Tryptophan-to-heme electron transfer in ferrous myoglobins

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It was recently demonstrated that in ferric myoglobins (Mb) the fluorescence quenching of the photoexcited tryptophan 14 (*Trp14) residue is in part due to an electron transfer to the heme porphyrin (porph), turning it to the ferrous state. However, the invariance of *Trp decay times in ferric and ferrous Mbs raises the question as to whether electron transfer may also be operative in the latter. Using UV pump/visible probe transient absorption, we show that this is indeed the case for deoxy-Mb. We observe that the reduction generates (with a yield of about 30%) a low-valence Fe–porphyrin π[Fe(porph••)]–anion radical, which we observe for the first time to our knowledge under physiological conditions. We suggest that the pathway for the electron transfer proceeds via the leucine 69 (Leu69) and valine 68 (Val68) residues. The results on ferric Mbs and the present ones highlight the generality of Trp–porphyrin electron transfer in heme proteins.

Electron transfer plays a fundamental role in many biological systems (1–3) ranging from photosynthetic proteins (4) to iron–sulfur (5), copper (6), and heme (7, 8) proteins. It was demonstrated that electron transfer can be used to produce from heme proteins in situ drugs with antimarialar activity (9) and it might have a role in protein folding (2). In general, electron transfer in proteins can occur over long distances (>10 Å) by hopping through different residues, thus reducing the time that would be needed for a single step tunneling from the donor to the acceptor (10–12). Aromatic amino acids and Tryptophan (Trp) in particular can act as a relay in such processes (13–19). Trp also acts as a phototriggered electron donor, e.g., in DNA repair by photolysis (16–18) and in cryptochromes (20, 21). When no obvious electron acceptors are present, excited Trp or (*Trp) still displays shorter lifetimes than its nanosecond decay times in solution (22, 23). This is due to its strong tendency to act as an electron donor, undergoing electron transfer toward the protein’s backbone as in the case of apo-myoglobin mutants (24), small cyclic peptides (25), and human γ-crystallin (26). It is interesting to note that in wild-type horse heart (WT-HH) apo-myoglobin the fluorescence lifetime of the two *Trp residues was reported to be comparable to that in water (27), demonstrating the absence of deactivation mechanisms, either by energy or by electron transfer.

The protein visible absorption spectrum is dominated by their cofactors, e.g., heme or flavins, whereas the UV absorption in the region between 250 nm and 300 nm is mainly due to the three aromatic amino acids, Trp, tyrosine (Tyr), and phenylalanine (Phe) (28), with Trp having the highest molar extinction coefficient. The high sensitivity of Trp to the local environment and the possibility to correlate it with its fluorescence response (28) have led to its widespread use as a local natural probe of protein structure and dynamics in time-resolved fluorescence resonance energy transfer (FRET) studies, and it has emerged as the “spectroscopic ruler” in such studies (28–30). FRET is mediated by dipole–dipole coupling between a donor *Trp and an acceptor molecule, and its rate is inversely proportional to the sixth power of the distance between them and to the relative orientation of their dipoles.

**Significance**

We demonstrate the occurrence of tryptophan (Trp) to heme electron transfer (ET) in ferrous myoglobins by ultrafast UV spectroscopy. The ET gives rise to the theoretically predicted, low-valence Fe(II)(porph••)–anion radical, which we observe for the first time to our knowledge under physiological conditions. These results highlight the generality of Trp–porphyrin electron transfer events in heme proteins and question the systematic use of Trp fluorescence in FRET studies of protein dynamics.

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Results and Discussion
The static absorption spectra of deoxy-Mb and Trp (SI Appendix, Fig. S1) show that at wavelengths <310 nm, the contributions of the heme and Trp overlap, excluding selective excitation of the Trp residues. To disentangle the Trp and heme contributions we performed TA measurements exciting deoxy-Mb at 315 nm, where only the heme absorbs, and compared them to those exciting at 290 nm where both Trp and heme absorb. Assuming that the heme response is similar for both excitation wavelengths, this allows separation of the heme and the Trp contributions in the transient absorption signal. It must be noted that in the probe range used in our experiments (390–730 nm), Trp excited-state absorption (ESA) and stimulated emission (SE) as well as an eventual Trp–photoproduction absorption may contribute to the transient signal (22).

Photocexcitation at 315 nm. Fig. 1A shows TA spectra, at selected pump–probe time delays, obtained upon 315-nm excitation of deoxy-Mb (more TA spectra are shown in SI Appendix, Fig. S5). Two negative features appear at ~430 nm and ~550 nm that are due to ground state bleach (GSB) of the Soret and Q bands, respectively (SI Appendix, Fig. S1). Positive features due to ESA are observed at 450 nm and 600 nm, respectively (46, 52), which shift to the blue within the first 10–15 ps, while becoming weaker. The apparent shift of the GSB features results from the dynamics of the overlapping ESA contributions. Two mechanisms were proposed to explain the heme photocycle, namely the system undergoes vibrational relaxation (46, 52) or relaxes by cascading through spin states (56). However, our purpose here is not to discuss these mechanisms as they occur in the first few picoseconds or so and do not influence the *Trp kinetics we are investigating.

The timescales related to the relaxation of deoxy-Mb were retrieved by both a singular value decomposition (SVD) analysis and a global fit (GF). The fit function, used to recover the involved timescales, is a sum of exponential decays convoluted with the instrumental response function (IRF) (~300 fs), assumed to be Gaussian. The timescales, obtained by a GF of the kinetic traces (SI Appendix, Fig. S3), are 280 ± 60 fs, 1.6 ± 0.2 ps, and 4.0 ± 0.4 ps, in agreement with the literature (49, 52, 56). The large error for the 280-fs contribution results from its proximity to the duration of the IRF. In Fig. 1B we show the decay-associated spectra (DAS) obtained from the SVD analysis. The DASs are due to the amplitudes of the exponential decay functions used to best fit the data points, allowing us to define whether a certain feature is decaying or rising. A DAS is related to a particular timescale and it can be read by comparing it with the transient spectrum at the corresponding time delay. If the amplitude of the DAS has the same sign as the spectrum, the feature is decaying (e.g., a negative DAS in the GSB region), whereas if the DAS has opposite sign with respect to the spectrum, the spectral feature is rising (e.g., a negative DAS in the spectral region corresponding to an ESA feature).

As mentioned above, the interpretation of the mechanism related to the heme relaxation is still a subject of debate (46, 52, 56–58). Our aim here is not to discuss these mechanisms. Important is that the longest timescale in the heme photocycle is ~4 ps, which is much shorter than the Trp decay times (~20 ps and ~120 ps).

Photoexcitation at 290 nm. Fig. 2A shows TA spectra at selected time delays, obtained upon 290-nm excitation (more TA spectra are shown in SI Appendix, Fig. S6). They display GSB features at ~430 nm and 550 nm due to the Soret and Q bands, respectively. The latter is overlapped with a very broad unstructured positive contribution that we assign to ESA of the photoexcited Trp residues (22). Additionally the two ESA features of the heme (~450 nm and ~600 nm, corresponding to the Soret band and the Q band, respectively) are observed for small pump–probe delays. For time delays <15 ps, the TA spectra exhibit the same behavior as upon 315-nm excitation, namely the ESA features shift to the blue and decrease in intensity while the GSB decreases in intensity. However, at longer pump–probe delays the TA signals show a persistent GSB feature at ~430 nm and two ESA features, at ~460 nm and at ~610 nm, respectively (Fig. 3). The TA spectra maintain the same shape from delay times of ~40 ps up to 1 ns, except for a small reduction in intensity (over the entire probe range) in the first 100 ps due to the disappearance of the *Trp14 and *Trp7 ESA. Fig. 3 compares the TA spectra at 900 ps upon 315-nm (Fig. 3A) and 290-nm (Fig. 3B) excitation. The shape and the amplitude of these transients differ significantly, suggesting the formation of a long-lived (LL) photoproduction for 290-nm excitation. As this photoproduction spectrum displays clear features in the Soret- and Q-band region, it indicates a modification...
of the heme group. Further, it cannot result from a *Trp–heme FRET process, because the heme photocycle is very short. The *Trp FRET rate would be the rate-limiting step and no LL photoproduct would be observed.

An SVD analysis and a GF (SI Appendix, Fig. S3) were performed to determine the kinetics of the spectral evolution. The fits, using six exponential components, yielded time constants of 230 ± 60 fs, 1.5 ± 0.2 ps, 4.4 ± 0.4 ps, 18 ± 2 ps, and 106 ± 12 ps and a long component (set to 5 ns) that accounts for the LL signal. All time constants (except for the 5 ns) were free parameters of the fit and the results are in excellent agreement with the decay times for the heme obtained above (see above) and with the literature values for the *Trp decay times (35, 37).

Fig. 2B shows the DASs obtained for the *Trp and *Trp⁴ decay times (106 ps and 18 ps, respectively). Additionally the DASs assigned to *Trp and *Trp¹ decay times were free significantly, indicating different relaxation pathways. The former contains a decay of the *Trp ESA as well as a response of the heme observed on the same timescale, because FRET is the rate-limiting step. The 18-ps DAS (Trp¹) is almost a mirror image of the LL DAS. More precisely, around 430 nm the positive feature in the Trp¹ DAS mirrors the negative feature present in the LL DAS, although it is somewhat narrower. Furthermore, the two DASs mirror each other in the entire range from 460 nm to 730 nm, bearing in mind an overall small positive offset in the Trp¹ DAS.

As mentioned above, the LL photoproduct must be related to a change of the heme group and is not due to a *Trp–heme FRET. Because a phototrigged Trp¹-to-heme electron transfer was already reported for ferric Mbs (40) and because the *Trp decay times are almost invariant for all Mbs (SI Appendix, Table S1), this suggests that a photoinduced Trp¹-to-heme electron transfer also occurs in the ferrous deoxy-Mb. The resulting low-valent heme could be either an Fe⁺ heme or an Fe⁳⁺ porphyrin π-anion radical [Fe⁺(porph)¹] complex, if the additional electron resides on the porphyrin ring (59–63). Several studies were performed, with a wide variety of techniques, on low-valent iron complexes, both as Fe⁺–porphyrin and Fe⁳⁺(porph)¹ (42, 53, 54, 60–64). However, a large part of these studies focuses on tetrapyrrol-porphyrins (TPP) and octaethylporphyrins (OEP) in organic solvents (61, 64). It was concluded that formation of Fe⁺–porph or Fe⁳⁺(porph)¹ depends sensitively on the relative energy of the iron d⁷ 2g orbitals and the porphyrin e₃ orbitals (SI Appendix, Scheme S1) (62). One way to experimentally affect the relative energies of these orbitals is substitution of the hydrogens in the porphyrin meso positions (e.g., in TPP and OEP) (62). If electron-withdrawing substituents are introduced in the ring, the energy of the e₃ orbitals will decrease, making the π-anion radicals more likely (62). On the other hand, if electron-donor groups are present in the ring, the energy of the e₃ orbitals will become higher, leading to Fe⁵ complexes (62, 63, 65). In the case of the Fe⁵(porph)¹ species, absorption spectra display a broad band centered at ~700 nm and ~450 nm, and the Q and Soret bands disappear (62).

The LL photoproduct absorption spectrum (Fig. 3C) is obtained by subtracting the GSB contribution to the transient signal at 900 ps (Fig. 3B). It is comparable to the absorption spectrum of the reduced Fe⁵(NO₂-OEP), which generates a porphyrin π-anion radical (62). However, it is likely that this comparison is qualitative as the porphyrin, the solvent, and the environment differ. In Fig. 3C, the Soret band and the Q band are nearly vanished and new bands arise around 450 nm and 600 nm. This comparison leads us to conclude that the anion radical Fe⁺(porph)¹ is formed.

This is further supported by cryo-radiolysis experiments (41, 42). EPR/ENDOR studies of Mb at ~70 K show that upon γ-ray irradiation, a mixture of Fe⁺–Mb and Fe³⁺(porph)¹–Mb is generated, in a 9:1 ratio (42). The authors suggested that different conformations in the frozen protein complexes might explain the simultaneous observation of both species. Annealing experiments hint at the possibility that the decay of Fe³⁺ species could involve intramolecular electron transfer, leading to the formation of Fe⁺.
Fe^{II}(porph^\bullet\bullet\bullet) (42). The latter results suggest that the \epsilon_g and \epsilon_{\pi-}\pi orbi-
tals are close in energy, leading to the Fe^{II}(porph^\bullet\bullet\bullet) when the system has the possibility to relax. Low-valent heme species, their nature, and relevance under physiological conditions were also investigated in theoretical studies (43–45), which also suggest formation of an Fe^{II}(porph^\bullet\bullet\bullet).

The photoproducts of Trp^{14}-to-heme electron transfer can be TrpH\bullet\bullet\bullet or *TrpH\bullet\bullet\bullet and Fe^{II}(porph^\bullet\bullet\bullet) or * Fe^{II}(porph^\bullet\bullet\bullet). Because the transient spectra at delay times >40 ps do not display any changes (except for a small vertical offset due to *Trp\bullet\bullet\bullet and *Trp^{14} ESA), it is safe to assume that the Fe^{II}(porph^\bullet\bullet\bullet) product is generated. In the opposite case [generation of *Fe^{II}(porph^\bullet\bullet\bullet)] different spectral features should have been present in the transient spectra, together with their evolution. Further, if *TrpH\bullet\bullet\bullet is generated, an ESA feature at 560 nm [absorption band of TrpH\bullet\bullet\bullet (66)] should arise with the *Trp\bullet\bullet\bullet decay time; if instead *TrpH\bullet\bullet\bullet is generated, some transient features should appear somewhere in the probing region (note that no information is available on *TrpH\bullet\bullet\bullet absorption bands, but an ESA feature should at least show up in the probed region). However, despite this fact, the transient spectra do not display any ESA feature around 560 nm, suggesting that the molar extinction coefficient of the generated TrpH\bullet\bullet\bullet (or *TrpH\bullet\bullet\bullet) is too small to detect the produced species. This is in line with the results on MbCN and met-Mb, in which the Trp radical cation was not detected either in its ground or in its excited state (40). Of course, if the signals of the products in their excited state fall outside the region of our probe and/or they decay to their ground state on a timescale that is too fast to be measurable with our setup, these considerations are no longer valid.

To estimate the quantum yield (QY) for electron transfer, the deoxy-Mb static spectrum (SI Appendix, Fig. S1) has been rescaled to the GSB amplitude of the LL transient spectrum at 900 ps, allowing a rough estimate of the proportion of MbS with a Fe^{II}(porph^\bullet\bullet\bullet) heme. The obtained value was divided by the total amount of excited Trp\bullet\bullet\bullet residues (details are given in SI Appendix). We found that ~30% of photoexcited Trp\bullet\bullet\bullet relax via electron transfer to the heme whereas the remaining ones relax via FRET. Because there are only these two parallel relaxation mechanisms, we can apply the relationship QY = k_{exo}(2, k_0) to obtain an estimate of the k_{exo} leading to k_{ex} = 1/\tau_{exc} = 1.7 \times 10^{-10} s^{-1} (\tau_{exc} = 60 ps). These values are similar to the ferric MbS, where the QY was found to be ~40% (40), implying k_{ex} = 1/\tau_{exc} = 3.3 \times 10^{-10} s^{-1} (\tau_{exc} = 30 ps). More insights into the *Trp^{14}-to-heme electron transfer reaction in ferric MbS could be obtained using Marcus theory (ref. 67 and SI Appendix, section SII.4). However, too many parameters are unknown in the present case, hindering a deeper analysis of the Marcus region in which this electron transfer occurs.

The reported absence of *Trp fluorescence quenching in apom-
Mb (27) highlights the strong affinity of *Trp to undergo energy and electron transfer to the heme. Given the similarity in yields and timescales for ferric and ferrous MbS, it seems that the *Trp-
to-heme electron transfer is not determined by the oxidation state of the iron ion, and therefore the interaction between *Trp and the porphyrin is the main cause for *Trp deactivation and thus for electron transfer to occur. We thus predict that in the ferric case, the prime acceptor of the electron is the porphyrin, even if at a subsequent stage it develops an Fe^{II} center.

Zhong and coworkers investigated Trp fluorescence quench-
ing in proteins by inter residue and interhelical electron transfer and identified carbonyl- and sulfur-containing residues as quench-
ing groups (24). Although a glutamic acid residue is present in the vicinity of Trp^{14} in Mb, it is oriented toward the solvent, so we do not consider it playing a role. Rather, we believe that a possible pathway for a single-step electron transfer from *Trp^{14} to the heme can involve Leu^{10}, which is in van der Waals contact with Trp^{14} and Val^{60} (SI Appendix, Fig. S7).

Having identified a Trp^{14}-to-heme electron transfer in ferric (40) and now in the ferrous deoxy-Mb and given the fact that the Trp fluorescence lifetimes are invariant in all MbS (SI Appendix, Table S1), we anticipate the process to also be present in ligated ferrous MbS. Hemoglobin (Hb) has six Trp residues: one in each a-subunit (\alpha 14) and two in each \beta-subunit (\beta 15 and \beta 37). Quite remarkably, their fluorescence lifetimes (68, 69) are comparable to the Mb values in SI Appendix, Table S1. As a matter of fact, in deoxy-, oxy-, and carboxy-Hb, the Trp residues are at typical distances of 13–18 Å from a heme porphyrin, which is comparable to the Trp^{14}-heme distance of 15.1 Å in Mb. We predict that a Trp-to-heme electron transfer also occurs in hemoglobins.

An extreme case of Trp fluorescence quenching in heme pro-
teins was found for ferrous and ferric cytochrome c (Cyt c) with, respectively, decay times of 350 fs and 770 fs (57, 58). Trp is at van der Waals distances of the porphyrin (3–5 Å; SI Appendix, Fig. S8) in Cyt c, but the Trp quenching was clearly identified as being due to FRET, at least to 85%. In this case the FRET may well be mediated by an exchange (Dexter) mechanism, in which the donor loses an electron from its excited state that is donated back to its ground state by the acceptor. In the light of these and the present results, it seems that electron transfer can compete with FRET only when the latter is less efficient (due to distance and orientation of the donor/acceptor dipoles) and/or when residues between the Trp and the porphyrin are present that can mediate it. In any case, more studies are needed to fully understand the competition between FRET and electron transfer in hemoproteins.

Conclusions

Femtosecond UV-visible transient absorption experiments were performed on deoxy-Mb for excitation wavelengths near 300 nm. They reveal the formation of a long-lived photoprod uct, which results from a *Trp^{14}-to-heme electron transfer with a quantum yield of ~30%. This species is an Fe^{II}–porphyrin \pi-anion radical that has a lifetime exceeding our measurement window of 1 ns. To our knowledge this is the first observation of such a low-
valent heme complex under physiological conditions, although their existence and biological importance as intermediates in the production pathway of active species in cytochrome P450 (43, as well as for CO2 reduction (44) were discussed earlier.

The similarity to our previous results on fer rous MbS (40) and the invariance of the *Trp lifetimes for all myoglobin suggests that the *Trp^{14}-to-heme electron transfer is likely operative in the ligated ferrous MbS.

We propose a single-step tunneling pathway for the electron transfer that involves the Leu^{10} and Val^{60} residues that lower the tunneling energy. Finally, as previously stressed (40, 70), care is advised when using *Trp fluorescence as a spectroscopic r ule, assuming that its fluorescence decay is due to FRET. This is surely an important tool in studies of protein dynamics but more often than previously thought, parallel electron transfer pathways may also contribute to its quenching.

Associated Content

Sample preparation, optical setup details, power dependence, data analysis details, and extra figures are available in SI Appendix.

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