000–19005

17. Nishi, H., Nakamura, K., and Watanabe, Y. (2003) Aromatic substrates for RPE65 isomerohydrolase activity. Biochemistry **42**, 7970–7978

18. Chen, H., Zheng, L., Nikolaeva, O., Floyd, R. A., Ma, J. X., and Anderson, R. E. (2011) NINA combines carotenoid oxygenase and retinoid isomerase activity in a single polypeptide. Proc. Natl. Acad. Sci. U.S.A. **108**, 13429–13434

19. Sanner, M. F., Olson, A. J., and Spehner, J. C. (1996) Reduced surface: an efficient way to compute molecular surfaces. Biopolymers **38**, 305–320

20. van Aalten, D. M., Bywater, R., Findlay, J. B., Hendlich, M., Hooft, R. W., and Vriend, G. (1996) PRODRG, a program for generating molecular topologies and unique molecular descriptors from coordinates of small molecules. J. Comput. Aided Mol. Des. **10**, 255–262

21. Schüttelkopf, A. W., and van Aalten, D. M. (2004) PRODRG: a tool for high-and-throughput crystallography of protein-ligand complexes. Acta Crystallogr. D Biol. Crystallogr. **60**, 1355–1363

22. Trott, O., and Olson, A. J. (2010) AutoDock Vina: improving the speed and accuracy of docking with a new scoring function, efficient optimization, and multitithreading. J. Comput. Chem. **31**, 455–461

23. Kleywegt, G. J., Harris, M. R., Zou, J. Y., Taylor, T. C., Wahlby, A., and Jones, T. A. (2004) The Uppsala Electron-Density Server. Acta Crystallogr. D Biol. Crystallogr. **60**, 2240–2249

24. Fischer, M., Zhang, Q. C., Dey, F., Chen, B. Y., Honig, B., and Petrey, D. (2011) MarkUs: a server to navigate sequence-structure-function space. Nucleic Acids Res. **39**, W357–W361

25. Kleywegt, G. J., Harris, M. R., Zou, J. Y., Taylor, T. C., Wahlby, A., and Jones, T. A. (2004) The Uppsala Electron-Density Server. Acta Crystallogr. D Biol. Crystallogr. **60**, 2240–2249

26. Fischer, M., Zhang, Q. C., Dey, F., Chen, B. Y., Honig, B., and Petrey, D. (2011) MarkUs: a server to navigate sequence-structure-function space. Nucleic Acids Res. **39**, W357–W361

27. Kleywegt, G. J., Harris, M. R., Zou, J. Y., Taylor, T. C., Wahlby, A., and Jones, T. A. (2004) The Uppsala Electron-Density Server. Acta Crystallogr. D Biol. Crystallogr. **60**, 2240–2249

28. Garcia-Martín, A., Pazur, A., Wilhelm, B., Silber, M., Robert, B., and Mandal, M. N., Moiseyev, G. P., Elliott, M. H., Kasus-Jacobi, A., Li, X., Chen, H., Zheng, L., Nikolaeva, O., Floyd, R. A., Ma, J. X., and Anderson, R. E. (2011) NINA combines carotenoid oxygenase and retinoid isomerase activity in a single polypeptide. Proc. Natl. Acad. Sci. U.S.A. **108**, 13429–13434

29. Mandal, M. N., Moiseyev, G. P., Elliott, M. H., Kasus-Jacobi, A., Li, X., Chen, H., Zheng, L., Nikolaeva, O., Floyd, R. A., Ma, J. X., and Anderson, R. E. (2011) α-Phenyl-N-tert-butylnitrone (PBN) prevents light-induced degeneration of the retina by inhibiting RPE65 protein isomerohydrolase activity. J. Biol. Chem. **286**, 32491–32501

30. Garcia-Martín, A., Pazur, A., Wilhelm, B., Silber, M., Robert, B., and Mandal, M. N., Moiseyev, G. P., Elliott, M. H., Kasus-Jacobi, A., Li, X., Chen, H., Zheng, L., Nikolaeva, O., Floyd, R. A., Ma, J. X., and Anderson, R. E. (2011) α-Phenyl-N-tert-butylnitrone (PBN) prevents light-induced degeneration of the retina by inhibiting RPE65 protein isomerohydrolase activity. J. Biol. Chem. **286**, 32491–32501

31. Schüttelkopf, A. W., and van Aalten, D. M. (2004) PRODRG: a tool for high-and-throughput crystallography of protein-ligand complexes. Acta Crystallogr. D Biol. Crystallogr. **60**, 1355–1363

32. Trott, O., and Olson, A. J. (2010) AutoDock Vina: improving the speed and accuracy of docking with a new scoring function, efficient optimization, and multitithreading. J. Comput. Chem. **31**, 455–461

33. Kleywegt, G. J., Harris, M. R., Zou, J. Y., Taylor, T. C., Wahlby, A., and Jones, T. A. (2004) The Uppsala Electron-Density Server. Acta Crystallogr. D Biol. Crystallogr. **60**, 2240–2249