Spectral Signatures of Canthaxanthin Translocation in the Orange Carotenoid Protein

Natalia B. Pigni, Kevin L. Clark, Warren F. Beck, and José A. Gascón*

ABSTRACT: The orange carotenoid protein (OCP) is involved in the photoprotective processes in cyanobacteria via nonphotochemical quenching. Triggered by blue-green light absorption, the carotenoid chromophore undergoes translocation, displacing around 12 Å from the C-terminal domain (CTD) to the N-terminal domain (NTD). The detailed molecular rearrangements that occur within the carotenoid and the protein during this process remain largely elusive. By using a combination of molecular dynamics, well-tempered metadynamics, and hybrid quantum mechanical/molecular mechanical (QM/MM) calculations, we were able to mimic the translocation of the carotenoid from the inactive OCP0 and obtain metastable red-shifted states in the photoactivation mechanism, replicating the $\lambda_{\text{max}}$ values of reference experimental spectra. In addition, our simulations give insight into the structure of the red-shifted form of the inactive state of OCP.

INTRODUCTION

Photosynthetic organisms have developed efficient ways for light energy capture and storage, accompanied by a refined defense mechanism against varying light conditions. Non-photochemical quenching (NPQ) protects these organisms from the potentially damaging effects of excess light. In cyanobacteria, NPQ is facilitated by the orange carotenoid protein (OCP).1 OCP is a 35 kDa water-soluble protein consisting of an all $\alpha$-helical N-terminal domain (NTD) and a mixed $\alpha$-helical/$\beta$-sheet C-terminal domain (CTD) (Figure 1A). Encapsulated between these two domains, a ketocarotenoid interacts with the protein environment through hydrogen bonds and hydrophobic contacts.2,3 Under normal light conditions, OCP is found in its inactive form (OCPO) characterized by an orange coloration in solution. After photoactivation by blue-green light, OCP undergoes a series of reversible structural modifications leading to the active form known as OCP*. Either spontaneously or facilitated by the fluorescent recovery protein (FRP), OCP* is then reconverted to OCP0.4 OCP can bind different carotenoids, including 3′-hydroxyechinenone (3′-hECN), echinenone, and canthaxanthin (CAN, Figure 1B). All of these ketocarotenoids have a common carbonyl group that stabilizes the $\beta1\beta$-ionone ring of OCP0 through hydrogen bonds with two key residues from the CTD, Tyr201 and Trp288.5 Stabilization and disruption of this pair of hydrogen bonds is apparently essential for the photoactivation cycle because these hydrogen-bonding interactions are highly conserved in OCP structures from different cyanobacteria.5

Recent studies of the fluorescence spectra from 3′-hECN and CAN in OCP0 show that the two ketocarotenoids differ in their ability to form an additional hydrogen bond between the $\beta2\beta$-ionone ring with Leu37 in the NTD. Interestingly, the referenced studies led to the conclusion that large-amplitude motions of the $\beta$-ring contribute to the earliest events following absorption of light that leads to photoactivation to the OCP* state.6

Protein structural details of the photoactivation mechanism have been recently described using time-resolved electronic and vibrational spectroscopy.7 Optical excitation of the mid-IR absorption band populates the $S_2$ state of the carotenoid followed by ultrafast internal conversion to the dark $S_1$ state, which is coupled to an intramolecular charge-transfer (ICT) state. The $S_1$/ICT state further decays to the ground state. With a yield of $\sim1.5\%$, this overall process results in the photoproduct P1 (Figure 1C), which is characterized by the rupture of hydrogen bonds between the carotenoid and Tyr201 and Trp288. Around 40% of this intermediate photoprocess is temporarily stabilized by protein rearrangements in the NTD side (photoprocess P2). These movements lead further on ($\sim10\mu$s) to a remarkable 12 Å translocation of the carotenoid toward the NTD.5 The resulting photoproduct ...

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(P3) shows a UV−vis spectrum with a close resemblance to that reported for OCP\textsuperscript{R}\textsuperscript{9}−\textsuperscript{13}. Lastly, on a time scale longer than 1 ms, the two domains separate, leading to the fully active OCP\textsuperscript{R} state.\textsuperscript{10}−\textsuperscript{13}

The structure of OCP\textsuperscript{O} bound to canthaxanthin in solution is characterized by an absorption spectrum with a $\lambda_{\text{max}}$ at $\sim$477 nm.\textsuperscript{14,15} After photoactivation, OCP\textsuperscript{R}.CAN shows a red-shifted $\lambda_{\text{max}}$ at $\sim$525−530 nm.\textsuperscript{9,15} The intermediate states (P1−P3) are also reported to have overall red-shifted spectra with respect to the OCP\textsuperscript{O} steady-state.\textsuperscript{8} There are several crystal structures of OCP\textsuperscript{O} available, such as OCP\textsuperscript{O} bound to CAN (PDB: 4XB5), but no structural information is available for any of the intermediate photoproducts or the fully active OCP\textsuperscript{R}. However, the structure of the isolated NTD bound to CAN, known as red carotenoid protein (RCP), was successfully crystallized and described in 2015 (PDB: 4XB4) by Kerfeld’s group, representing a valuable target model of OCP\textsuperscript{R}.\textsuperscript{9} In fact, RCP in solution displays a UV−vis spectrum almost identical to that of OCP\textsuperscript{R} with a $\lambda_{\text{max}}$ reported at $\sim$525−530 nm.\textsuperscript{9,14}

Despite the progress in describing many aspects of the photoactivation mechanism, there are still a number of open questions about the various stages of the cycle, especially regarding the detailed carotenoid rearrangement that disrupts the hydrogen-bond network, the structural dynamics of carotenoid translocation, and domain separation.\textsuperscript{8,16}

In this work, we elucidate the structural changes that might occur within the protein as the carotenoid emerges into the NTD domain. While the detailed structural changes that occur during the formation of P1 still remain elusive to experimental and theoretical characterization, we construct a path that directly connects the inactive OCP\textsuperscript{O} structure with a P2-like state, followed by a P3-like state, where the carotenoid is displaced by 12 Å toward the NTD side. This pathway represents the hypothetical process where the carotenoid translocates from its inactive state OCP\textsuperscript{O} to the red state OCP\textsuperscript{R} in the dark. To accomplish this, we perform well-tempered metadynamics simulation to steer the carotenoid from CTD to NTD. Metadynamics has been successfully applied to tackle questions in different fields including materials science, chemical reactions, and biomolecular dynamics.\textsuperscript{17} As an illustrative example, its application has provided insights to understand ligand translocation mechanisms in protein gorges/channels.\textsuperscript{18} With regard to the study of OCP, the use of this methodology has been recently reported to enhance the sampling of CAN conformational freedom in RCP, with an exclusive focus on the geometry of rings $\beta$1 and $\beta$2.\textsuperscript{16}

The metadynamics simulations are combined with QM/MM calculations to characterize the spectral signatures of the different states of the translocation process. We further offer an explanation for the experimentally observed heterogeneity of the inactive state of OCP\textsuperscript{O}, which presents a red-shifted spectral component.

### COMPUTATIONAL METHODS

#### System Preparation

The crystal structures of OCP\textsuperscript{O} (PDB: 4XB5) and RCP (PDB: 4XB4)\textsuperscript{9} containing the protein and the carotenoid were processed with Maestro (Schrödinger 2019-4). Water molecules and glycerol were deleted. For the dimeric structure of RCP, only one monomer was kept (chain A). The Protein Preparation application was used to preprocess the complex, assigning bond orders, adding hydrogens, and filling in missing loops with Prime (residues 51−59 in RCP). For the alternative positions of side chains reported in the crystal structures, the most populated ones were kept. The particular residues for each structure, with the respective occupancy value of the selected position given between parentheses, were the following: Glu118 (0.59) for OCP; Cys84 (0.79), Arg112 (0.53), and Ser147 (0.51) for RCP. For the alternative positions of side chains reported in the crystal structures, the most populated ones were kept. The particular residues for each structure, with the respective occupancy value of the selected position given between parentheses, were the following: Glu118 (0.59) for OCP; Cys84 (0.79), Arg112 (0.53), and Ser147 (0.51) for RCP. Optimizations of hydrogen-bond assignment was performed with default parameters (pH 7.0), and a restrained
minimization was applied for hydrogens only. A 10 Å buffer size, orthorhombic box was constructed using the TIP3P water model, and volume minimization was applied with a resulting number of water molecules equal to 5814 and 10 121 for RCP and OCP, respectively. A number of Na⁺ ions were added to neutralize the negative charge of the system. All preliminary minimizations and MD simulations used the force field OPLS3e.20

**Molecular Dynamics and Metadynamics.** MD simulations were run for solvated systems on Nvidia GPU hardware with Desmond (Schrödinger 2019-4). Prior to MD production, the default relaxation protocol was performed (detailed in SI). Classical MD simulations were run for a total time of 1000 ns using the NPT ensemble (300 K, 1 atm). Snapshots were extracted every 20 ns and fed into QSite for QM/MM calculations. Additional information about the MD analysis, including root mean square deviation (RMSD) calculation, description of protein–ligand interactions, ligand solvent accessible surface area (SASA), and torsions data, is reported in SI.

WT-metadynamics simulations were run using the solvated system prepared for OCPO. The collective variable (CV), to drive the displacement of CAN, was defined as the distance from the center of mass of 5 carbon atoms belonging to 5 residues on the NTD side (Glu34-Cβ, Thr80-Cβ, Cys84-Cβ, Pro126-Ca, Tyr129-Cα1) to C6′ on the β-ring of CAN (Figure 1B). A wall of 20 Å was set as a limit for the explored coordinate space. This CV serves as an effective reaction coordinate for the translocation process and the protein response. The width of the Gaussian potential was σ = 0.06, and the initial height W₀ = 0.03 kcal mol⁻¹. The deposition time was Δt = 0.2 ps, and the bias factor of WT-metadynamics was set to γ = 10. There were 20 repetitions run varying the seed randomly. The total time for each metadynamics simulation was 100 ns. The ensemble class and relaxation protocol were the same as those applied for classical MD. Snapshots were taken every 1 ns to perform the electronic structure calculations and to analyze the carotenoid geometry and protein–ligand interactions.

**Electronic Structure Calculations.** Quantum mechanical/molecular mechanical (QM/MM) calculations were run on snapshots extracted from MD and metadynamics simulations. Solvent and ions were removed. The QM region was defined as CAN while the rest of the protein was at the MM-OPLS3e level. Since the force field largely overestimates the bond length alternation (BLA) of the carotenoid, a local geometry optimization of CAN was performed leaving the rest of the protein frozen. This optimization was performed using QSite (Schrödinger 2019-3) at the DFT B3LYP/LACVP* level. Excited state energies were then calculated on the optimized structures with the Tamm–Dancoff approximation at the CAM-B3LYP/LACVP* level. No ad hoc corrections, shifts, or parametrizations of any kind were employed to fit the calculated TDDFT values with experimental data. An important reason for choosing CAM-B3LYP as the functional is that it is one of the best at reproducing the correct excitation energy of the isolated CAN in solvent, which ranges between 470 and 490 nm.21,22 Indeed, we made a comparison with a number of other functionals including SVWN5, BP86, VSXC, BHHLYP, LC-WHPE, X3LYP, and B3LYP, and we found that CAM-B3LYP was the best at reproducing the solution phase results. This benchmark was done using an optimized geometry at the B3LYP geometry and a conductor-like screening model with methanol as the implicit solvent.

Protein solvation effects were treated using the Poisson–Boltzmann implicit solvent method implemented in QSite. This combination of two different levels of DFT and/or basis sets, one for ground state optimization and the other for excited state energies, has been successfully used before, particularly in the context of carotenoids.23–25 The incorporation of solvent (via Poisson–Boltzmann in this case) is essential to reproduce the experimental values of UV–vis absorption. Such a substantial solvent effect has been pointed out recently by Guberman-Pfeffer et al. in the multi-chromophoric peridinin-chlorophyll α-protein (PCP).26 As shown in the SI, use of the explicit waters from MD produces very similar results (Table S2). The average excitation for both OCPO and RCP differs by approximately 4 nm, with the explicit case being slightly more blue-shifted in both OCP and RCP.

One-sample t tests were run to analyze the results from the optimized MD snapshots of the reference models, RCP and OCPO, calculating the confidence intervals at a level of 95% (N = 51, for each MD). Smoothing of metadynamics data was done by using a moving average of 5 points. The method used to calculate bond length alternation (BLA) is described in the Supporting Information.

### RESULTS AND DISCUSSION

**Validation of Reference Models: OCPO and RCP.** The crystal structures of OCPO-CAN (PDB: 4XB5) and RCP-CAN (PDB: 4XB4) were used as the starting points to run our MD simulations and QM/MM calculations. UV–vis experimental data previously reported in the literature were taken as the reference values to validate our methodology. Since the protein system exists in a dynamical equilibrium between multiple conformations, we aimed to obtain a finite-temperature thermodynamics average of the strongest Sₐ → S₇ absorption. In order to obtain a sampling of representative conformations of both complexes, we performed 1000 ns long classical MD simulations for the prepared systems of OCPO and RCP. Previous studies have applied especially developed force fields to describe carotenoid dynamics, allowing for a direct calculation of QM-derived electronic properties from MD snapshots.27–29 In our approach, we maintained the general parametrization of the OPLS3e force field, but we locally minimized only the carotenoid for each snapshot under a QM/MM treatment. Notwithstanding, the most recent version of OPLSA (OPLS3e) incorporates an extended torsional parametrization as well as optimized partial charge assignment.20 Taking into account only the polyene chain, the in-place RMSD before and after QM/MM minimization is, on average, 0.28 Å. Thus, the local QM/MM minimization produces very small displacements within the protein cavity.

The MD of OCPO shows a stable RMSD for the protein (∼1.5 Å) and the ligand (∼1 Å) during the entire simulation (Figure S1). The most frequent protein–ligand interactions are the hydrogen bonds with Tyr201 and Trp288, as well as hydrophobic contacts located mainly on the NTD side (Tyr44, Trp110) (Figure S3). On the other hand, in RCP, the RMSD stabilizes around 2.5 Å for both the ligand and the protein (Figure S2). In this case, the carotenoid interacts mostly through hydrophobic contacts between residues of the NTD cavity and the isoprenoid chain, while both rings are exposed to solvent (Figure S4), which explains the enhanced motional...
freedom of the protein and ligand. As expected from the visual inspection of both structures, the ligand SASA is significantly higher in RCP than in OCPO (Figure S5). From the resulting MD trajectories, we exported 51 snapshots evenly distributed along the simulation time (every 20 ns) for each model. After the deletion of solvent and ions, the structures were optimized using QSite, with the carotenoid included in the QM part of the system, and with the protein frozen. Subsequently, single points were run on each optimized snapshot to obtain excitation energies. Remarkably, despite the variability observed, the confidence intervals are within 2% of the computed average and reproduce the experimental values reported in the literature (Figure 2, Table 1). These results help validate our computational methodology.

The degree of conjugation of a carotenoid's isoprenoid chain is known to have a strong influence on its electronic properties and excitation energies. Bond length alternation (BLA) is a parameter that describes such conjugation. In addition, the torsions connecting the isoprenoid chain with the end-rings indirectly affect conjugation. Our results confirm the direct correlation between the BLA and the calculated λ_max for both models, showing a linear correlation with R^2 = 0.85 for OCPO^3, and R^2 = 0.95 for RCP (Figure S6). Dihedrals on both end-rings (Figure 3C) have a higher conformational freedom in RCP, where both ends of the carotenoid are exposed to solvent, while in OCPO the protein environment restricts those torsions affecting the electronic properties of the system and lowering the degree of conjugation. In our MD trajectory, RCP is mostly trapped in one of the free-energy minima showing average values of β_1-dihedral = 45° ± 7° and β_2-dihedral = 159° ± 27° for the optimized snapshots. This configuration corresponds to a nominal s-cis for β_1 and s-trans for β_2 (as observed in the crystal structure). The large deviations observed for the β_2-dihedral are caused by a small fraction of states (~6%) exploring the s-cis and ~90° configurations. In particular, it has been previously demonstrated that the combination of an s-cis β_1-dihedral with an s-trans β_2-dihedral represents only one of several possible

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**Table 1. QM Calculations on Optimized Structures from MD for OCPO and RCP Models**

<table>
<thead>
<tr>
<th>model</th>
<th>λ_max (nm)</th>
<th>confidence interval (95%)</th>
<th>SD</th>
<th>BLA</th>
<th>expt λ_max (nm)</th>
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<td>OCPO-CAN</td>
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<td>474.5 483.0</td>
<td>15.1</td>
<td>0.073 ± 0.004</td>
<td>477.0</td>
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<tr>
<td>RCP-CAN</td>
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<td>519.0 532.0</td>
<td>23.0</td>
<td>0.056 ± 0.007</td>
<td>525.0</td>
</tr>
</tbody>
</table>

*Results are expressed as the mean of 51 snapshots. A one-sample t test was used to obtain the confidence intervals. Bond length alternation (BLA) is expressed as an average ± standard deviation.*

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**Figure 2.** Structures of OCPO-CAN (A) and RCP-CAN (B). The lower panels show the results of the calculated λ_max for each locally optimized snapshot extracted from the MD. Confidence intervals (95%) for both models are shown with orange and red shaded areas, respectively. Dashed lines indicate the experimental λ_max reported in the literature for both structures.
thermodynamically accessible conformations that CAN adopts in RCP. On the other hand, our results for the OCP\(^{16}\) optimized snapshots from the MD show a predominant \(\text{s-cis}\) configuration for \(\beta2\)-dihedral \((-45^\circ \pm 5^\circ)\), together with a \(\beta1\)-dihedral showing a higher variation but with a more prevalent \(\text{s-trans}\) conformation \((112^\circ \pm 21^\circ)\). Previous anisotropy measurements by Konold et al. have also shown that the \(\beta1\)-dihedral assumes an \(\text{s-trans}\) conformation in OCP\(^{16}\). Interestingly, our MD simulation indicates that this nominally \(\text{s-trans}\) \(\beta1\)-dihedral is restricted by the protein environment at the CTD of OCP\(^{16}\) to a maximum angle of \(145^\circ\). This prevents the enhancement of conjugation in the chromophore, reflected in the orange coloration of the inactive state. Table S1 presents additional analysis of dihedrals and a comparison with simulations in solution.

**Metadynamics to Drive Carotenoid Translocation.** As pointed out in the Introduction, the photoactivation mechanism in OCP involves a 12 Å carotenoid translocation into the NTD, followed by domain separation. The results from our MD simulation, as well as inferences reported by other authors,\(^{16}\) clearly indicate that OCP\(^{16}\) is a thermodynamically stable complex requiring enhanced sampling methodologies to force the system away from this local energy minimum. Hence, we designed a metadynamics experiment with the aim to drive CAN translocation in OCP bypassing the creation of the photoproduct P1.

In metadynamics, a history-dependent bias potential is built during the simulation as a sum of repulsive Gaussians in the defined collective variable/s (CV) space. Moreover, WT-metadynamics adds a bias factor \((\gamma)\) that limits the exploration to thermodynamically relevant configurations.\(^{31}\) The careful selection of the CV is one of the most important considerations in metadynamics. It can include diverse geometrical descriptors, such as distances, angles, or dihedrals. In the case of OCP, the availability of two reference models representing initial (OCP\(^{0}\)) and final (RCP) states of the system facilitates the definition of the CV. In order to force the displacement of the carotenoid toward the NTD using OCP\(^{0}\) as the starting point, we needed to find a CV able to drive carotenoid translocation through the tunnel connecting the two domains. The goal was to apply a subtle force to steer CAN into the NTD. A close inspection of the carotenoid in RCP led us to select the CV of our system as the distance between the center of mass of 5 residues on the \(\beta2\)-side of the NTD (shown in green in Figure 3A), and the atom C6\(^{\prime}\) of CAN’s \(\beta2\)-ring (Figure 3C).

It is important to state that we made no attempt to obtain a converged free-energy profile for such a complex reaction path. We used metadynamics only to create reasonable structural models that mimic the intermediate and final states of the translocation process. Indeed, out of 20 repetitions of the experiment, varying the initial seed randomly, the carotenoid did not move from its initial pocket in seven cases. The 13 remaining experiments, for which the translocation of CAN toward the NTD was observed, shared common structural features representing different states along the translocation pathway. Only in two of these experiments did the carotenoid reach a full 12 Å displacement without showing high distortions of the polyene chain. The infrequent occurrence of reasonable paths for translocation suggests that the protein environment effectively limits the movement of the carotenoid. The analysis of characteristic structural features at different points of the simulation is reported here for one of these two paths. The other path, which leads to the same general conclusions, is presented in the Supporting Information (Figure S7). Both of these paths show a similar energy barrier \((\sim 10 \text{ kcal/mol})\) for the overall process of translocation (Figure S8).

Snapshots along the selected path were extracted from the resulting metadynamics trajectory every 1 ns. Each snapshot was optimized at the QM/MM level, using the same procedure described before for the MD snapshots, followed by the calculation of \(\lambda_{\text{max}}\). The upper graph in Figure 4 shows that, during the first 48 ns, the excitation energies fluctuate around values similar to those calculated for OCP\(^{16}\). The solid black line represents the moving average of 5 adjacent snapshots displayed to guide the eye. It is important to mention that metadynamics time does not correspond to the real kinetics of the process. We only use it here to reference different states of the system throughout the path. A significant rise of \(\lambda_{\text{max}}\) is observed at 49 ns, with average values reaching the interval of variation obtained for RCP. At this stage, the carotenoid has been displaced halfway between the starting position and the
expected final configuration, \( \sim 5–6 \) Å from the first snapshot considering C6 and C6’ as the reference atoms (Figure 4, lower panel). Between 60 and 80 ns, a marked blue shift of the absorbance maximum is observed. Finally, after 80 ns, the system reaches the full 12 Å translocation, with \( \lambda_{\text{max}} \) values reflecting a red-shifted OCP-like state.

Spectroscopic features of the carotenoid are essentially affected by the conjugation of the system. BLA is a good indicator of the conjugation, but in the case of \( \beta \)-carotene-like molecules, torsions of terminal rings further influence conjugation. In fact, rotation of the terminal rings of CAN around the single bonds C6–C7 (in \( \beta_1 \)) and C6’–C7’ (in \( \beta_2 \)) have been previously described as one of the main degrees of freedom modulating the excitation energy. Results of our metadynamics demonstrate that while the \( \beta_2 \)-dihedral fluctuates mainly within the absolute values \( 45–60^\circ \), RCP-like \( \lambda_{\text{max}} \) values are only observed when the \( \beta_1 \)-dihedral values are over \( \sim 150^\circ \) (s-trans). When either of these two dihedral angles is \( \sim 90^\circ \), the conjugation of the system decreases, showing blue-shifted states. As mentioned in the previous section, it is clear that the protein environment in different locations inside OCP allows different conformations of the rings, directly affecting the electronic structure of the chromophore.

The resulting WT-metadynamics trajectory can be divided in four main stages representing different locations of CAN in OCP. Two of these states present higher average \( \lambda_{\text{max}} \) values, typical of the OCP photoproducts (highlighted by the pink vertical stripes in the lower graphs of Figure 4). Representative snapshots of each state are shown in Figure S5, where the nearest residues (up to 4 Å) defining the ligand pocket are highlighted. The protein surface helps visualize the shape of the cavities as well as solvent accessibility areas in the proximities of CAN. Ligand SASAs at each stage of the metadynamics trajectory are included in the S1 (Table S3). The position represented by the snapshot at 52 ns shows that when the carotenoid is located halfway between initial and final states of the translocation, with the key hydrogen bonds already broken, the protein environment can easily accommodate the chromophore allowing a more planar configuration of the system. We propose this state to be a representative of the OCP(P2) photoprodut. On the contrary, at a distance of \( \sim 10 \) Å from the starting point (60–80 ns), most of the snapshots show a carotenoid highly distorted from planarity, as exemplified by the snapshot at 70 ns. At this point, isomerization (s-cis) of different bonds along the central torsions of the polyene chain is observed (Table S4) due to constraints by the local protein environment, causing a marked decrease in the conjugation of the system. We propose that such states represent the bottleneck of the translocation process. Finally, after 80 ns, the system reaches a location where, again, a more planar configuration of the carotenoid is allowed. At this point, the location of CAN within the cavity is very similar to that observed in the RCP crystal (Figure 6). The outstanding difference found between both models is related to the configuration of both end-ring torsions. Interestingly, as has been demonstrated by Mennucci et al., there are several thermodynamically stable conformations found for CAN in RCP besides the one observed in the crystal structure. In summary, we propose that states like these are models of the OCP(P3) photoprodut.

Red-Shifted Form of OCP\(^{10} \). The heterogeneity observed experimentally in the UV-vis absorbance spectra of OCP\(^{15} \) bound to different carotenoids, showing two bands of similar strength (at 477 and 500 nm for OCP-CAN), has been initially attributed to either the presence of OCP\(^{16} \) in the sample, or to the existence of an alternative form of OCP\(^{10,13} \). More recent experimental evidence supports the second hypothesis establishing the existence of a “red-shifted” form of OCP\(^{14,15} \). According to these authors, this form should be characterized by a looser binding pocket, with a higher degree of conjugation of the carotenoid’s isoprenoid chain. Despite these inferences, a detailed structural characterization of this “red-shifted” state has remained elusive. The results of our WT-metadynamics trajectory show a subpopulation of red-shifted snapshots distributed along the time range between 0 and 48 ns which are still bound to Tyr201 and Trp288 via hydrogen bonds. This observation leads us to propose this...
subpopulation of structures as the hypothetical “red-shifted” form of OCP$^0$. The snapshot at 17 ns (Figure 7) represents an example of this alternative form where the protein environment is reorganized enabling a more planar configuration of the conjugated chain when compared to the main form of OCP$^0$ (0 ns). As can be observed, the key hydrogen-bonding interactions with Tyr201 and Trp288 are kept unaltered in such a state. However, the occurrence of this “red-shifted” form involves an excursion into the NTD side allowing for the β2-end to stick out of the protein. This is consistent with the observation of the N-terminal domain being much more flexible than the C-terminal domain.

**CONCLUSIONS**

Using metadynamics, we have obtained insight into the process of carotenoid translocation in the photoactivation mechanism of OCP. We have proposed approximate molecular models for the two intermediate photoproducts (P2 and P3). To the best of our knowledge, this is the first study that simulates chromophore displacement and its relationship to absorption spectroscopy. The close correlation of the calculated values of the proposed intermediates with the UV−vis features reported from experimental studies gives a reasonable validation to our models. The finding of a single geometric variable capable of producing a plausible reaction path offers an important contribution for future characterization of OCP with various carotenoid chromophores. We have developed a computational methodology combining MD and QM/MM calculations which allowed us to reproduce the experimental $\lambda_{\text{max}}$ values of the reference models (OCP$^0$ and RCP). In addition, we were able to propose a model of a red-shifted form of OCP$^0$.
The OCP(P3) model shows a high resemblance to RCP and indirectly modulates the electronic properties. Previously inferred by spectroscopy experiments, although the primary event of OCP photoactivation (the formation of the P1 photoproduct) is still elusive, our work presents new insight into how the protein environment exerts distortions on the carotenoid and indirectly modulates the electronic properties. The OCP(P3) model shows a high resemblance to RCP and presents a useful starting point to explore the following step of the mechanism, protein domain separation. Current work in our laboratory is aiming to model this process.

Figure 7. Representative snapshot of the hypothetical “red-shifted” form of OCP<sup>0</sup> (at 17 ns). NTD residues are colored in orange, and CTD residues, in cyan. Tyr201 and Trp288 are highlighted with thicker sticks, and hydrogen-bonding interactions are indicated with magenta dashed lines. The conformation of CAN in the standard blue-shifted state (at 0 ns) is shown for comparison.

**REFERENCES**


