Essential Role of an Unusually Long-lived Tyrosyl Radical in the Response to Red Light of the Animal-like Cryptochrome aCRY

Cryptochromes constitute a group of flavin-binding blue light receptors in bacteria, fungi, plants, and insects. Recently, the response of cryptochromes to light was extended to nearly the entire visible spectral region on the basis of the activity of the animal-like cryptochrome aCRY in the green alga *Chlamydomonas reinhardtii*. This finding was explained by the absorption of red light by the flavin neutral radical as the dark state of the receptor, which then forms the anionic fully reduced state. In this study, time-resolved UV-visible spectroscopy on the full-length aCRY revealed an unusually long-lived tyrosyl radical with a lifetime of 2.6 s, which is present already 1 μs after red light illumination of the flavin radical. Mutational studies disclosed the tyrosine 373 close to the surface to form the long-lived radical and to be essential for photoreduction. This residue is conserved exclusively in the sequences of other putative aCRY proteins distinguishing them from conventional (6–4) photolyases. Size exclusion chromatography showed the full-length aCRY to be a dimer in the dark at 0.5 mM injected concentration with the C-terminal extension as the dimerization site. Upon illumination, partial oligomerization was observed via disulfide bridge formation at cysteine 482 in close proximity to tyrosine 373. The lack of any light response in the C-terminal extension as evidenced by FTIR spectroscopy differentiates aCRY from plant and *Drosophila* cryptochromes. These findings imply that aCRY might have evolved a different signaling mechanism via a light-triggered redox cascade culminating in photooxidation of a yet unknown substrate or binding partner.

Cryptochromes represent a group of diverse sensory photoreceptors present in all kingdoms of life (1, 2). Together with the UV-light-dependent DNA repair enzymes, the photolyases (3), they constitute the cryptochrome/photolyase family. Members of this family share a highly conserved photolyase homology region (PHR), which comprises ~500 amino acids and carries a non-covalently bound flavin adenine dinucleotide (FAD) as a chromophore. The C-terminal extension (CCT) present in many cryptochromes and photolyases is strongly variable in amino acid composition and length and has been shown to be crucial for signal transduction in the *Arabidopsis* cryptochrome AtCRY1 (4). The diverse subfamilies of cryptochromes comprise proteins acting as central blue light sensors in bacteria, fungi, plants, and insects (animal type I CRY) (1). Moreover, CRYs are also found as the light-independent, central part of the oscillator of the biological clock in mammals (animal type II CRY) (5) and as a mediator for light-dependent magnetosensitivity in flies (6). Opposed to these findings, DASH cryptochromes have been found to repair lesions in single-stranded DNA and double-stranded loop-structured DNA *in vitro* (7, 8). Therefore, DASH cryptochromes are more similar in terms of functionality to photolyases than cryptochromes. Among the photolyases, two different types are separated depending on their ability to either repair cyclobutane pyrimidine dimers (CPD) (9) or (6–4) photoproducts (10). Animal type II CRY are closely related to eukaryotic (6–4) photolyases whereas plant cryptochromes are homologous to CPD photolyases. In contrast, prokaryotic (6–4) photolyases (11) and bacterial cryptochromes (12) form a distant subfamily.

The paradigm of cryptochromes and other flavoproteins as classical blue light receptors has been challenged recently by the finding that the animal-like cryptochrome (aCRY) from the green alga *Chlamydomonas reinhardtii* strongly influences gene expression not only in response to blue but also to yellow and red light *in vivo* (13). These genes code for proteins involved in chlorophyll and carotenoid biosynthesis, light-harvesting complexes, nitrogen metabolism, cell cycle control, and the circadian clock. The extended spectral sensitivity was explained by the presence of the flavin neutral radical (FADH*) in the dark form of the receptor (Fig. 1A) (13) as opposed to the oxidized flavin (FAD*ox*) found in plant cryptochromes (14, 15).

The abbreviations used are: PHR, photolyase homology region; FAD*ox*, oxidized flavin adenine dinucleotide; CCT, C-terminal extension; aCRY1, *Arabidopsis thaliana* cryptochrome 1; CPD, cyclobutane pyrimidine dimers; aCRY, animal-like cryptochrome; FADH*, FAD neutral radical; FADH*, anionic fully reduced state of FAD; UV-vis, UV-visible; Trp*, tryptophan neutral radical; TyrO*, tyrosyl radical; SEC, size exclusion chromatography; RNR, ribonucleotide reductase; PSI1, photosystem II.

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The absorption spectrum of FADH' covers almost the full visible spectrum of light extending up to 680 nm in agreement with the action spectrum of aCRY (13). In vitro, the predominant part of aCRY was found in the oxidized state after purification. Accordingly, formation of FADH' requires a pre-illumination with blue light which was not applied in the in vivo experiments (13). However, FTIR spectroscopy revealed structural changes in the protein moiety only in the transition from FADH' to the anionic fully reduced state (FADH)|, of aCRY providing further support for FADH| as the dark state of the chromophore (16). Strikingly, these changes in turn structures were not detected in the closely related *Xenopus laevis* (6–4) photolyase (17). The lifetime of FADH' state *in vitro* was strongly sensitive to alterations of the pH but not of the oxygen level in contrast to other subfamily of cryptochromes and photolyases (18–20).

The molecular mechanisms underlying these extraordinary characteristics of aCRY have remained undisclosed up to now because of missing time-resolved information on the early events after red light illumination. Previously, time-resolved UV-visible (UV-vis) spectroscopy on cryptochromes and photolyases has revealed the involvement of transient radical species formed from FAD, tryptophan, and tyrosine as part of the electron transfer cascade (21–27). Tryptophan neutral radicals (Trp') exhibit a broad band centered at around 510 nm (28), whereas tyrosyl radicals (TyrO') are characterized by two sharp, adjacent bands at 388 and 408 nm (29, 30). Moreover, the formation of these amino acid radicals in the electron transfer cascade has been disclosed by time-resolved EPR studies (31, 32). However, to date the photoreduction process of the FADH' has not been studied time-resolved in the cryptochrome as well as (6–4) photolyase families.

In this study, we focused on resolving the events in aCRY starting from the FADH' state by its selective induction using illumination with red light, because the blue light-induced conversion of FADox in aCRY is not considered to be physiologically relevant (13). To elucidate the red light-induced processes, we employed transient UV-vis spectroscopy, size exclusion chromatography (SEC), and FTIR difference spectroscopy on the full-length aCRY as well as on point mutants Y393A, Y373F, C482A (Fig. 1B), and aCRYΔCCT lacking the 99 residues of the CCT. We identified the contribution from a remarkably long-lived TyrO' formed by Tyr-373 close to the surface (Fig. 1B), which is conserved in aCRY homologues but not in any other subfamily of cryptochromes and photolyases. Furthermore, we investigated the role of the CCT in the light-induced structural response and the oligomerization of aCRY.

### Experimental Procedures

**Generation of aCRY Mutants—aCRYΔCCT** coding for amino acids 1 to 496 of the aCRY gene, lacking the 99 amino acids of its CCT, was codon-adapted for *E. coli* (synthesized by Geneart) and cloned into pET28a(+) (Novagen), providing a 6x His-Tag at the C terminus by using the restriction enzyme sites Ncol and HindIII. Mutation Y393A was inserted into the full-length codon-adapted sequence of aCRY (synthesized by Geneart (13)) by the replacement of a 739 bp BstEII and HindIII fragment with a fragment containing the codon-adapted sequence for the substituted amino acid Y393A. Full-length aCRY-Y393A was ligated via Ncol and HindIII restriction sites into vector pET28a(+). The Y373F and C482A mutations were inserted into the full-length aCRY gene in the pET28a(+) vector using phosphorylated back-to-back primers, of which one
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primer contained the mutation. To amplify the whole plasmid, Phusion DNA polymerase (New England Biolabs) was used in the polymerase chain reaction. The reaction products were ligated afterward. The amino acid exchanges in all resulting plasmids were verified by dideoxy sequencing.

Expression and Purification—Expression and purification of aCRY and its variants were conducted following published procedures (13). Finally, the proteins were obtained in a 50 mM sodium phosphate buffer, pH 7.0, 100 mM NaCl, 20% (v/v) glycerol.

Millisecond Time-resolved UV-Vis Spectroscopy—The concentration of the sample was adjusted to $A_{447} = 0.3$. An HR2000+ spectrometer with DH-2000-BAL light source (Ocean Optics) was used for experiments in the millisecond to second time regime modified with a mesh filter with 35% transmission to avoid sample conversion by the probe light. For illumination, a 451 nm LED (Luxeon Star, Lumileds) with an intensity of 67 milliwatt/cm² (FWHM of 20 nm) and a 632 nm LED (Luxeon Star, Lumileds) with an intensity of 95 milliwatt/cm² (FWHM of 15 nm) at the sample was pre-illuminated for 10 s at 451 nm in a 10 mm fluorescence cuvette (Helma). FADH$^-$ was produced by illuminating the sample for 2 s or 10 s at 632 nm. A continuous series of spectra were recorded before, during and after illumination with an integration time of 2 ms and a time resolution of ~40 ms. Difference spectra were calculated and summarized on a logarithmic time scale to increase the signal-to-noise ratio using MATLAB (The Mathworks).

Nanosecond Time-resolved UV-Vis Spectroscopy—The concentration of the sample was adjusted to $A_{447} = 0.5$. The experimental setup for time-resolved UV-vis spectroscopy on slowly recovering systems has been described previously (29). aCRY was pre-illuminated for 15 s under stirring to generate FADH$^+$ using a 455 nm LED (Luxeon Star, Lumileds) with an intensity of 10 milliwatt/cm² at the sample (FWHM of 20 nm). For excitation of FADH$^+$, aCRY was illuminated for $10 \text{s}$ at 451 nm in a 2 × 10 mm fluorescence cuvette (Helma). FADH$^-$ was produced by illuminating the sample for 2 s or 10 s at 632 nm. A continuous series of spectra were recorded before, during and after illumination with an integration time of 2 ms and a time resolution of ~40 ms. Difference spectra were calculated and summarized on a logarithmic time scale to increase the signal-to-noise ratio using MATLAB (The Mathworks).

Size Exclusion Chromatography—SEC was performed using an Akta purifier (GE Healthcare) with a Superdex200 10/300 GL column (GE Healthcare) at 4 °C. For equilibration and elution, 50 mM phosphate buffer at pH 7.0 and 150 mM NaCl were used. Aliquots of 100 μl with a protein concentration of 0.5 mM were centrifuged at 21,400 × g for 10 min at 4 °C. For the investigation of aCRY carrying FAD$_{ox}$ and FADH$^+$, loading of the samples and the SEC were performed in the dark. FADH$^-$ was obtained by illuminating the sample for $10 \text{s}$ with two 451 nm LEDs (Luxeon Star, Lumileds) with an intensity of 67 milliwatt/cm² each at the sample. The elution profiles were recorded at 447 and 630 nm to ensure that only protein with the flavin bound contributes. For the generation of FADH$^-$, samples were centrifuged at 21,400 × g for 10 min at 4 °C. For the investigation of aCRY carrying FAD$_{ox}$ and FADH$^+$, loading of the samples and the SEC were performed in the dark. FADH$^-$ was obtained by illuminating the sample for $10 \text{s}$ with two 451 nm LEDs (Luxeon Star, Lumileds) with an intensity of 67 milliwatt/cm² each at the sample. The elution profiles were recorded at 447 and 630 nm to ensure that only protein with the flavin bound contributes. For the generation of FADH$^-$, samples were centrifuged at 21,400 × g for 10 min at 4 °C. For the investigation of aCRY carrying FAD$_{ox}$ and FADH$^+$, loading of the samples and the SEC were performed in the dark. FADH$^-$ was obtained by illuminating the sample for $10 \text{s}$ with two 451 nm LEDs (Luxeon Star, Lumileds) with an intensity of 67 milliwatt/cm² each at the sample. The elution profiles were recorded at 447 and 630 nm to ensure that only protein with the flavin bound contributes. For the generation of FADH$^-$, samples were centrifuged at 21,400 × g for 10 min at 4 °C. For the investigation of aCRY carrying FAD$_{ox}$ and FADH$^+$, loading of the samples and the SEC were performed in the dark. FADH$^-$ was obtained by illuminating the sample for $10 \text{s}$ with two 451 nm LEDs (Luxeon Star, Lumileds) with an intensity of 67 milliwatt/cm² each at the sample. The elution profiles were recorded at 447 and 630 nm to ensure that only protein with the flavin bound contributes. For the generation of FADH$^-$, samples were centrifuged at 21,400 × g for 10 min at 4 °C. For the investigation of aCRY carrying FAD$_{ox}$ and FADH$^+$, loading of the samples and the SEC were performed in the dark. FADH$^-$ was obtained by illuminating the sample for $10 \text{s}$ with two 451 nm LEDs (Luxeon Star, Lumileds) with an intensity of 67 milliwatt/cm² each at the sample. The elution profiles were recorded at 447 and 370 nm. Standard marker proteins (GE Healthcare) were used to determine the apparent molecular mass of the sample by calibration. Representative traces are shown from the experiments, which were repeated at least three times for each of 2–3 independent preparations.
Phylogenetic Analysis—Phylogenetic analyses were conducted using the MEGA software package version 6 (33). The protein sequences used for the alignment were selected according to the results of the NCBI BLASTP 2.3.0 (34) using the sequences with the highest identity to aCRY. Additionally, sequences of closely related and well characterized cryptochromes and photolyases were included (supplemental Table S1). The protein sequences were aligned with the ClustalW algorithm.

Results

The UV-vis spectrum of aCRY recorded after purification revealed FADox to be present in aCRY in vitro (Fig. 1C). Blue light illumination then generated FADH+/H+ with characteristic bands at 585 and 633 nm. We noticed an additional small band at 416 nm with a shoulder at 410 nm in the absorption spectrum after illumination, which was only present for a few seconds (Fig. 1C). Such contribution is indicative for transient accumulation of an amino acid radical, but usually detected on a much shorter time scale. Therefore, this finding was scrutinized in a time-resolved manner in the context of the physiologically more relevant reduction of FADH+/H+ to FADH−.

Microsecond Time-resolved Response to Red Light of aCRY—Time-resolved UV-vis experiments have revealed the involvement of tryptophan and tyrosyl radicals in the photoreaction of cryptochromes in vitro starting from FADox. Here, flash photolysis was applied to detect possible amino acid radicals involved in the red light-induced photoreduction of FADH+/H+ in aCRY. Previous steady-state experiments have shown only the formation of FADH− (16).

Samples containing FADox were pre-illuminated with blue light for 15 s to generate a sufficient amount of FADH+/H+, which was stabilized by adjusting the pH to 7.0 (16). To selectively convert FADH+/H+ to FADH−, nanosecond laser pulses at 630 nm were used. Time-resolved UV-vis difference spectra were recorded at time points from 1 μs to 2 ms (Fig. 2A). All difference spectra show the characteristic negative band pattern of the bleaching of FADH+/H+ with two broad maxima at 590 nm and 630 nm. Additionally, a small positive contribution from the formation of FADH−/H+ is detected at 400 nm. For comparison, spectra after illumination for 2 s with an LED were obtained in the absence and in the presence of 0.1 mM ascorbic acid. This strong reducing agent quenches all contributions by amino acid radicals within a few microseconds. The time-resolved spectra show a pronounced positive band at 416 nm, which is still present 2 s after illumination but is missing in the presence of ascor-
bic acid. This additional contribution is present to an equal extent relative to flavin at all time points.

To reveal the identity of the species contributing at 416 nm, the double difference spectrum of the 1/H9262s spectrum minus the steady-state spectrum was calculated. The resulting spectrum shows a well-defined peak with a maximum at 416 nm and a shoulder at 400 nm (Fig. 2B). The comparison to reference spectra (29, 30) revealed that such a band is characteristic for TyrO/H18528. In contrast to the reference spectra obtained in water, TyrO/H18528 in aCRY is red-shifted by the protein environment by 9 nm (inset, Fig. 2B). Any hint for the presence of a tryptophan radical at 1/H9262s or later was not found.

**Kinetic Analysis of the Decay of TyrO in aCRY after Red Light Illumination—**Tyrosyl radicals in proteins usually exhibit maximal lifetimes in the range of a few milliseconds (21, 23, 29, 35). Here, the time-resolved studies on aCRY indicated a much higher lifetime. Therefore, the decay of TyrO and FADH− after red light illumination was analyzed on a time scale of seconds at 20 °C. The sample was pre-illuminated with blue light and then exposed to 632 nm for 2 s. A continuous series of difference spectra with a time resolution of ~40 ms was recorded for 15 s (Fig. 3). The decay of FADH− was analyzed indirectly using the negative marker bands of the recovery of FADH2. The absorbance at 615–643 nm was averaged and subsequently analyzed with a monoexponential fit yielding a time constant of 2.6 s. For the decay of TyrO, changes in absorbance at 395–420 nm were averaged. The monoexponential fit yielded a time constant of 2.6 s for TyrO, implying a mutual decay with FADH2.

**Candidates for the Tyrosine Residue Involved in the Red Light Response of aCRY—**To reveal the identity of the tyrosine residue contributing to the difference spectra, two different candidates for the Tyrosine Residue Involved in the Red Light Response of aCRY—

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**FIGURE 4. Effects of mutations in tyrosine residues on the red-light response of aCRY.** A, red light-induced UV-vis difference spectra of the wild type, the Y393A and Y373F mutants of aCRY. Pre-illuminated samples were illuminated for 10 s with 632 nm. In both the wild type and the Y393A mutant, FADH− is converted to FADH2 accompanied by the formation of TyrO. The Y373F mutant does not show any light response. B, time-resolved absorption spectra at 500 ns of the Y373F mutant and wild type aCRY in the presence of 3 mM ascorbic acid. Both the Y373F mutant and the wild type show the conversion of FADH− to FADH2. In contrast to the wild type, the Y373F mutant does not show an additional contribution of TyrO at 416 nm identifying Tyr-373 as the origin of this absorption. The conversion of FADox to FADH2 and FADH2 is observed in aCRY-Y373F only in the presence of ascorbic acid (inset).

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**FIGURE 5. Size exclusion chromatography of wild-type aCRY, aCRYΔCCT and the C482A mutant at injected concentrations of 0.5 mM.** A, samples were illuminated for 10 s with 451 nm to convert FADox to FADH2, which was selectively detected at 630 nm. Wild-type aCRY and the C482A mutant eluted at 13.2 ml, which corresponds to an apparent molecular mass of 130 kDa. Considering a theoretical molecular mass of 66.4 kDa of aCRY, the bands can be assigned to dimers. aCRYΔCCT predominantly eluted at 14.8 ml, which corresponds to an apparent molecular mass of 60 kDa and can accordingly be assigned to a monomer. B, samples were illuminated for 45 s with 451 nm and 630 nm to convert FADox to FADH2 and FADH2, which were detected at 370 nm. Wild-type aCRY shows an additional, small peak at 11.4 ml (314 kDa), which indicates a tetrameric species. This light-induced oligomerization can be abolished by the addition of the reducing agent TCEP. C, samples were treated as in B. In the C482A mutant, light-induced oligomerization is not observed. In contrast, aCRYΔCCT shows an additional peak at 13.3 ml, which can be assigned to a dimeric species. Therefore, some light-induced oligomerization was found for both wild-type aCRY and aCRYΔCCT.
mutants of aCRY, Y373F and Y393A were produced. The residues Tyr-373 and Tyr-393 were selected because they are not conserved in other cryptochrome or photolyase families and might therefore provide a basis for the unique photoreceptor properties of aCRY (Fig. 1B). Both mutants were expressed and purified as soluble yellow proteins. Samples were pre-illuminated for 10 s with 451 nm to generate FADH$^\ast$, but in Y373F, FADH$^\ast$ was not formed. Differences spectra were then recorded after illumination of the samples for 10 s with 632 nm (Fig. 4A). For comparison, the same procedure was repeated with wild-type aCRY. Both the wild type and the Y393A mutant showed the typical pattern of the conversion of FADH$^\ast$ to FADH$^-$ accompanied by the formation of TyrO$^\ast$. In contrast, the Y373F mutant did not show any light-induced difference in absorption. These findings lead to the conclusion that in aCRY-Y373F the light sensitivity is strongly reduced by the exchange of the tyrosine residue, whereas Tyr-393 seems not to have any influence on the photoreaction. However, it remained unclear whether Tyr-373 is the residue forming the radical or whether the mutation negatively affected the structural integrity of aCRY.

**Identification of Tyr-373 Forming the Long-lived Radical**—We have investigated whether there are conditions under which a photoreduction of the Y373F mutant was observable. The protein could be photoreduced from FAD$_{ox}$ to FADH$^-$ and further to FADH$^-$ by the addition of ascorbic acid, albeit after a much longer light exposure (Fig. 4B, inset). However, under these harsh conditions TyrO$^\ast$ in the wild type was quenched preventing a direct comparison with the mutant (Fig. 2). Therefore, time-resolved experiments were conducted to capture TyrO$^\ast$ in the wild type before its reaction with ascorbic acid. Conditions had to be established under which the Y373F mutant was sufficiently photoactivated while the wild type showed the signal of TyrO$^\ast$, resulting in time-resolved spectra at 500 ns in the presence of 3 mM ascorbic acid. After pre-illumination with blue light, both samples were excited with laser pulses at 532 nm. The intensity was adjusted so that the difference spectra of the wild type and the Y373F mutant showed both the formation of the FADH$^-$ to a similar extent (Fig. 4B). The spectrum of the wild type showed the same band pattern as a sample in the absence of ascorbic acid besides an additional band at around 370 nm originating from a reaction intermediate in the oxidation of ascorbic acid. The Y373F mutant, however, does not produce the band at 416 nm of TyrO$^\ast$. In conclusion, Tyr-373 was identified to form the long-lived radical in aCRY.

**Size Exclusion Chromatography on aCRY and Its Variants**—The oligomerization state of aCRY and its dependence on the redox state of the flavin was studied by SEC at injected concentrations of 0.5 mM. aCRY containing FADH$^\ast$ was analyzed using blue light-illuminated samples. Absorbance was recorded at 630 nm to selectively detect protein with bound FADH$^\ast$. The SEC showed a single peak with a maximum at an elution volume of 13.2 ml (Fig. 5A), which corresponds to an apparent molecular mass of 130 kDa. Considering a theoretical molecular mass of 66.4 kDa, the peak can be assigned to a dimer of aCRY. To resolve the functional role of the CCT, aCRY was truncated after amino acid 496. In contrast to the full-length receptor, aCRY$\Delta$CCT showed a peak at an elution volume of 14.8 ml, which corresponds to an apparent molecular mass of 60 kDa (Fig. 5A). With a theoretical mass of 57.7 kDa of aCRY$\Delta$CCT, the peak can be assigned to a monomer. This difference points to the fact that the CCT is necessary for the dimerization of aCRY carrying FADH$^\ast$.

The effects of the formation of FADH$^-$ in aCRY were investigated using blue and red light-illuminated samples. Due to the decay of FADH$^-$ back to FADH$^+$ within a few seconds during the SEC, the detected absorbance at 370 nm mainly represents FADH$^+$ in aCRY, but FADH$^-$ and FAD$_{ox}$ might absorb as well. Interestingly, aCRY showed the same profile as before, but with an additional peak at 11.4 ml (314 kDa) possibly from a tetrameric species (Fig. 5B). Similarly, aCRY$\Delta$CCT showed an additional peak at 13.3 ml (122 kDa) after the conversion, which is assignable to the formation of a dimer (Fig. 5C). Therefore, it can be concluded that the red light-induced changes in aCRY are persistent even after the fast recovery of FADH$^+$ from FADH$^-$.

The formation of FADH$^-$ is accompanied by light-induced oligomerization of aCRY to some extent. This process is abolished by the presence of 1 mM of the strong reducing agent TCEP (Fig. 5B). As TCEP is known to reduce disulfide bonds, a potential candidate for such an intermolecular bond was identified as Cys-482 on the basis of its close proximity to

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**FIGURE 6.** Red light-induced FTIR difference spectra of aCRY$\Delta$CCT as compared with wild-type aCRY. Samples were pre-illuminated with blue light and then exposed to 632 nm for 10 s to selectively convert FADH$^\ast$ to FADH$^-$.

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**TABLE 1.** Properties of aCRY and Its Variants

| Protein   | Molecular Mass (kDa) | Oligomerization State |
|-----------|----------------------|-----------------------|
| aCRY      | 130                  | Dimer                 |
| aCRY$\Delta$CCT | 60                   | Monomer               |

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**FIGURE 2.** Blue light-induced SDS-PAGE analysis of aCRY and its variants. aCRY was incubated with blue light for 10 s and then subjected to SDS-PAGE and Coomassie Blue staining. Wild-type aCRY and aCRY$\Delta$CCT show bands corresponding to dimer and monomer, respectively.
the reactive tyrosine residue Tyr-373 (Fig. 1B) as well as its position close to the protein surface. The replacement of Cys-482 with alanine led to a disappearance of the additional peak in the SEC profile of aCRY after red-light illumination (Fig. 5C), whereas the profile of aCRY without red light treatment was not altered in the C482A mutant (Fig. 5A). Therefore, light-induced oligomerization in aCRY most likely proceeds via disulfide bridge formation at Cys-482.

Red Light-induced FTIR Difference Spectroscopy on aCRYΔCCT—The previously recorded FTIR difference spectrum of the wild type (16) showed changes in turn structures in the light-induced conversion from FADH to FADH− that are not present in (6 – 4) photolyase (17). To further investigate a possible involvement of the CCT in the light red light response of aCRY, difference spectra of aCRYΔCCT were taken after red light exposure. The reference was recorded after the sample was pre-illuminated for 1 s with blue light. Immediately afterward the sample was illuminated for 10 s at 632 nm to induce the formation of FADH− (Fig. 6A). In the resulting difference spectrum, only bands of vibrational modes changing upon red light illumination are detectable. The negative bands represent the dark state with bound FADH−. Marker bands for FADH− (16) are present in the spectrum of aCRYΔCCT at 1626, 1620, 1515, and 1395 cm−1. The positive band in the difference spectrum of aCRYΔCCT are also in good agreement with those of the wild type, most of which can be assigned to FADH−. Marker bands for FADH− (16) are present in the spectrum of aCRYΔCCT at 1676 cm−1. Only the band at 1564 cm−1 originating from both flavin and the amide II band shows an upshift of 5 cm−1 as compared with the wild type spectrum. In conclusion, the difference spectrum of aCRYΔCCT is almost identical to that of the wild type taking into account a slight shift of several bands by ~1 cm−1. It can be concluded that the CCT in aCRY does not respond to light by a change in secondary structure, in contrast to the CCT in plant cryptochromes (36, 37). So far, structural changes detected in aCRY are therefore limited to the PHR domain.

Discussion

Unusual Stability of TyrO in aCRY—Tyrosyl radicals have been observed as part of the electron transfer cascade in other cryptochromes and photolases than aCRY (21, 23, 29, 32). Lifetimes of these tyrosyl radicals have been determined to range from 1 ms up to >140 ms (21, 23, 29). In contrast, an unusually long-lived TyrO with a lifetime of 2.6 s is formed in aCRY, implying a high stability of the radical.

A very similar lifetime of TyrO has been achieved in α3Y, a de novo model protein specifically designed for investigating the stabilizing effect of the protein matrix on the lifetime of tyrosyl radicals (38). α3Y does not contain any cofactors and places the reactive tyrosine in a three helix bundle scaffold. Similar to aCRY, α3Y does not carry a covalent modification of tyrosine or a metal center for stabilization of TyrO (39). However, the residue is buried deeply in a hydrophobic environment in α3Y without any hydrogen bonding partner or access to the solvent (38). Therefore, the surrounding protein matrix is rather different compared with the one present in aCRY, considering the relatively exposed position of Tyr-373 in a cavity on the surface of the protein, as derived from the modeled structure (16).

Interestingly, a system more similar to aCRY was found in the D2 subunit of photosystem II (PSII). Here, TyrD forms a radical that is stable for hours under physiological conditions without a metal cluster in close proximity (40). TyrD was reported to be deprotonated upon oxidation, because phenolic compounds become strong acids upon oxidation. Such a proton-coupled electron transfer has also been found for other tyrosyl radicals (39). For TyrD, it is believed that the kinetics of reprotonation is of considerable importance for the stability of TyrO (41). The radical is found in a very well-ordered hydrogen bonding network with rather hydrophobic residues, which was postulated to prevent any uncontrolled protonation (41). Additional stability might be provided by the movement of either the proton acceptor (41) or the tyrosine residue away from the proton-donating network upon oxidation, as found for the stable TyrO coupled to a diiron o xo site in subunit R2 of class I ribonucleotide reductases (RNR) (42). Moreover, it might be important to consider that after oxidation and deprotonation, the phenolic oxygen of the radical might be hydrogen-bonded to an adjacent residue or a coordinated water molecule (43, 44). This hydrogen bond may provide an additional stabilization by delocalizing the spin density of the oxygen, as it has been suggested for the rather hydrophilic environment in mouse RNR (44).

Transferring these findings to aCRY, it seems plausible that TyrO is also coordinated in a very stable, non-flexible hydrogen bonding network comparable to the one found for TyrD in PSII, even if there are not that many adjacent hydrophobic residues in aCRY. In PSII, arginine and aspartate close to TyrD form a salt bridge, facilitating the proton transfer during oxidation (41). In the modeled structure of aCRY (16), arginine (Arg-485) and aspartate (Asp-321) also form a salt bridge close to Tyr-373, possibly creating a similar environment as in PSII. Still, it is unclear whether an amino acid residue or a coordinated water molecule is the proton acceptor in aCRY. Furthermore, it needs to be resolved if the phenolic oxygen of TyrO is also hydrogen bonded as in mouse RNR and PSII-D2. In summary, the lifetime of TyrO in aCRY can be considered to be unusually long because the protein does not provide the classical environment for stabilization such as a metal center or hydrophobic residues. Furthermore, TyrO in aCRY does not benefit from possibly stabilizing effects of a membrane-bound complex such as PSII.

Red Shift of TyrO Absorbance—Another striking feature of TyrO found in aCRY is its red-shifted absorption pattern. Typically the radical shows absorption maxima at ~390 and ~410 nm (29, 30). In contrast, the maxima in aCRY are found at ~400 and 416 nm, similar to what has been reported previously in mouse RNR (45). An explanation for the shift in both aCRY and the mammalian RNR might be the existence of the hydrogen bond between the phenolic oxygen and a hydrogen bond donor. Interestingly, TyrO in the Escherichia coli RNR is not hydrogen-bonded and shows a non-shifted absorption spectrum in the direct comparison (45). A covalent crosslink of the cysteine to the tyrosine as observed in galactose oxidase (46) can be
ruled out as the absorption difference spectrum of the aCRY-C482A mutant is identical to that of the wild type (data not shown).

**Formation of TyrO from Trp**—Another surprising aspect of the time-resolved UV-vis spectra is the absence of any bands from tryptophan radicals after 1 μs. In other UV-vis-spectroscopic studies on cryptochromes, significant Trp contributions have been identified in this time region up to milliseconds (23, 24). To our knowledge, the only similar finding has been reported for Xenopus laevis (6–4) photolyase, in which TyrO is already formed within 100 ns after illumination (32). For aCRY, the question arises whether the Trp decays ultrafast after TyrO or if the Trp is not formed in the first place. The latter suggestion is unlikely because a tryptophan triad (23) (Trp-322, Trp-376, Trp-399) is also conserved in aCRY. As such a triad is considered to represent an optimal electron transfer pathway, the electron is most probably guided via this “electron wire” over the distance of ~16 Å, as derived from the model, from the Tyr-373 on the surface to the chromophore. Therefore, the most likely reason for the Trp band not to be present in the spectra after 1 μs is its ultrafast decay in a time range of pico- to nanoseconds. This short lifetime might be the consequence of a high reactivity of Tyr-373.

**Oligomerization State of aCRY**—In the dark, SEC at 0.5 mM injected concentration revealed aCRY to form a homodimer whereas the truncated ΔCCT mutant remains predominantly as a monomer. These findings indicate that the CCT may represent the oligomerization site of aCRY in the dark. In the light, SEC revealed some oligomerization for both proteins upon FADH\(^{-}\) formation at a concentration of 0.5 mM. Therefore, the CCT does not play a significant role during red light-induced oligomerization. This conclusion is underlined by the FTIR experiments which do not show any significant differences between red light-induced difference spectra of the wild type and the ΔCCT mutant. These findings are in contrast to events observed in other cryptochromes. In the Arabidopsis cryptochrome AtCRY1, large conformational changes upon photoexcitation were noted by a rearrangement of the CCT (36, 37). In the Drosophila cryptochrome dCRY, an increase of proteolytic susceptibility of the CCT upon illumination indicates the involvement of this domain in the light response (47).

Further SEC experiments imply that the light-induced oligomerization originates from the formation of disulfide bridges most likely by Cys-482. From the structural model, Cys-482 was derived to be in close proximity with a 4 Å distance to the reactive Tyr-373. It can be concluded that red light illumination of aCRY triggers a redox cascade with a partial electron transfer from a cysteine to TyrO. Such transfer has already been observed in class I RNR, in which the cascade does not result in disulfide bridge formation but in the reduction of ribonucleotides to deoxyribonucleotides (48). In aCRY this reaction is inefficient, because the lifetime of TyrO is not increased in the C482A mutant but even slightly shortened (Fig. 3B). Therefore, the formation of disulfide bridges might be regarded as a nonphysiological reaction or as a consequence of a missing in vivo redox substrate/partner, which might explain the low extent of oligomerization. Alternatively, we may speculate that the covalent linkage of the aCRY dimers constitutes a pathway present under oxidative stress and high light conditions only, which might lead to an inactivation of aCRY signaling by inhibiting any binding to its signaling partner (49).

**Tyr-373 in the Context of Other Cryptochromes and Photolyases**—The postulate of a central role of Tyr-373 in the sensory mechanism of aCRY would imply that this residue is conserved in other potential red light receptors. A BLAST search (Fig. 7, supplemental Table S1) revealed that indeed the two closest relatives of aCRY, the putative photolyases from the algae Volvox carteri and Dunaliella salina share this residue as well as a sequence from Chlamydomonas reinhardtii but not from Chlorella variabilis. These sequences form a subgroup that is well separated in the phylogenetic analysis from other candidates for conventional (6–4) photolyases. Even more convincingly, the next 40 candidates in the alignment of the aCRY homologues and all well characterized members of the cryptochrome/photolyase family to our knowledge do not possess a tyrosine at this position. We consider this finding to be significant as a further support for the crucial role of Tyr-373 in the function of aCRY as light receptor with an extended sensitivity range.

**Conclusions**—To date, only small conformational changes have been identified for aCRY in the red-light induced photoreduction. In contrast to other full-length cryptochromes, these
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Tyr-373 within 1 oxidation of a yet unknown substrate or binding partner. a light-triggered redox cascade, which culminates in a photoimply that aCRY might have evolved a signaling mechanism via tyrosine 373. These unexpected findings point to an important tochrome/photolyase superfamily. Partial light-induced oligo- ligation. This residue is only conserved within a small group of sines 373 was disclosed to be responsible for the radical forma-

changes resulting from illumination do not involve the CCT. This finding raises the question of how the signal proceeds from the flavin to the protein surface in aCRY if not by such confor- mational changes. An alternative model would involve a redox cascade starting with the reduction of flavin and proceeding to the surface, where a signal partner or a substrate may then be converted (50).

This study provides insights into the early events of the red light response of aCRY (Fig. 8). We revealed the rapid formation within 1 µs of a tyrosine radical with an unusually long lifetime of 2.6 s, which decays concomitant with FADH+. Tyrosine 373 was disclosed to be responsible for the radical formation. This residue is only conserved within a small group of proteins that might be regarded as a new subfamily of the cryptochrome/photolyase superfamily. Partial light-induced oligo-

merization of the protein was observed and attributed to disulis-

dide bridge formation via cysteine 482 in the absence of external reductant.

FIGURE 8. Model of the response to red light of aCRY in vitro. aCRY forms a homodimer (represented by two disks) via its CCT. aCRY carrying FADH+ is reduced to FADH− by light with λ < 680 nm. As a result, TyrO+ is formed by Tyr-373 within 1 µs with a red-shifted absorption in aCRY compared with water. The lifetime of 2.6 s of TyrO+ is unusually high and might allow for an efficient oxidation of a signaling partner in vivo finally leading to the changes in the levels of multiple transcripts. The decay of TyrO+ proceeds concomitant with that of FADH−. Some of the light-activated aCRY oligomerizes via disulfide bridge formation at Cys-482 in the absence of external reductant.

Author Contributions—T. K., M. M., and L. O. E. designed the study. S. O. performed and analyzed all the experiments with advice by T. K. S. F. and S. W. generated plasmids of aCRY variants. S. O. and T. K. wrote the manuscript with advice by S. F., S. W., L. O. E., and M. M. All authors reviewed the results and approved the final version of the manuscript.

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