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BIOCHEMISTRY

Photoactivation of *Drosophila melanogaster* cryptochrome through sequential conformational transitions

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Cryptochromes are blue-light photoreceptor proteins, which provide input to circadian clocks. The cryptochrome from *Drosophila melanogaster* (*DmCry*) modulates the degradation of Timeless and itself. It is unclear how light absorption by the chromophore and the subsequent redox reactions trigger these events. Here, we use nano- to millisecond time-resolved x-ray solution scattering to reveal the light-activated conformational changes in *DmCry* and the related (6-4) photolyase. *DmCry* undergoes a series of structural changes, culminating in the release of the carboxyl-terminal tail (CTT). The photolyase has a simpler structural response. We find that the CTT release in *DmCry* depends on pH. Mutation of a conserved histidine, important for the biochemical activity of *DmCry*, does not affect transduction of the structural signal to the CTT. Instead, molecular dynamics simulations suggest that it stabilizes the CTT in the resting-state conformation. Our structural photocycle unravels the first molecular events of signal transduction in an animal cryptochrome.

INTRODUCTION

The cryptochrome protein family is widespread in plants and animals (1, 2). Cryptochromes are important for the entrainment of the circadian rhythm (3–5) and have also been suggested to be involved in magnetoreception (6–9). The blue-light-sensing flavoproteins stem from the DNA repair enzymes known as photolyases but have since evolved to blue-light photoreceptors with little or no DNA repair ability (10). Cryptochromes and photolyases have a noncovalently bound light- and redox-active flavin adenine dinucleotide (FAD) as an essential chromophore. The FAD can be reduced twice, and whereas fully reduced FAD is required for DNA repair in DNA photolyases, the singly reduced semiquinone FAD triggers photoreceptor activity in cryptochromes (10, 11). Structurally, photolyases and cryptochromes are highly similar, but cryptochromes contain additional amino acids at their C-terminal end (1). The length of this C-terminal tail (CTT) varies between different cryptochromes, from tens to several hundreds of amino acids (1, 2).

The fruit fly *Drosophila melanogaster* contains one type I insect cryptochrome (*DmCry*). This protein is the primary light receptor of the *D. melanogaster* circadian clock (3). In the organism, it coordinates interactions between Timeless (TIM) and the E3-ubiquitin ligase Jetlag (JET) in a light-dependent manner. The interaction between *DmCry*, TIM, and JET depends on conformational changes that occur at the CTT of *DmCry* (12). It appears that the CTT blocks the interaction surface in the dark, and deletion of the CTT abolishes the light dependence of these interactions (13, 14). In *DmCry*, the CTT is about 20 amino acids long, and in the dark, the CTT forms a 10-residue helix

that is bound to the photolyase homology domain, placing phenylalanine 534 of the conserved Phe-Phe-Trp (FFW) motif at the position where photolyases bind DNA (15, 16).

Blue-light absorption triggers FAD excitation and nanosecond electron transport along four tryptophan residues to generate a radical pair $\text{FAD}^{\bullet-} \cdots \text{W394}^{\bullet+}$ (Fig. 1) (17, 18). After a few microseconds, $\text{W394}^{\bullet+}$ is deprotonated (19), and on a millisecond time scale, the $\text{FAD}^{\bullet-} \cdots \text{W394}^{\bullet}$ radical pair recombines. If the intermediate W394^{\bullet} radical is reduced by an external electron donor before recombination, then the remaining flavin radical is reoxidized over the course of several minutes, depending on the concentration of electron acceptors such as oxygen. Although the anion radical ($\text{FAD}^{\bullet-}$) is the predominant photoproduct, some neutral radical (FADH^{\bullet}) is also formed (20).

Despite this knowledge about the redox reactions of the FAD chromophore, reliable information about protein structural changes is sparse. After continuous light excitation, limited proteolysis assays and small-angle x-ray scattering (SAXS) measurements have suggested that the CTT of *DmCry* undocks or gets unstructured to open up a binding site for TIM (21). It appears that the FFW motif is directly involved in this process. The molecular rearrangements that lead from light absorption to release of the CTT remain less clear. A focal point of interest is a conserved histidine (378 in *DmCry*), which is located between the isoalloxazine moiety of the chromophore and the CTT (Fig. 1) (16, 22). In (6-4) photolyases, the equivalent histidine has been shown to be crucial for DNA repair activity (23, 24) and light-induced conformational changes (25). A recent study of *DmCry* suggested that flavin reduction causes proton uptake at H378 and that the induced changes in the hydrogen bonding network result in the detachment of the CTT (20). All of this information was obtained at photoequilibrium. Hence, it is currently not even clear at which point during the photoreaction the signaling state confirmation is reached. To make progress in understanding the mechanism of signal transduction in cryptochromes, it is necessary to investigate the protein using techniques that can resolve structural changes as a function of time.

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Here, we use nano- to millisecond time-resolved x-ray solution scattering (TRXSS) (26–32) and molecular dynamics (MD) simulations to investigate the light-induced conformational changes in *DmCry* after light excitation. Both the wild-type protein and a histidine 378 to alanine (H378A) mutant were investigated. As control experiment, we also investigated the (6-4) photolyase from *Xenopus laevis* (*XIPho*). *XIPho* is structurally similar to *DmCry* and undergoes similar light-induced electron transfer reactions, but it lacks the CTT.

RESULTS

TRXSS reveals a series of light-induced conformational changes from nano- to milliseconds

We used TRXSS to probe the conformational changes during photoactivation of *DmCry*. In the experiment, the protein solution is excited by a nanosecond laser pulse, and the x-ray solution scattering is re-

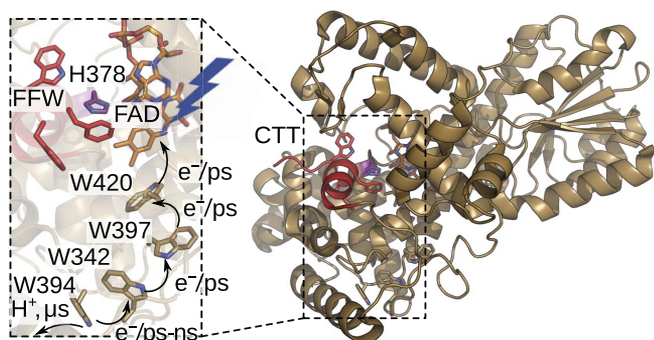


Fig. 1. Cryptochrome structure and photoreduction. The overall structure of *DmCry* [Protein Data Bank (PDB) ID: 4GU5]. The inset depicts an enlarged view of the area around the chromophore, H378, the FFW motif, and the tryptophan tetrad.

corded at a series of defined delay times. We recorded TRXSS data covering delay times from 10 ns to 10 ms for *DmCry*, from 3 μ s to 10 ms for *DmCry*(H378A), and from 100 ns to 10 ms for *XIPho* (Fig. 2). Difference scattering (light-dark) signals were observed for all three samples. The shapes of the signals for the two cryptochromes evolve over time with two changes of sign for the slope at $q < 0.1 \text{ \AA}^{-1}$ (q is the modulus of the scattering vector defined as $q = 4\pi\sin(\theta)/\lambda$, where 2θ is the scattering angle and λ is the x-ray wavelength) observed at 3 μ s and between 3 and 10 ms and the grow-in of a positive peak at $q = 1 \text{ \AA}^{-1}$ at approximately 1 ms. In contrast, the data for *XIPho* show only one signal at $q < 0.1 \text{ \AA}^{-1}$, which rises at 300 ns and decays between 1 and 3 ms.

We decomposed the difference scattering signal into time-independent species-associated difference scattering curves, assuming a sequence of transitions following first-order kinetics (Fig. 3 and Table 1). Five species were required to model the data for *DmCry*. Fewer intermediates resulted in systematic deviations between the model and the data, whereas additional components resulted in unreasonable values for the refined rate constants or did not notably improve the fit (fig. S1 and table S1). The five species will be referred to as *DmCry* $_{\alpha}$, *DmCry* $_{\beta}$, *DmCry* $_{\gamma}$, *DmCry* $_{\delta}$, and *DmCry* $_{\epsilon}$. The data for *DmCry*(H378A) could be modeled using four species. The scattering profiles of the four intermediates closely resembled *DmCry* $_{\beta}$ to *DmCry* $_{\epsilon}$ of the wild-type cryptochrome (Fig. 3, E and F). The difference scattering of the β intermediate of *DmCry*(H378A) may be convoluted with scattering for an α state, which is not resolved because of the limited time range probed. The time-resolved x-ray scattering of *XIPho* was well described by the rise and the decay of a single intermediate state.

Note that the transient difference scattering signal observed for *XIPho* closely resembles one of the early *DmCry* states, namely, *DmCry* $_{\beta}$ (Fig. 3, E and G). Since the two proteins are structurally comparable and undergo similar electron transfer reactions after illumination (1),

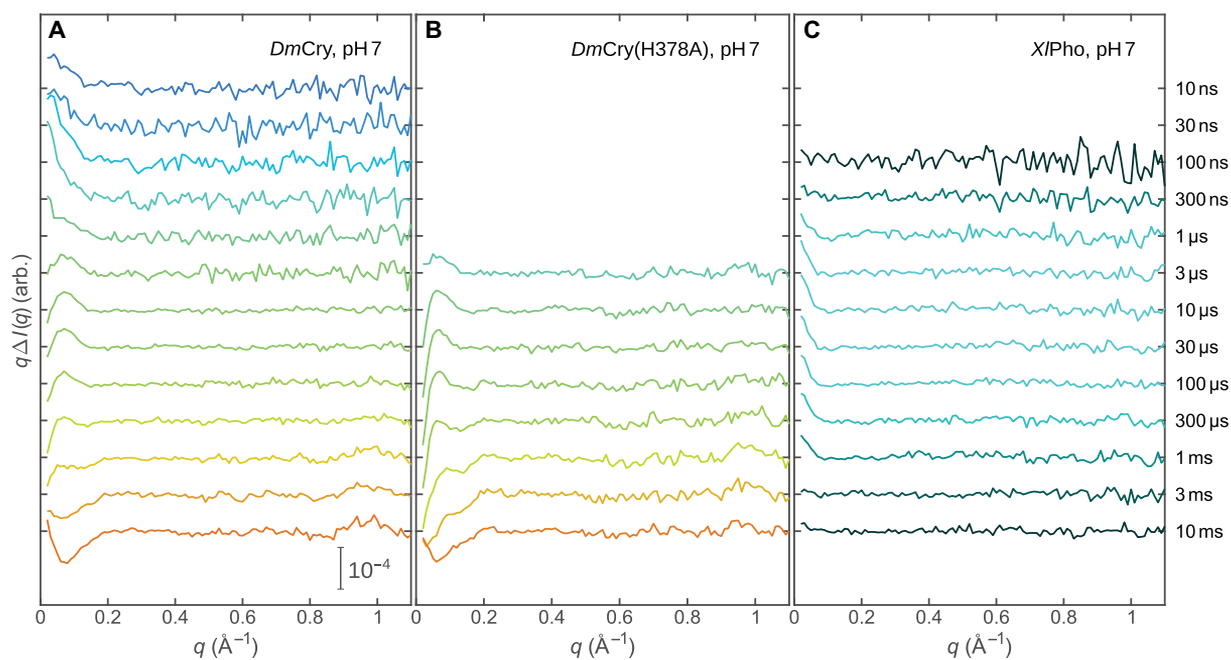


Fig. 2. TRXSS data. TRXSS data covering delay times from 10 ns to 10 ms for wild-type *DmCry* (A), for the *DmCry*(H378A) mutant (B), and for *XIPho* (C). Coloring follows the transient population of the different intermediates shown in Fig. 3. Different delay time ranges were probed for the samples because of restrictions in beamtime and available sample volumes.

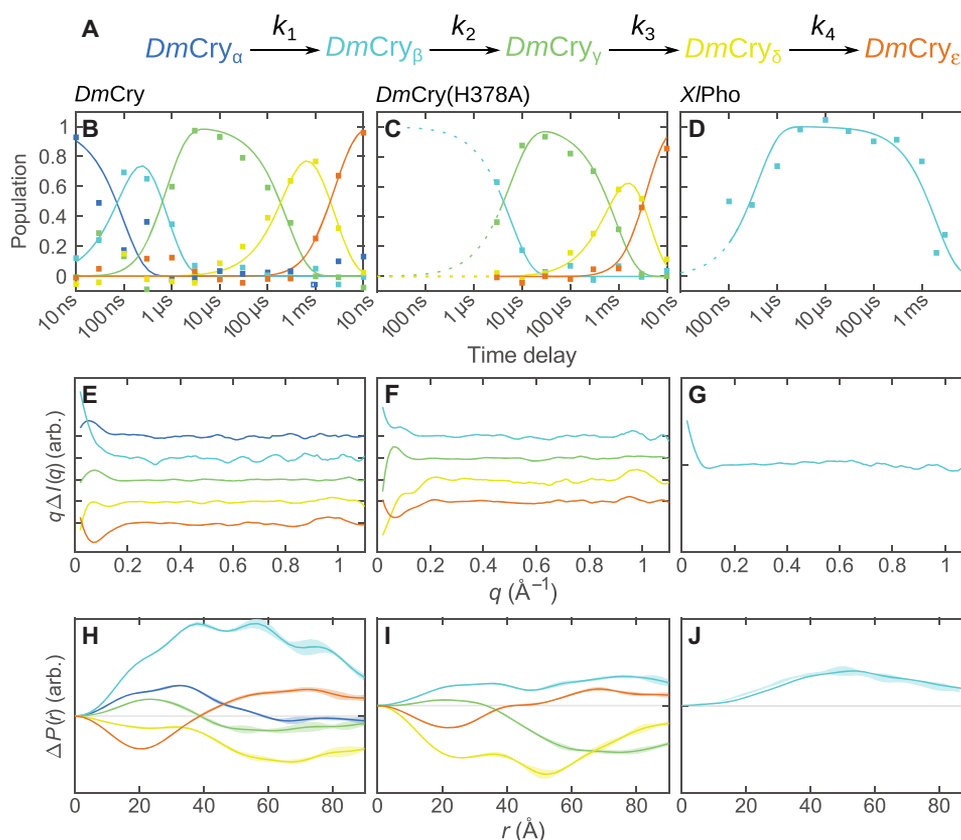


Fig. 3. Kinetic analysis of TRXSS data. (A) The kinetic model used for the analysis of the TRXSS data. Time-dependent populations of the different structural intermediates are shown in (B) to (D) for *DmCry* wild type (B), *DmCry*(H378A) (C), and XIPho (D). The fitted model is shown as lines. The squares represent the optimized populations obtained by fitting the experimental difference scattering curves with the species-associated difference scattering curves in (E) to (G). The model outside the probed time region is shown as dotted lines. Species-associated difference scattering curves for *DmCry* (E), *DmCry*(H378A) (F), and XIPho (G). Change in the pair distance distribution $[P(r)]$ for *DmCry* (H), *DmCry*(H378A) (I), and XIPho (J). The shaded area represents ± 1 SD. The first intermediate state of *DmCry*(H378A) is not well resolved because of the limited time range probed and appears to be a mixture of *DmCry* $_\alpha$ and *DmCry* $_\beta$.

Table 1. Rate constants for structural transitions by TRXSS.

TRXSS (pH 7)	k_1 (μs^{-1})	k_2 (μs^{-1})	k_3 (ms^{-1})	k_4 (ms^{-1})
<i>DmCry</i>	10.05 ± 0.64	1.30 ± 0.06	3.86 ± 0.30	0.40 ± 0.03
<i>DmCry</i> (H378A)	no data	0.17 ± 0.01	1.15 ± 0.07	0.30 ± 0.04

it is likely that the XIPho intermediate and *DmCry* $_\beta$ represent analogous structural changes.

TRXSS measures the change in distances between the scattering electrons in the sample. Generally, large-scale conformational rearrangements, such as domain movement, partial unfolding, mass transit, or changes in the hydration layer, are observed at low q ($q < 0.1 \text{ \AA}^{-1}$) (26, 29, 31, 33, 34). Clear difference signals can be observed at low q for all delay times. We estimated the change of the radius of gyration (ΔR_g ; Eq. 3) to be less than $\pm 0.2 \text{ \AA}$ for all states (table S2). This indicates concerted but small changes to the global protein structure. Signals in the intermediate q range ($0.2 < q < 1 \text{ \AA}^{-1}$) are associated with tertiary structural changes (26, 27, 30, 31). All three samples exhibit fairly flat difference scattering in the intermediate q range, thereby suggesting that

there is no substantial change of the tertiary structure of the protein. We observe a peak at $q = 1 \text{ \AA}^{-1}$ for *DmCry* and *DmCry*(H378A), which appears with the *DmCry* $_\delta$ state. The q range and the magnitude of the signal suggest that it originates from a coordinated change of interatomic distances around 6 \AA ($r = 2\pi/q$). Considering the pitch of α helices of 5.4 \AA , the signal would be consistent with structural changes of the partly α -helical CTT. This assignment is supported by the fact that the peak is absent in XIPho, which lacks the CTT.

To gain more structural insight, we computed the difference pair distance distribution $[\Delta P(r)]$ from the difference scattering curves $[\Delta I(q)]$ using a sine Fourier transform (Eq. 2). The $\Delta P(r)$ curve corresponding to *DmCry* $_\gamma$ shows a decrease of long ($>40 \text{ \AA}$) and an increase of short distances, which suggest a compacted state. *DmCry* $_\epsilon$ has more long and less short distances, characteristics of an elongated structure. This is also confirmed by ΔR_g , which we calculated to be $+0.09$ and $+0.07 \text{ \AA}$ for *DmCry* $_\epsilon$ and *DmCry*(H378A) $_\epsilon$, respectively (table S2). We conclude that *DmCry* $_\epsilon$ has an extended conformation with a detached CTT (21).

The $\Delta P(r)$ of *DmCry* $_\alpha$, *DmCry* $_\beta$, and *DmCry* $_\delta$ is positive or negative at all distances. This cannot be explained by a structural change solely within the protein, where an increase in long distances would always be accompanied by a corresponding decrease in short distances and vice versa. The protein, however, is surrounded by a solvent layer,

which has a different density to that of the bulk solvent. The density of this layer could increase or decrease, leading to all-positive or all-negative $\Delta P(r)$ functions.

We confirmed that this is a reasonable explanation by calculating the x-ray scattering of the *DmCry* crystal structure (22) with *CRY SOL* (35). We varied the contrast of the hydration layer corresponding to addition or removal of 20 water molecules, and we referenced the data to scattering computed for the default value of $0.03 \text{ e}^-/\text{\AA}^3$ (Fig. 4B). The resulting $P(r)$ curves reveal an increase or a decrease in all distances (Fig. 4C). We therefore conclude that there is a significant contribution of such solvent layer remodeling for *DmCry*_α, *DmCry*_β, and *DmCry*_δ. Considering that substantial charge redistribution occurs over the course of the photoreaction, changes in the ion cloud (36) or in the dynamics of the surface water seem reasonable. Overall, the structural analysis indicates that the tertiary structure of the *DmCry* is stable over the course of the photoreaction but that a number of local and small structural changes lead to formation of the final conformational state *DmCry*_ε.

The formation of the signaling state is pH dependent

To find out which of the states corresponds to the signaling conformation, we conducted a regular SAXS experiment in which *DmCry*

was kept either in the dark or under continuous blue-light exposure (fig. S2 and table S3). The results are in agreement with previously published SAXS data from dark- and light-adapted *DmCry* (21). The difference between our “light” and “dark” SAXS curves [$\Delta I(q)$] is highly similar to the scattering pattern detected for the last state, *DmCry*_ε (Fig. 4A). We therefore consider this state, which is reached within 10 ms of photoexcitation, to be the signaling state.

In the previous SAXS study, the light state was associated with a detached CTT (21). Our data are in agreement with this assignment. An extended conformation in *DmCry*_ε is indicated by the increase in R_g (table S2) and by the $\Delta P(r)$ function (Fig. 3, H and I). Moreover, the peak at $q = 1 \text{ \AA}^{-1}$ in *DmCry*_δ and *DmCry*_ε and the absence of a scattering signal in *XlPho* equivalent to that of *DmCry*_ε are consistent with conformational changes at the CTT (Fig. 3, E and F).

To further investigate the signal transduction pathway, we recorded TRXSS data of *DmCry* and *DmCry*(H378A) in a more basic solution with pH 9 (fig. S4, A and B). We decomposed the data using a sequential model as described for the measurements at pH 7 (table S4). By comparison of TRXSS patterns and $\Delta P(r)$ functions of the intermediates at pH 9 to those obtained at pH 7, we identified states at pH 9, which were equivalent to *DmCry*_β, *DmCry*_γ, and *DmCry*_δ at pH 7. We could not identify an equivalent to *DmCry*_α possibly because of altered fast kinetics at pH 9, which could shift the appearance of this intermediate outside the experimental time window.

An equivalent of the final conformational state *DmCry*_ε was not observed at pH 9. The final x-ray scattering pattern at pH 9 contains the positive peak at $q = 1 \text{ \AA}^{-1}$, but the signals at $q < 0.2 \text{ \AA}^{-1}$ differed from those at pH 7. The $\Delta P(r)$ of the final states (Fig. 3, H and I, and fig. S4, G and H) reveals that the positive feature at long distances, indicating an extension of the protein, is absent at pH 9. Because of technical limitations, we could only probe delay times up to 10 ms, and it is possible that an equivalent to *DmCry*_ε is developed outside the probed time window at pH 9. In either case, the data show that the formation of the signaling state *DmCry*_ε depends on pH and requires uptake of a proton.

The role of H378 in *DmCry* signal transduction

Placed directly between the isoalloxazine moiety of the chromophore and the CTT, the conserved H378 has been implicated in pH-dependent signal transduction between the chromophore and the CTT (20, 23, 24). We therefore recorded the TRXSS data on *DmCry*(H378A), but to our surprise, we found that the TRXSS patterns and kinetics in *DmCry* and *DmCry*(H378A) are highly similar (Fig. 3, E and F, and Table 1). The $\Delta P(r)$ curves for *DmCry*(H378A)_{γ-ε} show this most clearly, as they exhibit the same patterns of positive and negative signals as for *DmCry* (Fig. 3, H and I). We note that the states α and β are not resolved for *DmCry*(H378A) because of the limited time range of the data. In both samples, the signaling state is reached within 10 ms. This confirms that the H378A mutation hardly alters the structural changes after blue-light photoexcitation. Moreover, our TRXSS data show that even at pH 9, *DmCry* and *DmCry*(H378A) display highly similar structural rearrangements (fig. S4). Thus, we conclude that protonation reactions of H378 are not required for the pH-dependent formation of the signaling state.

To investigate the role of H378 in *DmCry* in greater detail, we performed classical MD simulations of the H378A mutant and the wild-type protein, with the FAD modeled with parameters for the oxidized (FAD) and semireduced (FAD^{•-}) states. H378 was protonated at the N^e (N3) position as suggested by the crystal structures (16, 22) and

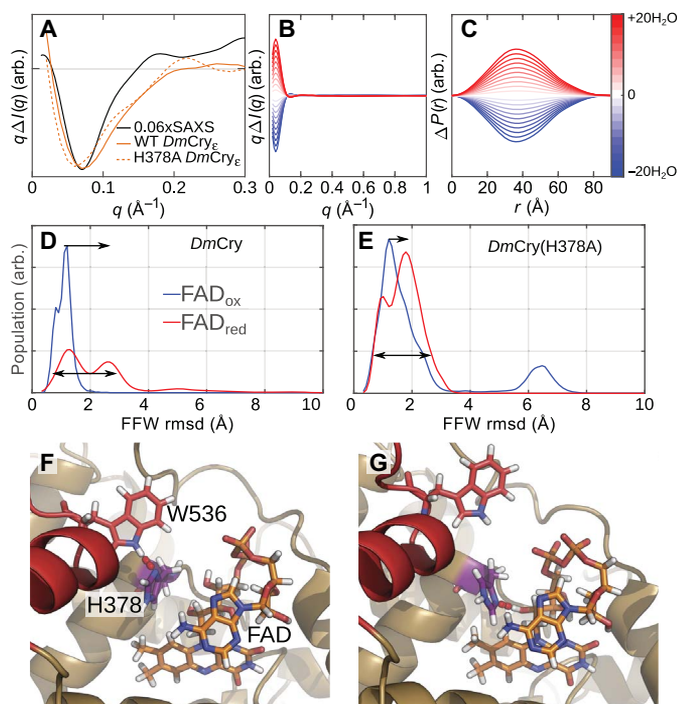


Fig. 4. Steady-state difference SAXS, modeling of hydration layer, FFW flexibility in MD simulations, and active-site hydrogen bonding network. (A) Difference scattering from a steady-state SAXS measurement compared to *DmCry*_ε. Note that it is the regularized difference intensity that is shown for the SAXS measurement. Predicted difference scattering for *DmCry* crystal structure (PDB ID: 4GU5) by only varying the hydration layer contrast by the equivalent of a few water molecules (B) and the corresponding change in $P(r)$ (C). Simulations with the oxidized chromophore parameters display overall lower FFW RMSD compared to simulations with the reduced chromophore parameters (D and E). Arrows indicate shift and widening of the RMSD distribution when going from an oxidized to reduced chromophore. When the chromophore is oxidized, H378 preferentially forms a hydrogen bond with W536 (F), but when the FAD becomes reduced, H378 preferentially binds to the FAD (G). arb., arbitrary unit.

Ganguly *et al.* (20). Four trajectories, each 250 ns long, were generated for each situation and will be referred to as WTFAD_{ox}1-4, WTFAD_{red}1-4, H378AFAD_{ox}1-4, and H378AFAD_{red}1-4. The mobility of the CTT was evaluated by the root mean square deviation (RMSD) of the FFW motif with respect to the crystal structure. As in (20), we removed rotational and translational degrees of freedom before calculating the RMSD by aligning the protein backbone on the quartile of residues that exhibited the lowest root mean square fluctuations (RMSFs).

For the wild-type protein, we find that the mobility of the FFW is elevated when FAD is reduced compared to when it is oxidized (compare the blue to the red line in Fig. 4D and fig. S3, A and B). This trend has been observed previously (20). We also find that the hydrogen bonding network between the FAD, H378, and W536 changes in response to the additional charge on the FAD. When FAD is oxidized, H378 predominantly hydrogen-bonds to W536 of the CTT (Fig. 4F and fig. S3E), and when FAD is reduced, the N^δ of H378 bonds to the ribose hydroxyl (OS2) of the chromophore (Fig. 4G and fig. S3E). We note that the described trends are observed in most trajectories but that WTFAD_{red}3 is an exception as H378 forms a hydrogen bond to W536, as in the oxidized trajectories. Also, the CTT completely detaches in WTFAD_{red}4 after 60 ns, and no hydrogen bond is observed (fig. S3, B and E).

The observations from the *DmCry* trajectories could be seen as an indication that H378 is actively involved in signal transduction. However, when we repeated the simulation after mutation of H378 into an alanine, we observed an upshift of the FFW-RMSD distribution, similar to the trend observed for the wild-type protein (compare the blue line and the red line in Fig. 4E and fig. S3, C and D). This underpins our conclusions from the TRXSS data and suggests that H378 is not the sole critical factor for relaying the change of redox state of the FAD chromophore to the CTT.

When comparing the mobility of *DmCry* and *DmCry*(H378A) with FAD in the oxidized (resting) state, we find that the peak of the FFW RMSD distribution remains unchanged at 1.2 Å between the samples but that the distribution is wider in the case of *DmCry*(H378A) (compare blue lines in Fig. 4, D and E). For one of the mutant simulations (H378AFAD_{ox}1), we observe pronounced CTT detachment, manifesting as a secondary peak around 6 Å RMSD. Together, these results suggest that the main role of H378 is to reduce the dynamics of the CTT when FAD is in the resting (oxidized) configuration. It achieves this by anchoring the CTT to the photolyase homology domain. Tight attachment of the CTT may be critical *in vivo* as it would prevent the cryptochrome from leaking signals in the dark.

DISCUSSION

Here, we show (i) that blue-light absorption triggers a series of conformational rearrangements in *DmCry*, resulting in the formation of the signaling state and detachment of the CTT on a millisecond time scale, (ii) that the last transition into the signaling state depends on pH, and (iii) that the conserved H378 is not required for signal transduction and is not the sole cause of the pH dependence. Instead, we propose that it stabilizes the CTT in the resting-state conformation.

Structural changes are already observed within 10 ns of excitation, and the protein undergoes a sequence of conformational rearrangements before reaching the signaling conformation after a few milliseconds. The protein retains its overall globular shape during the photoreaction, and changes in the water or ion density around the

protein occur. The resulting photoreaction scheme of *DmCry* contains more structural than spectroscopic intermediates (Fig. 5A). This is not surprising, considering the fact that ultraviolet-visible (UV-vis) spectroscopy reports selectively on the redox state and the immediate surroundings of the FAD chromophore but that TRXSS is sensitive to structural changes in the entire protein as well as the surrounding hydration layer. Structural intermediates have also been identified in a plant cryptochrome by step-scan infrared spectroscopy (37), which also probes structural changes in the entire protein.

Comparing kinetics from TRXSS and transient optical spectroscopy [see Fig. 5A and (17, 19)] reveals that formation of the FAD^{•-}...W394⁺ radical pair triggers the start of conformational changes of the protein. However, the rate constant for deprotonation of the tryptophanyl cation radical does not seem to correlate directly with the TRXSS kinetics (table S5). The rate constant for recombination is slightly lower than the formation rate of the signaling state as measured by TRXSS (Table 1 and table S4; see Materials and Methods for a detailed discussion). This supports our view that the protein structural changes, which are probed by TRXSS, are caused by changes in the oxidation state of FAD but that they otherwise are kinetically uncoupled from the reactions of the radical pair.

Our data support the notion that the signaling state *DmCry_s* has a detached CTT (12, 16, 21). The transition into the final state is slowed down or prevented at elevated pH (fig. S4 and table S4). We therefore conclude that a proton is taken up by the protein to facilitate this transition (Fig. 5). Currently, we are not able to identify the residue at which the protonation reaction occurs. H378 is located directly between the chromophore and CTT and has been proposed to undergo a protonation reaction during photoactivation of *DmCry* (20). However, this proposal is not consistent with our TRXSS data as the H378A mutation does not alter the formation of the signaling state at two different pH values. If H378 would be the primary residue responsible for the pH dependence, then we should have observed a change in rate constant for the formation of the signaling state upon removal of the residue by mutation. Alternative candidates are the residues S526 and

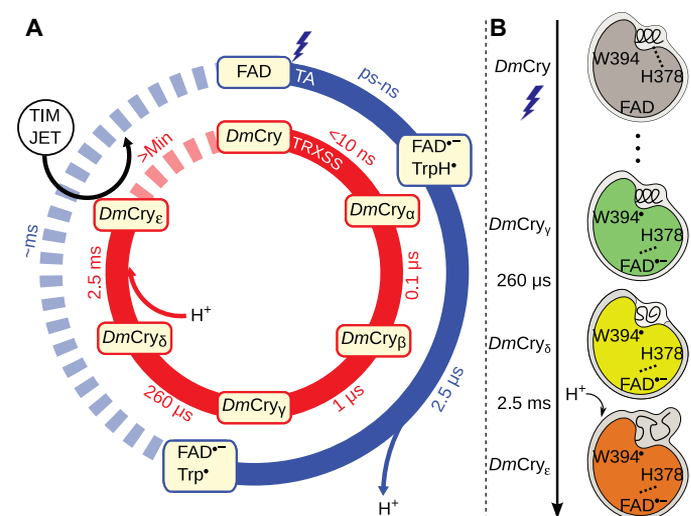


Fig. 5. Structural photocycle for *DmCry*. (A) Proposed structural photocycle for *DmCry* based on TRXSS data, shown as the inner cycle in red. The outer cycle in blue is the photochemical photocycle as determined by transient absorption (TA) spectroscopy. (B) The proposed structural rearrangements associated with the *DmCry₀* to *DmCry_e* states.

E530, which coordinate a network of hydrogen bonds, important for structural activation of *DmCry* (16). It is possible that this network may be disrupted by the protonation of one of the involved residues. To investigate this issue further, it would be good to conduct experiments, which can probe the protonation state of H378 and other residues directly.

As an alternative role for H378, we conclude from our MD simulations that H378 stabilizes the CTT in the resting state. This result is in agreement with activity assays in cell cultures of different H378 mutants of *DmCry*, from which similar conclusions were drawn (20). Thus, in the dark, the CTT is docked to the photolyase homology domain through hydrogen bonds between H378 and the CTT (Fig. 4, D to G). Stabilization of the CTT in the resting state may ensure that *DmCry* does not leak any signal in the dark, a feature that may be as important as efficient photoactivation. In the circadian clock in *D. melanogaster*, low levels of the TIM:PER (Period) complex have to be able to effectively block the expression of themselves in the cell nucleus at late night (3). If *DmCry* would have a residual dark activity, then it would mark TIM for degradation in the dark, hindering the clock cycle.

So how does the signal relay from the chromophore to the CTT? We consider the possibility that CTT detachment is caused by long-range Coulomb interactions between the semireduced $\text{FAD}^{\cdot-}$ and residues in the CTT. This is supported by the similarity of the structural changes that we detected when comparing wild-type *DmCry* and the H378A mutant. If the interaction between the FAD and the CTT is direct and through space, then the type of amino acid residing in between can be expected to play a minor role for signal transduction. Another alternative is that the signal is transduced from the chromophore to the CTT by a collective action of several amino acids that reside between the two sites. The alternatives are not mutually exclusive, and both are consistent with the cellular activity assays of various H378 mutations of *DmCry*, where it was found that none of the mutants completely disabled the light-induced change in activity of the protein (20).

Adding to the current understanding of cryptochromes, our findings provide hallmarks of the conformational photoactivation of *DmCry* (Fig. 5B). H378 stabilizes the resting-state conformation by tightly binding the CTT to the photolyase homology domain (gray structure). When photoexcited by blue light, the $\text{FAD}^{\cdot-} \cdots \text{W394}^{\cdot}$ radical pair is formed within a few microseconds. Structurally, the protein is in the state *DmCry_γ* at this point of the reaction (green structure), which is slightly contracted compared to the resting state (Fig. 3E). After about 0.2 ms, the next state (*DmCry_δ*) is reached, and we propose that the CTT loosens up or becomes unstructured as a direct consequence of the charge on the FAD (yellow structure). The loosening up of the CTT is a prerequisite for protonation of a yet unknown residue. This protonation reaction causes the CTT to completely detach from the N-terminal part of the protein (orange structure). The signaling conformation (*DmCry_ε*) is formed, primed for binding of TIM.

MATERIALS AND METHODS

Protein production and purification

The N-terminally His-tagged *DmCry* was produced in *Escherichia coli* BL21 cells following published procedures (19). *E. coli* cells were grown overnight at 303 K in the dark, and the protein synthesis was induced at an A_{600} (absorbance at 600 nm) of 0.4 to 0.6 by using 1 mM isopropyl- β -D-thiogalactopyranoside. After cell disruption, the lysate in buffer A

[50 mM Hepes (pH 7.0), 100 mM NaCl, and 10% (v/v) glycerol] was initially purified on a nickel-Sepharose column using binding buffer [50 mM Hepes (pH 7.0), 200 mM NaCl, 10% (v/v) glycerol, and 20 mM imidazole] and elution buffer [50 mM Hepes (pH 7.0), 200 mM NaCl, 20% (v/v) glycerol, and 500 mM imidazole]. Additional purification steps were carried out using gel filtration chromatography with buffer [50 mM Hepes (pH 7.0), 100 mM NaCl, and 20% (v/v) glycerol] and anion exchange chromatography using binding buffer [50 mM Hepes (pH 7.0), 50 mM NaCl, and 10% (v/v) glycerol] and elution buffer [50 mM Hepes (pH 7.0), 500 mM NaCl, and 10% (v/v) glycerol]. Yellow-colored *DmCry*-containing fractions were pooled and subjected to desalting column and concentration with microconcentrators. The histidine-to-alanine mutation was performed using the FastCloning method (38) with two overlapping primers in combination with a high-fidelity polymerase (KAPA HiFi/PEQLAB Biotechnologie) and verified by sequencing. The pET28a vector containing the mutated gene was transformed into *E. coli* SoluBL21 cells. *DmCry*(H378A) was produced in the same way as the wild-type protein. *XlPho* was produced using published procedures (39). A buffer consisting of 50 mM Hepes (pH 7.0 or pH 9.0), 100 mM NaCl, 5 mM ferricyanide, and 10% (v/v) glycerol was used in TRXSS experiments. The same buffer was used in the transient absorption experiments, and the ferricyanide concentration was reduced to 1 mM to account for the fivefold reduced protein concentration. In the conventional SAXS measurements, a buffer at pH 7 without ferricyanide and with a glycerol concentration of 20% (v/v) was used. Before any experiment, samples were filtered through a 0.2- μm filter.

Nonreducing SDS–polyacrylamide gel electrophoresis

An SDS–polyacrylamide gel electrophoresis (SDS-PAGE) under non-reducing conditions (no β -mercaptoethanol was added to the loading buffer) was performed to exclude the presence of any multimerization reactions due to disulfide linkages in any of the experiments. Both *DmCry* and *DmCry*(H378A) were investigated, and two parameters were varied: Samples were either kept in 50 mM Hepes (pH 7.0), 100 mM NaCl, and 10% (v/v) glycerol or, in the same buffer, supplemented with a five-times molar excess of ferricyanide. In addition, both proteins were kept either in the dark or were loaded on the gel directly after 5 min of blue light illumination using a lamp power of 9.7 mWcm^{-2} . As can be seen in fig. S5A, all band patterns were highly similar, and neither ferricyanide nor blue light illumination resulted in the formation of *DmCry* dimers or other degradation products.

Transient absorption measurements

Transient absorption measurements were performed as described in (19). In detail, time-resolved optical absorption spectroscopy was performed at 277 K with a commercially available laser flash photolysis spectrometer (LP920K, Edinburgh Instruments Ltd., Kirkton Campus, UK), and data were recorded with a digital oscilloscope (TDS-3012C, Tektronix, Beaverton, OR). The protein sample was placed in a synthetic quartz (Suprasil) semimicro cell (108F-QS, Hellma, Muellheim, Germany). The temperature was regulated to 277 ± 1 K by a temperature controller (Alpha RA 8, LAUDA, Lauda-Koenigshofen, Germany). Optical excitation was carried out using an OPO system (OPO PLUS, Continuum, San Jose, CA) pumped with a Nd:YAG laser (Surelite I, Continuum) at a wavelength of 460 nm, a pulse width of 6 ns, and a pulse energy of 4.0 ± 0.2 mJ. The repetition rate of the spectrometer was set to 6.67 MHz. To account for background signals, transients were measured alternately with and without optical excitation and used for

calculation of difference absorbance spectra with Beer-Lambert's law. We recorded two datasets per sample, one on a microsecond time scale to probe the deprotonation of the terminal tryptophan and the other on a millisecond time scale to probe the recombination of the radical pair.

Conventional SAXS data collection

SAXS data were collected at BM29 at the European Synchrotron Radiation facility at protein concentrations of 0.5, 1, 2, and 4 mg/ml. The measurements were performed at 10°C. The x-ray energy was 12.5 keV, and the detector used was Pilatus 1M. For each concentration, 10 images with 1-s x-ray exposure were collected with 10 images of the buffer before and after. Two sets of measurements were performed, one with the chromophore in the oxidized state and the other with the chromophore in the reduced anion radical state; this was achieved by illumination with 455-nm light and verified by UV-vis spectroscopy. The dark measurements were performed under red light, and the light-state measurements were performed under blue light. The data were averaged and subtracted using software at the beamline. Low-concentration (0.5 mg/ml) and high-concentration (4 mg/ml) data were merged at $0.05 < q < 0.10 \text{ \AA}^{-1}$. Additional data processing was performed using the ATSAS package (40), and the molecular weight was estimated as described by Rambo and Tainer (41).

TRXSS data collection

Time-resolved scattering experiments were performed at the BioCARS beamline at the Advanced Photon Source (APS). All measurements were performed at ambient temperatures (ca. 22°C). The protein concentration was 24 to 30 mg/ml (0.4 to 0.6 mM), and the buffer was supplemented with 5 mM potassium ferricyanide (ca. 10 times molar excess) to ensure that the chromophore is reoxidized during the experiment. With transient optical absorption spectroscopy, we established that the millisecond recombination of the radical pair did not change significantly by adding ferricyanide (table S5). To evaluate whether the high concentrations used for the TRXSS measurements distorted the difference scattering curve, the packing structure factor $S(q)$ (see fig. S5B) was estimated on the basis of a dilution series (24, 12, and 6 mg/ml and extrapolated to infinite dilution). No significant variation of this was observed ($\leq \pm 5\%$ at each point). We used the pink beam profile with a nominal x-ray energy of 12 keV, and the x-rays were focused to a ca. 50 μm by 50 μm spot. The sample was excited using a 475-nm 7-ns laser pulse with an energy density of 2 mJ/mm²; the laser spot size was adjusted so that it engulfed the x-ray spot. The x-rays and laser were overlapped on a 1-mm-diameter quartz capillary, holding the sample. The protein was delivered through inert polyether ether ketone tubing using a peristaltic pump at a constant pump speed of 2 $\mu\text{l/s}$. To collect data at different time points after excitation, the temporal offset between laser and x-rays was adjusted. Every fourth image recorded corresponded to a negative time delay (the laser pulse arrives after the x-ray pulse). The fingerprint of solvent heating was recorded by adding azo dye (0.5 mg/ml; Fast Yellow, Sigma-Aldrich) to the buffer solution. This dye absorbs the same wavelength as the protein sample, but most of the energy is transferred to the solvent as heat. This experiment was performed in the same way as just described. Detector images were radially integrated using our own software and normalized to the average scattering in the region $1.97 < q < 2.17 \text{ \AA}^{-1}$, where water scattering displays an isosbestic point with respect to heating. Absolute scattering data deviating from the median by more than 3 SDs and difference data deviating more than 2 SDs from the median in the region $2 < q < 2.5 \text{ \AA}^{-1}$ and $1.5 < q < 2.1 \text{ \AA}^{-1}$, respectively, were

regarded as outliers and were removed. In total, more than 90% of the images were kept. To generate difference scattering curves, the data of the closest (before and after) negative time delay were averaged and subtracted from each scattering curve. Difference scattering curves corresponding to the same x-ray laser delay time were averaged together. The contribution of the solvent heating was subtracted by scaling the pure heating curve to each time delay in the range $1.7 < q < 3 \text{ \AA}^{-1}$.

Kinetic decomposition and basic structural modeling of the scattering data

The x-ray solution difference scattering data were considered to be described as a linear combination of several time-independent species-associated difference scattering curves (Fig. 3 and fig. S1). A sequence of transitions described by first-order reactions was used to model the time-dependent concentration of the different intermediate states. The general case for five components (A to E) is described by Eqs. 1A to 1E. The species-associated difference scattering curves were retrieved by solving the system of linear equations $C \cdot SADS = \Delta I$, where C are the time-dependent concentrations and $SADS$ are the species-associated difference scattering curves. For wild-type *DmCry* at pH 7, five species were needed, and at pH 9, four states were needed. The measurements on the H378A mutant did not cover nanosecond time scales, and the data could be described using four or three states at pH 7 and 9, respectively. Fewer intermediates resulted in systematic deviations, and additional species did not improve the fit and sometimes resulted in unreasonable values for the rate constants (fig. S1). The rate constants of the model were refined by a least-squares minimization of the residual between the experimental and reconstructed data. The estimated error of the rate constants is retrieved from the Jacobian at the solution

$$\frac{d[A]}{dt} = -k_1[A] \quad (1A)$$

$$\frac{d[B]}{dt} = k_1[A] - k_2[B] \quad (1B)$$

$$\frac{d[C]}{dt} = k_2[B] - k_3[C] \quad (1C)$$

$$\frac{d[D]}{dt} = k_3[C] - k_4[D] \quad (1D)$$

$$\frac{d[E]}{dt} = k_4[D] \quad (1E)$$

By transforming the retrieved basis spectra to a difference in pair distance distribution $[\Delta P(r)]$, a structural evaluation of the different intermediates could be performed. The transformation is done according to Eq. 2

$$\Delta P(r) = \frac{r}{2\pi^2} \int_{q_{\min}}^{q_{\max}} q \Delta I(q) \sin(qr) \cdot e^{-q^2 \alpha^2} dq \quad (2)$$

$\Delta P(r)$ was evaluated from 0 to 90 \AA using difference scattering for $0.02 \leq q \leq 0.4 \text{ \AA}^{-1}$, and the regularization parameter α was set to 0.01. The statistical uncertainty of this transformation was estimated by

adding white Gaussian noise, with an amplitude similar to that observed in the raw experimental data, to the difference scattering curves associated with each species. One hundred curves with different noise were generated for each basis spectrum and used to estimate the SD of the reconstructed $\Delta P(r)$. The radius of gyration (R_g) for each state is calculated according to Eq. 3

$$R_g = \sqrt{\frac{\int_0^{r_{\max}} r^2 P(r) dr}{2 \int_0^{r_{\max}} P(r) dr}} \quad (3)$$

Here, $P(r)$ was computed by adding the $P(r)$ of the dark-adapted *DmCry* from a conventional SAXS measurement to the $\Delta P(r)$ for each state, accounting for the photoconversion yield of ca. 6%. The photoconversion yield was estimated by adjusting the absolute scattering curves from the TRXSS experiment to the same scale as the conventional SAXS experiment. The difference scattering of the *DmCry_e* state was then compared to the difference scattering (light-dark) of the conventional SAXS measurement, which, based on UV-vis measurements, is considered to represent complete conversion to the reduced anion radical form. The ΔR_g was calculated by subtracting the R_g of the dark-adapted state. The error of ΔR_g was estimated by calculating it using 100 $\Delta P(r)$ with simulated noise, as described above.

MD simulations

The starting structure used was based on the coordinates of a *D. melanogaster* cryptochrome crystal structure [Protein Data Bank (PDB) ID: 4GU5] (15, 22). The parameters for the oxidized and anion radical FAD were based on previous work (42). The simulations were run using GROMACS 5.0.4 (43) and the CHARMM27 force field. The simulation box was replicated in all directions with periodic boundary conditions, all bonds were constrained using the linear constraint solver algorithm, and a time step of 2 fs was used. Particle mesh Ewald (PME) electrostatics with fourth-order interpolation and with a grid spacing of 0.12 nm were used. The cutoff scheme was Verlet with a 1.0-nm cutoff, and cutoffs for short-range electrostatic and van der Waals interactions were 1.0 nm as well. During the production run, pressure control was achieved using the Parrinello-Rahman barostat ($\tau_p = 2$ ps, $P = 1$ bar), and temperature control was achieved via the modified Berendsen (velocity-rescale) thermostat ($\tau_T = 0.1$ ps, $T = 300$ K). The protein was placed in a cubic box, 1 nm larger than the protein in all directions and solvated with transferable interparticle potential with 3 points (TIP3P) water molecules. The system was neutralized by replacing water molecules with sodium and chloride ions to a final concentration of 150 mM. After adding ions, the system underwent initial energy minimization until all forces were below $1000 \text{ kJ mol}^{-1} \text{ nm}^{-1}$. Subsequently, the system was minimized for 100 ps in the canonical (NVT) and isothermal-isobaric (NPT) ensembles. During equilibration all nonhydrogen atoms were position restrained with force constants of $1000 \text{ kJ mol}^{-1} \text{ nm}^{-2}$. In total, 16 simulations were run for 250 ns each. These simulations used two different force-field parameters for the FAD, and in half of the simulations, histidine 378 was mutated to an alanine. In the simulations with H378 present, it was protonated on N^ε since this is suggested to be the protonation state of H378 in the dark-adapted *DmCry* (20). Eight simulations each for the wild type and mutant were using the parameters for oxidized FAD and the anion radical, each with four replicates.

The RMSD of the FFW motif was used as a reporter of the CTT flexibility. Before computing the RMSD, rotational and translational degrees of freedom were removed by aligning the protein on the backbone atoms of residues that displayed the lowest RMSF. The average RMSF for all simulations was computed, and the protein was aligned on the quartile of the residues with the lowest RMSF. After this, the RMSD of the FFW motif was calculated.

Comparison of spectroscopic time constants and those obtained by TRXSS

To find out whether any of the observed structural states correlate with spectroscopic events, we compared the rate constants for deprotonation of W394 and recombination of the radical pair for the two cryptochrome samples (table S5) to the ones obtained from TRXSS (Table 1 and table S4). The deprotonation of the terminal tryptophan (W394) was observed on microsecond time scales, as previously shown for *DmCry* (19). The time constant of the recombination reaction is mainly dependent on the oxidation potential in the buffer. In the TRXSS experiments, the oxidant was potassium ferricyanide, which was added to prevent FAD^{•+} accumulation during data acquisition. Transient absorption measurements were performed using identical molar ratios of protein and potassium ferricyanide as in scattering experiments. Two transient absorption datasets were recorded per sample, one with a 10- μ s time window and one with a 10-ms time window. The 10- μ s datasets were fitted by four exponential decays using Glotaran (44) as described in (19). The 10-ms datasets were fitted with two exponential decays (table S5). k_3 has previously been assigned as the rate constant for deprotonation of the tryptophanyl cation radical (19), and k_5 and k_6 describe the recombination reaction of the radical pair. It is obvious from table S5 that the deprotonation rates are similar for *DmCry* and *DmCry* (H378A) are comparable to published data (19), and are slightly pH dependent. The recombination of the radical pair required two rate constants in *DmCry* samples, and their rates are slightly decreased as compared to samples without ferricyanide (19). *DmCry*(H378A) samples also required two rate constants, and their values were increased with respect to *DmCry*. A detailed investigation and analysis of this behavior is ongoing.

SUPPLEMENTARY MATERIALS

Supplementary material for this article is available at <http://advances.sciencemag.org/cgi/content/full/5/7/eaaw1531/DC1>

Table S1. TRXSS kinetic modeling.

Table S2. Changes in R_g .

Table S3. SAXS parameters.

Table S4. Rate constants for TRXSS measurements of *DmCry* at pH 9.

Table S5. Rate constants for TA measurements of *DmCry*.

Fig. S1. TRXSS data and the reconstructed data using a kinetic model with different numbers of components.

Fig. S2. SAXS scattering profiles for *DmCry* in the dark and under blue-light illumination.

Fig. S3. The RMSD of the FFW motif at each simulation frame.

Fig. S4. TRXSS data and kinetic modeling for wild-type *DmCry* and H378A at pH 9.

Fig. S5. Stability analysis of the used samples via SDS-PAGE and SAXS.

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