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### Recommended Citation

Yukl, Erik T.; Liu, Fange; Krzystek, J.; Shin, Sooim; Jensen, Lyndal M. R.; Davidson, Victor L.; Wilmot, Carrie M.; and Liu, Aimin, "Diradical intermediate within the context of tryptophan tryptophylquinone biosynthesis" (2013). *Faculty Bibliography 2010s*. 4914.  
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Source: *Proceedings of the National Academy of Sciences of the United States of America*, Vol. 110, No. 12 (March 19, 2013), pp. 4569–4573

Published by: National Academy of Sciences

Stable URL: <https://www.jstor.org/stable/42583309>

Accessed: 13-12-2018 14:22 UTC

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# Diradical intermediate within the context of tryptophan tryptophylquinone biosynthesis

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Edited by JoAnne Stubbe, Massachusetts Institute of Technology, Cambridge, MA, and approved February 1, 2013 (received for review August 30, 2012)

Despite the importance of tryptophan (Trp) radicals in biology, very few radicals have been trapped and characterized in a physiologically meaningful context. Here we demonstrate that the diheme enzyme MauG uses Trp radical chemistry to catalyze formation of a Trp-derived tryptophan tryptophylquinone cofactor on its substrate protein, premethylamine dehydrogenase. The unusual six-electron oxidation that results in tryptophan tryptophylquinone formation occurs in three discrete two-electron catalytic steps. Here the exact order of these oxidation steps in the processive six-electron biosynthetic reaction is determined, and reaction intermediates are structurally characterized. The intermediates observed in crystal structures are also verified in solution using mass spectrometry. Furthermore, an unprecedented Trp-derived diradical species on premethylamine dehydrogenase, which is an intermediate in the first two-electron step, is characterized using high-frequency and -field electron paramagnetic resonance spectroscopy and UV-visible absorbance spectroscopy. This work defines a unique mechanism for radical-mediated catalysis of a protein substrate, and has broad implications in the areas of applied biocatalysis and understanding of oxidative protein modification during oxidative stress.

cofactor biosynthesis | tryptophan radical | heme | posttranslational modification | electron transfer

Protein-based radicals, particularly on tyrosine (Tyr) and tryptophan (Trp) residues, have been implicated in a large number of catalytic and electron transfer reactions in biology (1), including the long-range electron transfer reactions required for photosynthesis (2), respiration (3), and DNA synthesis (4) and repair (5). Aberrant formation of protein radicals during oxidative stress is also of special importance. Evidence for the involvement of protein-based and substrate-based radicals in enzyme-catalyzed reactions has increased substantially in recent years, and expanded our knowledge of the scope of chemical reactions accessible to enzymes. The ability of enzymes to catalyze what were previously thought to be unattainable reactions is exemplified by MauG. The biosynthesis of the tryptophan tryptophylquinone (TTQ) cofactor in methylamine dehydrogenase (MADH) requires posttranslational modification of two tryptophan residues. MADH from *Paracoccus denitrificans* is a 119-kDa  $\alpha_2\beta_2$  heterotetramer that contains two active sites and two TTQ cofactors derived from residues  $\beta$ Trp57 and  $\beta$ Trp108 (6). MauG is a *c*-type diheme enzyme that catalyzes the conversion of a MADH precursor (preMADH) that has one oxygen atom already inserted into the  $\beta$ Trp57 indole ring ( $\beta$ Trp57-OH of preTTQ) (7), to mature TTQ-containing MADH (8) (Fig. 1).

Catalysis by MauG proceeds via a *bis*-Fe(IV) redox state, which may be generated by reaction of di-Fe(II) MauG with O<sub>2</sub> or di-Fe(III) MauG with H<sub>2</sub>O<sub>2</sub> (9). Catalytically active crystals of the MauG-preMADH protein complex show that the site of TTQ formation on  $\beta$ -preMADH is 40.1 Å from the MauG high-spin heme iron that reacts with H<sub>2</sub>O<sub>2</sub> or O<sub>2</sub> (Fig. S1A), demonstrating that catalysis involves long-range electron transfer (10). The overall reaction is a six-electron oxidation, and thus

three cycles of MauG *bis*-Fe(IV) formation are required to complete TTQ biosynthesis. The three two-electron steps must equate to (i) formation of the cross-link between  $\beta$ Trp57-OH and  $\beta$ Trp108, (ii) insertion of the second oxygen atom into  $\beta$ Trp57-OH, and (iii) oxidation of the quinol to the quinone. The oxidation of the cross-linked quinol to quinone is believed to be the final step in the reaction (11). Here the order of the first two steps of the reaction is resolved using X-ray crystallography and mass spectrometry, with high-frequency and -field electron paramagnetic resonance (HFEP) measurements providing details of an unprecedented radical mechanism for the first step. This study fills gaps in the MauG reaction mechanism, and leads to a better understanding of the remarkable chemistry that occurs in this system.

## Results

It was previously shown that addition of excess H<sub>2</sub>O<sub>2</sub> to crystals of the MauG-preMADH complex resulted in formation of TTQ *in crystallo* (10). X-ray crystallography cannot resolve quinol to quinone oxidation, but can distinguish cross-link formation from oxygen insertion. To determine the order of these events in MADH maturation, the residual peroxide contaminants within commercial PEG were exploited. PEG solutions slowly accumulate organic peroxides through autooxidation of the polyoxyethylene bonds, and can release hydrogen peroxide as part of this process (12). The MauG-preMADH crystals were grown using 22–26% (wt/vol) PEG 8000 as precipitant, and thus as the crystals aged within the crystallization trays they were exposed to the slow release of H<sub>2</sub>O<sub>2</sub>. As a result, catalytic turnover in MauG-preMADH crystals was observed over time. X-ray diffraction data from crystals harvested after 10, 20, 30, 40, 50, and 130 d clearly show a progression of changes at the preTTQ site that is consistent with the cross-link forming first, followed by addition of the second oxygen to  $\beta$ Trp57-OH (Fig. 2A; Table S1 and S2). Electron density changes indicate cross-link formation between  $\beta$ Trp57-OH and  $\beta$ Trp108 is complete after ~40 d, and requires  $\beta$ Trp57-OH to undergo a rotation of ~20° (Fig. 2B). Significant positive-difference electron density appears for the second O atom of TTQ only after cross-linking has occurred.

To ensure these events were indeed MauG-dependent reactions and not simply due to direct reaction with H<sub>2</sub>O<sub>2</sub>, a crystal of

Author contributions: L.M.R.J., V.L.D., C.M.W., and A.L. designed research; E.T.Y., F.L., J.K., and S.S. performed research; E.T.Y., F.L., J.K., S.S., V.L.D., C.M.W., and A.L. analyzed data; and E.T.Y., F.L., V.L.D., C.M.W., and A.L. wrote the paper.

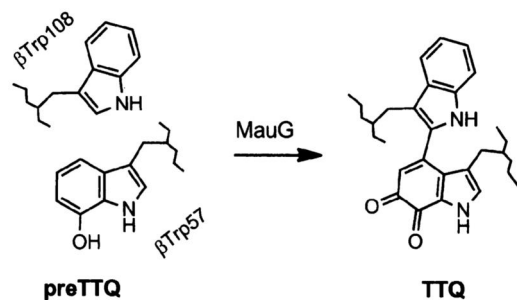
The authors declare no conflict of interest.

This article is a PNAS Direct Submission.

Data deposition: The atomic coordinates and structure factors have been deposited in the Protein Data Bank, www.pdb.org (PDB ID codes 4FA1, 4FA4, 4FA5, 4FA9, 4FAN, and 4FAV).

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This article contains supporting information online at www.pnas.org/lookup/suppl/doi:10.1073/pnas.1215011110/-DCSupplemental.



**Fig. 1.** Chemical structure of the MADH-based substrate and product of MauG catalysis.

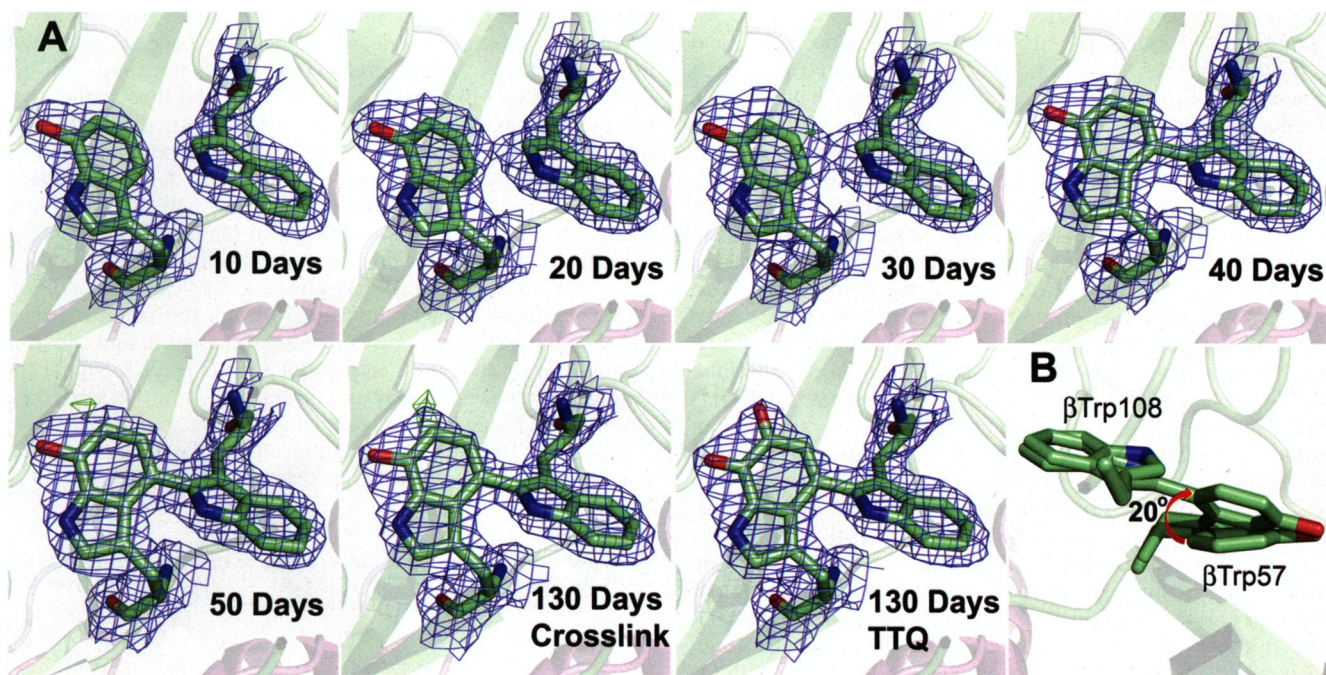
the complex between inactive W199F MauG and preMADH was aged. The W199F MauG mutant can form the *bis*-Fe(IV) state on reaction with H<sub>2</sub>O<sub>2</sub>, but cannot catalyze TTQ biosynthesis, as hole hopping through Trp199 is required for preTTQ oxidation (Fig. S14) (13, 14). In contrast to the native MauG–preMADH crystals, the W199F MauG–preMADH crystals showed no changes at the preTTQ site after 60 d (Fig. S1B; Table S3). Therefore, the sequential reactions of preMADH observed in the crystals are MauG dependent.

To demonstrate that the *in crystallo* results reflect the order of the reaction in solution, whole-protein mass spectrometry of MauG, preMADH, and H<sub>2</sub>O<sub>2</sub> in a 2:1:2 stoichiometry was compared with a control sample without H<sub>2</sub>O<sub>2</sub>; this equates to one MauG:one preTTQ:one H<sub>2</sub>O<sub>2</sub>, because preMADH contains two active sites. Reverse-phase chromatography on a C4 column allows for the separation of the  $\beta$ -subunit of MADH from both MauG and  $\alpha$ -MADH before mass spectrometry (Fig. S2). Comparison of the chromatogram of the MauG–preMADH

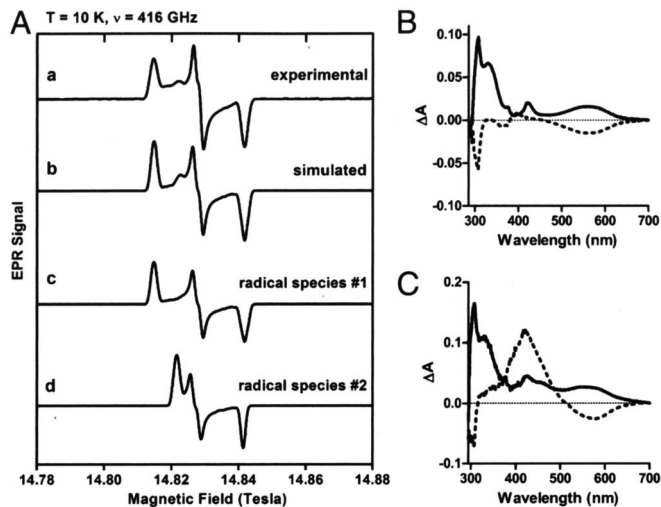
sample to the H<sub>2</sub>O<sub>2</sub> treated sample shows the appearance of a new peak eluting earlier than  $\beta$ -preMADH (Fig. S2A). Mass spectrometry demonstrated the presence of two distinct species within this peak (Fig. S2B). The majority species is consistent with the mass of TTQ-containing  $\beta$ -MADH (15,009.73  $\pm$  0.20 Da), and the assignment is confirmed by this chromatographic fraction having an absorbance maximum at 440 nm (Fig. S2D) (15). The other species present is 2-Da lighter (14,995.60  $\pm$  0.05 Da) than  $\beta$ -preMADH in the control (14,997.54  $\pm$  0.02 Da) and H<sub>2</sub>O<sub>2</sub>-treated samples (14,997.32  $\pm$  0.26 Da; Fig. S2B), which is consistent with a  $\beta$ -MADH species containing the cross-link between  $\beta$ Trp57-OH and  $\beta$ Trp108 in the absence of the second oxygen atom. None of the chromatographic peaks from the H<sub>2</sub>O<sub>2</sub>-treated sample contained a species with a mass consistent with either quinol or quinone, but no cross-link (Table S4); this confirms that cross-link formation occurs first, followed by insertion of a second hydroxyl group into  $\beta$ Trp57-OH, and finally oxidation to the quinone.

Previous work demonstrated that when *bis*-Fe(IV) MauG was treated with stoichiometric preMADH, a protein-based radical was observed by electron paramagnetic resonance spectroscopy (9). To characterize this radical species in the context of TTQ formation, X-band EPR and HFEPR were used. After mixing MauG, preMADH and H<sub>2</sub>O<sub>2</sub> in a 1:1:1 ratio, regardless of the order of addition, the diferric EPR signal of MauG returns, and a radical intermediate is detectable with a spin quantitation of 1.4–1.6 equivalents by X-band EPR, suggesting that two electrons were transferred to the *bis*-Fe(IV) site resulting in formation of two protein-based radicals.

Trp and monohydroxylated Trp radicals, as well as other aromatic radicals such as Tyr, are indistinguishable by conventional X-band EPR spectroscopy (frequency  $\sim$ 9 GHz). Therefore, HFEPR at 108–416 GHz and correspondingly high fields (up to 15 T) was used to determine the nature of the radical intermediate



**Fig. 2.** (A) Electron density for preTTQ site in MauG–preMADH crystals of different ages. Electron density maps for the crystals aged for 10, 40, 50 and 130 d were generated from the respective refined structures. The 20- and 30-d electron density maps were calculated using the refined 10-d model (preTTQ) to enable visualization of the development of cross-link density. The two final refined models for the 20- and 30-d data contain a mix of preTTQ and cross-linked preTTQ (Tables S1 and S2). The 130-d cross-link electron density image was calculated using the refined 50-d model (cross-linked preTTQ) to enable visualization of the positive-difference electron density for the second oxygen. The final refined model at 130 d contains 100% TTQ. 2Fo–Fc, blue, contoured at 1.0 $\sigma$ ; Fo–Fc, green, contoured at +4 $\sigma$ . (B) The observed rotation in  $\beta$ Trp57-OH during cross-link formation. The figure was produced using PyMOL ([www.pymol.org](http://www.pymol.org)).



**Fig. 3.** (A) 10 K HFEPR spectrum at 416 GHz and correspondingly high field (14–15 Tesla) of the preMADH diradical intermediate (signal a). Spectral simulation (signal b) shows two free radical species, one with  $g$  tensors of 2.00216, 2.00398, 2.00581, and  $\Delta g = 0.00365$  (signal c) and the other 2.00216, 2.00402, 2.00486, and  $\Delta g = 0.00270$  (signal d). The final simulated spectrum presents a decomposition of 85% radical species 1, and 15% radical species 2. Absorbance difference spectra of the diradical intermediate of preMADH obtained under single turnover (B) and pre-steady state (C) conditions. Solid line is the difference of spectra taken after addition of  $H_2O_2$ —before addition. Broken line is the difference of spectra taken after addition of hydroxyurea—after  $H_2O_2$  addition.

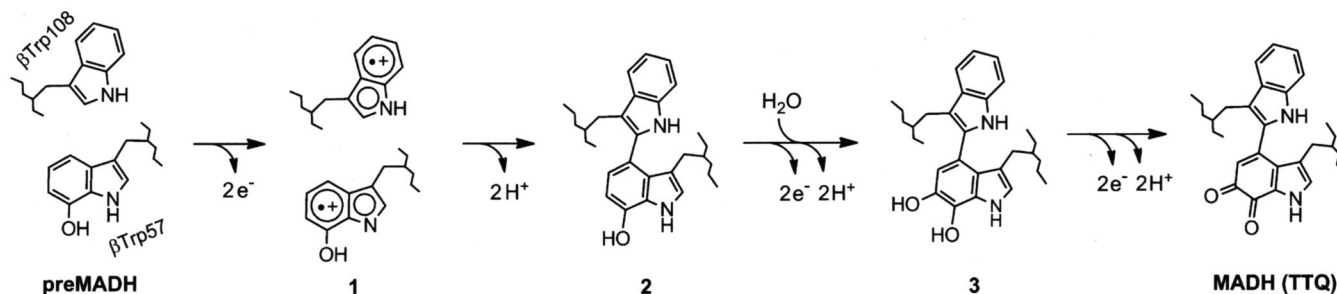
preceding cross-link formation in the reaction of preMADH with *bis*-Fe(IV) MauG (Fig. S3). Trp and Tyr radicals are differentiable by HFEPR because oxygen substitution on the phenyl ring of tyrosine significantly increases the  $g$ -tensor anisotropy ( $\Delta g = |g_x - g_z|$ ) (16). Specifically, Tyr radicals exhibit a much larger  $g$ -anisotropy ( $\Delta g > 5 \times 10^{-3}$  for isolated radicals) than Trp radicals ( $\Delta g < 1.5 \times 10^{-3}$ , Table S5) (16–21). A similar  $g$ -tensor anisotropy effect would be expected for a hydroxylated Trp radical because the radical spin density is typically delocalized over the entire indole ring (22–27). Therefore,  $\beta$ Trp57-OH of preMADH would be expected to have a  $\Delta g$  value greater than that of Trp radicals and closer to that of Tyr radicals. Fig. 3A describes a diradical intermediate composed of two distinct radicals at 416 GHz ( $T = 10$  K). A single radical species cannot account for all of the resonances of the HFEPR data (Fig. S4). One radical species (species 1) exhibits principal  $g$  values of  $g_x = 2.00216$ ,  $g_y = 2.00398$ ,  $g_z = 2.00581$ , and  $\Delta g = 0.00365$ ; the other (species 2) has  $g$  tensors of  $g_x = 2.00216$ ,  $g_y = 2.00402$ ,  $g_z = 2.00486$ , and  $\Delta g = 0.00270$ . Radical species 2 is consistent with a Trp radical, whereas species 1 is consistent with the anticipated  $\beta$ Trp57-OH radical. The presence of two distinct radical species is further

verified by HFEPR data measured at a higher temperature (Fig. S5). In the HFEPR spectrum of the frozen solution, the two radicals are not in a 1:1 ratio. The solution kinetics associated with the two-electron oxidation of preMADH ( $k_{lim} = 0.8 \text{ s}^{-1}$ ) shows that full diradical formation will occur within the dead time of mixing the HFEPR sample (28). Therefore, any inequality must arise from differing decay rates or stabilities of the two radicals; this is most likely due to the two radicals having different stabilities during the freezing process. The pH of common biological buffers is temperature dependent (29–31). Even a modest change in pH could differentially affect the stability of one of the radicals. A similar mismatched ratio has been previously observed in the *bis*-Fe(IV) intermediate of MauG, likely for the same reason (9). It is also possible that the radicals may have different power saturation properties, which cannot be tested due to technical limitations of the HFEPR technique.

Evidence for the Trp-based diradical species was also obtained by monitoring solution single-turnover and steady-state kinetic reactions of MauG and preMADH by UV-visible absorption spectroscopy. When MauG plus preMADH were mixed with stoichiometric  $H_2O_2$  for 30 s, the formation of absorption features at 560 and 310 nm characteristic of Trp-based cation radicals in solution (32) and in protein (33) were observed (Fig. 3B). Subsequent addition of hydroxyurea, a radical scavenger, quenched these features, confirming that they represent radical species. The shoulder at 330 nm, which was not sensitive to hydroxyurea, is likely a small amount of quinol MADH (15). The reaction was also studied with excess  $H_2O_2$  present to generate steady-state conditions (Fig. 3C). Conditions were such that after 36 s the reaction was still in the pre-steady-state phase. At this time, the 560- and 310-nm absorption features were again observed, and subsequent addition of hydroxyurea quenched these features. Appearance of an absorption feature at 440 nm following hydroxyurea treatment is likely a small amount of quinone MADH that formed during the  $\sim 10$ -s interval between recording the initial difference spectrum and addition of the di-Trp radical in solution on a time scale that parallels the mixing/freezing time to prepare the EPR samples. The pre-steady-state Trp-based cation radical spectral features disappear with the appearance of the 440-nm absorbance that indicates TTQ formation, and confirms the relevance of the intermediate to the catalytic cycle.

**Discussion**

The radical intermediate associated with the first two-electron oxidation of preMADH was characterized by HFEPR, which identified two distinct radical species; one of these is consistent with a preMADH  $\beta$ Trp57-OH radical, whereas the other appears to be a Trp radical. The characterization of this radical intermediate, and the order of the reaction *in crystallo*, allows a working chemical reaction mechanism to be proposed for



**Fig. 4.** Proposed chemical reaction mechanism for the six-electron oxidation of preMADH leading to the formation of TTQ and mature MADH.

TTQ biosynthesis (Fig. 4). In this mechanism, the first two oxidizing equivalents from the *bis*-Fe(IV) intermediate oxidize preMADH to produce an unprecedented Trp-based diradical (1), which subsequently loses two protons and combines to form a cross-link (2). The second cycle of oxidation by a regenerated *bis*-Fe(IV) MauG incurs an attack by a water molecule, leading to oxygen insertion, as defined by X-ray crystallography, to form the quinol MADH (3). The final *bis*-Fe(IV) MauG catalyzes oxidation of reduced quinol TTQ to the oxidized and catalytically active TTQ cofactor. Although a  $\beta$ Trp108 radical makes sense from a mechanistic standpoint, the possibility that the second Trp radical may be localized on another Trp residue cannot be excluded.

The ability to resolve the *g*-values for the two radicals by HFEPR signifies that there is little or no magnetic coupling between them; this may have various origins, such as geometric factors, the specific nature of coupling (through space or via bonds), and the fact that both radicals are indicated by absorbance data to be cationic. We know through theoretical work that the spin distribution on a cation radical (of 3-methylindole) is different from that on a neutral radical (34), which may also affect the magnetic coupling between them.

Generally, for the cross-linking reaction between the cation radicals to proceed, they need to become deprotonated and couple. These processes are likely interconnected and essentially concurrent. Although the precise mechanism of deprotonation remains to be elucidated, it is known to be a spontaneous process for Trp-based protein radicals (20). It should be noted that the Fe ions in MauG would not affect the coupling within the diradical, analogous to the situation described for the R2 protein of *Escherichia coli* ribonucleotide reductase (33). The nearest heme Fe in MauG is 19 Å away from  $\beta$ Trp108 and  $\beta$ Trp57-OH, which is much longer than the Trp radical and di-iron center separation in R2. A hydrogen bond connection postulated to play a role in R2 is also absent in the MauG–preMADH complex. Rather, the electron transfer from preMADH to MauG has been shown to occur through hole-hopping (13, 14).

The present study begins to unravel the remarkable chemistry that occurs during this biosynthetic reaction. The occurrence and stability of the di-Trp radical intermediate is unprecedented. The longevity may be a consequence of the required rotation of Trp57-OH to react with Trp108 (Fig. 2B), or a rate-limiting proton loss that precedes radical coupling. The nature of the proton acceptors is as yet unknown. The role of the protein environment surrounding the two Trp residues in preMADH clearly plays an important role in facilitating the reaction; this may be considered an unusual form of substrate-assisted catalysis coupled with long-range electron transfer to the MauG high-valent hemes.

## Methods

Detailed methods can be found in *SI Methods*.

**Protein Expression and Purification.** Recombinant MauG (35) was purified from *P. denitrificans* as described previously. PreMADH (7) was expressed in *Rhodobacter sphaeroides* and purified as described previously (36).

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**UV-Visible Absorbance Spectroscopy.** The single-turnover reactions contained 10  $\mu$ M preMADH, 1.0  $\mu$ M MauG, and 10  $\mu$ M H<sub>2</sub>O<sub>2</sub>. Hydroxyurea was then added at a concentration of 2.0 mM. The pre-steady-state reaction contained 30  $\mu$ M preMADH, 3.0  $\mu$ M MauG, and 1.0 mM H<sub>2</sub>O<sub>2</sub>. Hydroxyurea was added to 3.0 mM.

**EPR Spectroscopy.** The HFEPR sample was prepared in 10 mM potassium phosphate buffer (pH 7.5) containing 5–10% (vol/vol) glycerol by first adding MauG (135  $\mu$ L, 1.43 mM) to preMADH (170  $\mu$ L, 1.17 mM). Then, H<sub>2</sub>O<sub>2</sub> (final concentration = 1.22 mM) was added and the mixture frozen. HFEPR spectra were recorded at the EMR Facility at the National High Magnetic Field Laboratory in Tallahassee, FL (37). HFEPR simulations were performed using DOUBLET. The samples for continuous-wave X-band EPR quantitative analyses had final protein concentrations of 150  $\mu$ M for both MauG and preMADH with addition of 1 equivalent of H<sub>2</sub>O<sub>2</sub>. The X-band EPR spectra were recorded on a Bruker ER200D spectrometer at 100-kHz modulation frequency using a 4116DM resonator.

**Mass Spectrometry.** Reaction mixtures contained 25–55  $\mu$ L of 20  $\mu$ M preMADH, 40  $\mu$ M MauG, and 0–120  $\mu$ M H<sub>2</sub>O<sub>2</sub>. Samples were incubated for 1 h before injection onto a C4 column (Phenomenex) connected to a HPLC system (Waters). Fractions were collected, dried, and reconstituted in 50% acetonitrile, 0.1% formic acid for mass spectrometry. Data were acquired on a QSTAR XL (AB Sciex) quadrupole TOF mass spectrometer with the IonSpray electrospray source.

**Crystallization, X-Ray Data Collection, and Structure Determination.** The WT MauG and W199F MauG complexes with preMADH were crystallized as previously reported (10, 14). WT MauG–preMADH crystals were harvested every 10 d for 50 d, and once at 130 d following crystallization tray setup. The W199F MauG–preMADH crystal was harvested after 60 d. X-ray diffraction data were collected at the Advanced Photon Source (APS), Argonne National Laboratory, Argonne, IL. The data were processed with HKL2000 (38) (Tables S1–S3). Refinement was carried out using REFMAC (39) in the CCP4 program suite (40) starting from the model of WT MauG–preMADH (PDB ID code 3L4M). Model-building was carried out in COOT (41, 42), and calculated electron density maps were generated using PHENIX (43) (Tables S1–S3).

**ACKNOWLEDGMENTS.** We thank Ed Hoeffner for KSBL support; LeeAnn Higgins and Bruce Witthun for CMSP support; and the staff at APS sector 23 for their support, especially Michael Becker, Steve Corcoran, Venugopalan Nagarajan, and Derek Yoder; and Dr. Andrew Ozarowski of the National High Magnetic Field Laboratory for providing the EPR simulation program used in this study. HFEPR studies were supported by the National High Magnetic Field Laboratory, which is funded by the National Science Foundation (NSF) through Cooperative Agreement DMR 1157490, the State of Florida, and the Department of Energy. This work was supported by National Institutes of Health Grants GM-66569 (to C.M.W.), GM-41574 (to V.L.D.), GM-97779 (to E.T.Y.), NSF Grant MCB-0843537 (to A.L.), the Georgia Cancer Coalition Distinguished Scholar Program (A.L.), and Minnesota Partnership for Biotechnology and Medical Genomics Grant SPAP-05-0013-P-FY06 (to C.M.W.). Mass spectrometry was conducted at the Center for Mass Spectrometry and Proteomics at the University of Minnesota. Computer resources were provided by the Basic Sciences Computing Laboratory of the University of Minnesota Supercomputing Institute. X-ray data were collected at the Kahlert Structural Biology Laboratory at the University of Minnesota and the National Institute of General Medical Sciences and the National Institute of Cancer Institute Structural Biology Facility–Collaborative Access Team (GM/CA-CAT) at the APS, Argonne National Laboratory. GM/CA-CAT has been funded in whole or in part by National Cancer Institute Grant Y1-CO-1020 and National Institute of General Medical Science Grant Y1-GM-1104. Use of the APS was supported by the US Department of Energy, Basic Energy Sciences, Office of Science Contract DE-AC02-06CH11357.

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