

Imino-oxy Acetic Acid Dealkylation as Evidence for an Inner-Sphere Alcohol Intermediate in the Reaction Catalyzed by Peptidylglycine α -Hydroxylating Monooxygenase

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Abstract: Peptidylglycine α -hydroxylating monooxygenase (PHM, EC 1.14.17.3) catalyzes the stereospecific hydroxylation of a glycol α -carbon in a reaction that requires O₂ and ascorbate. Subsequent dealkylation of the α -hydroxyglycine by another enzyme, peptidylamidoglycolate lyase (PAL, EC 4.3.2.5), yields a bioactive amide and glyoxylate. PHM is a noncoupled, type II dicopper monooxygenase which activates O₂ at only a single copper atom, Cu_M. In this study, the PHM mechanism was probed using a non-natural substrate, benzaldehyde imino-oxy acetic acid (BIAA). PHM catalyzes the O-oxidative dealkylation of BIAA to benzaldoxime and glyoxylate *with no involvement of PAL*. The minimal kinetic mechanism for BIAA was shown to be steady-state ordered using primary deuterium kinetic isotope effects. The $D(V/K)_{\text{APPARENT, BIAA}}$ decreased from 14.7 ± 1.0 as $[O_2] \rightarrow 0$ to 1.0 ± 0.2 as $[O_2] \rightarrow \infty$ suggesting the dissociation rate constant from the PHM·BIAA complex decreases as $[O_2]$ increases; thereby, reducing the steady-state concentration of $[PHM]_{\text{free}}$. BIAA was further used to differentiate between potential oxidative Cu/O species using a QM/MM reaction coordinate simulation to determine which species could yield product O-dealkylation that matched our experimental data. The results of this study provided compelling evidence for the presence of a covalently linked Cu^{II}-alkoxide intermediate with a quartet spin state responsible BIAA oxidation.

Introduction

Peptidylglycine α -amidating monooxygenase (PAM, E.C. 1.14.17.3) catalyzes the final reaction in the biosynthesis of α -amidated peptide hormones and may also have a role in the production of oleamide and other bioactive primary fatty acid amides.^{1,2} This chemistry is a two-step process with the formation of an intermediate that has been stereospecifically hydroxylated at the α -carbon of a glycine. PAM is bifunctional consisting of two catalytic units: peptidylglycine α -hydroxylating monooxygenase (PHM) and peptidylamidoglycolate lyase (PAL). PHM catalyzes the glycine hydroxylation while PAL catalyzes the carbinolamide dealkylation (Scheme 1a). The relationship between PHM and PAL is multifaceted and species specific. Mammalian PAM is coded for by a single gene; alternative splicing of the PAM mRNA and proteolytic processing of PAM yields a mixture of the bifunctional enzyme and the monofunctional enzymes, PHM and PAL. In insects and cnidarians, bifunctional PAM is not found; only monofunctional

PHM and PAL are known, each coded by a distinct gene.^{3–5} In the mollusk, *Lymnae stagnalis*, a pentafunctional enzyme is produced with four PHM domains per PAL domain.⁶ The complexities in the various forms of the PAM/PHM/PAL system are not well understood, but hint at a novel regulatory mechanism for the *in vivo* production of the amide and carbinolamide products of this chemistry.

Of the two monofunctional enzymes in the PAM/PHM/PAL system, PHM has been the most extensively studied, due to its structural and mechanistic similarity to dopamine.

β -monooxygenase (D β M).⁷ Both PHM and D β M catalyze an O₂-, copper-, and ascorbate-dependent hydrogen atom transfer and substrate hydroxylation resulting in the stereospecific hydroxylation of their respective substrates (Scheme 1a and 1b). Both enzymes contain a solvent filled active site separating two essential, noncoupled copper atoms.^{1,8–12} In PHM, the two protein bound copper atoms have different roles in catalysis. One copper atom, Cu_M, has two N ϵ -histidines ligands (His₂₄₀

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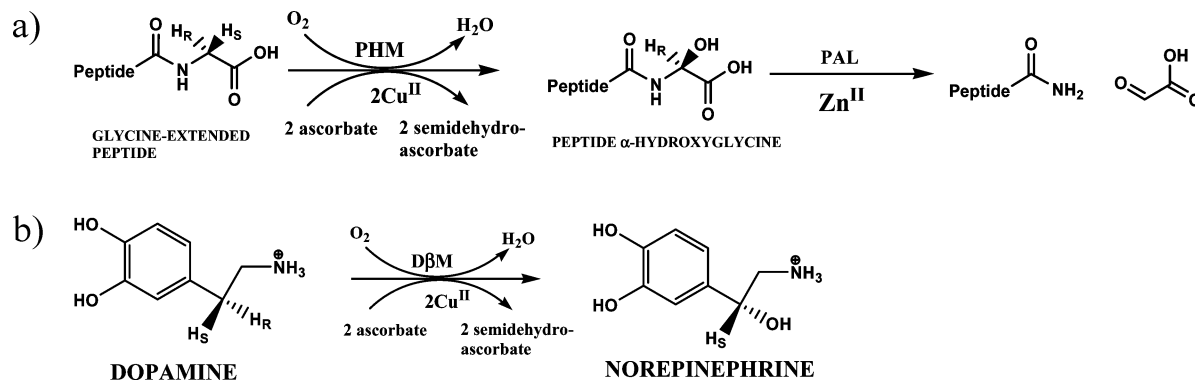
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Scheme 1. (a) Peptidylglycine α -Amidating Monooxygenase (PAM) and (b) Dopamine β -Monooxygenase (D β M) reactions

and His₂₄₂) and a methionine sulfur ligand (Met₃₁₄) and is directly involved in dioxygen activation and substrate oxidation. The other copper atom, Cu_H, has three N δ -histidine ligands (His₁₀₇, His₁₀₈, and His₁₇₂) and is involved in electron transfer. The PAL domain, unique to the PAM system, is zinc- and calcium-dependent and catalyzes the dealkylation of the carbinolamide to the corresponding amide and glyoxylate (Scheme 1a).^{13,14}

Intriguingly, D β M, without a PAL-like partner, will catalyze oxidative dealkylation similar to the sequential reactions catalyzed by PHM and PAL in bifunctional PAM.¹⁵ We report herein the PAL-independent oxidative dealkylation of iminoxyacetic acids to the corresponding oximes and glyoxylate by PHM. For PHM and D β M, the dealkylation reactions result solely from the oxidation chemistry of their monooxygenase domains. This similarity in dealkylation chemistry provides a novel framework to study the mechanism of PAM catalysis.^{15–17}

The role of oxygen activation in PHM and D β M has been the subject of much study.^{18–22} The presumed nucleophile responsible for hydrogen abstraction is predominantly considered to be an end-on/ η^1 copper-superoxo radical species.^{11,23} Both η^1 and η^2 copper-superoxo species had been prepared in models, although none exhibited the propensity for both H-abstraction and oxidation chemistry.^{24–29} Recently, the Karlin group prepared end-on/ η^1 Cu^{II}-superoxo model complexes able to orchestrate both H-abstraction and oxidation chemistry.^{30–33} Characterization of Cu^{II}-superoxo species was a critical advance; however, ambiguities remain as to the identity of the actual oxidant species.³⁰

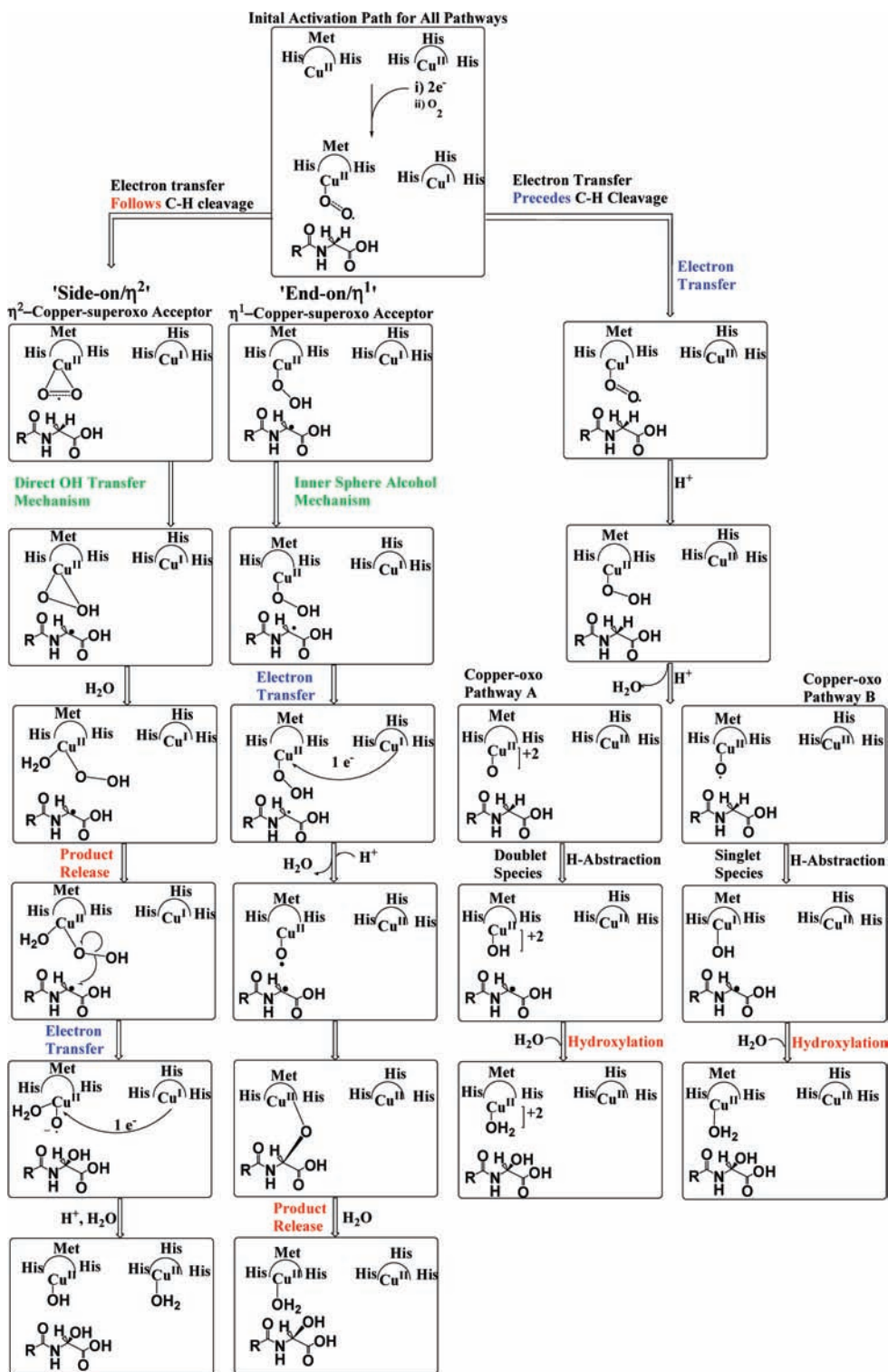
The two PHM mechanisms which involve a Cu^{II}-superoxo nucleophile for substrate activation differ in the initial coordination geometry of the dioxygen species to copper (Scheme 2). The ‘side-on/ η^2 ’ mechanism is supported by spectroscopic evidence from side-on/ η^2 Cu/O species model studies and is predicted to have an antiferromagnetically coupled singlet ground state (see Scheme 3).^{25,34} When compared with the end-on/ η^1 species, the side-on/ η^2 copper^{II}-superoxo was calculated to be thermodynamically preferred for H-transfer.^{19,35} Evidence for the end-on/ η^1 species comes from the PHM crystal structure¹¹ and the work of Blackburn et al.,³⁶ suggesting that this species is the nucleophile responsible for C α –H abstraction. Both mechanisms include an end-on/ η^1 Cu^{II}-hydroperoxo species following C α –H bond cleavage. The ‘side-on/ η^2 ’ mechanism involves a direct hydroxylation of the C α -substrate radical, through a ‘water-assisted’ radical recombination reaction resulting in the simultaneous reduction of the Cu^{II}-hydroperoxo and C α -OH product release. Conversely, the Cu^{II}-hydroperoxo species from the ‘end-on/ η^1 ’ mechanism is reduced via an

intramolecular electron transfer from the other Cu atom in the active site, Cu_H, yielding a CuII-O• radical. Radical recombination of the substrate and Cu-O radical species produces an inner-sphere alcohol intermediate. Subsequent hydrolysis of the inner-sphere alcohol generates the hydroxylated product.

A highly reduced copper-oxo species is postulated for substrate C α –H bond cleavage in both ‘copper-oxo’ mecha-

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Scheme 2. Collection of Mechanisms Postulated for the Hydrogen Abstracting and Oxidation Reactions of Peptidylglycine α -Amidating Monooxygenase (and Dopamine β -Monooxygenase), Respectively

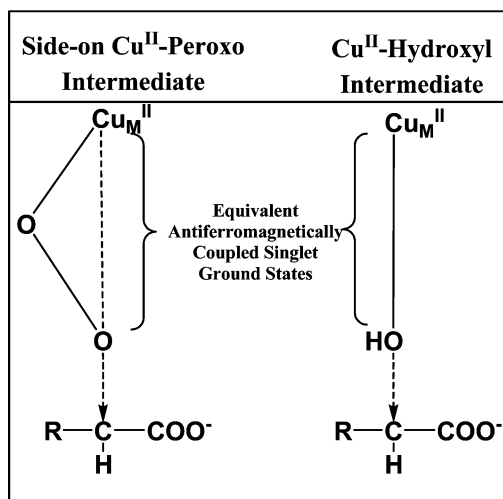


nisms, which uncouple dioxygen reduction from substrate activation. Hybrid QM/MM simulations suggest that a Cu^{III} -oxide/ Cu^{II} -oxyl species is thermodynamically preferred relative to Cu^{II} -superoxo or Cu^{II} -hydroperoxo nucleophiles.^{37–39} The Cu^{III} -oxide/ Cu^{II} -oxyl species is analogous to the ferryl oxidant (CpI) responsible for C–H activation in cytochrome P450.^{37,40} Both 'copper-oxo' mechanisms proceed from an initial end-on/ η^1 Cu^{II} -superoxo species to a highly reduced copper-oxo species by coupling intramolecular electron transfer with the acquisition

of two protons leading to water release. In 'copper-oxo pathway A', the reduced copper-oxo species is Cu^{II} -oxyl with two unpaired electrons ferromagnetically coupled to an unpaired electron delocalized within the Cu_M domain yielding a quartet spin state.³⁷ Here, H-abstraction and hydroxylation are in concert with spin inversion to a doublet ground state allowing substrate

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Scheme 3. Degenerate Electronic Ground-State Structures of the Cu^{II} -Hydroperoxo and Cu^{II} -Hydroxyl Oxidant Species Proposed for Direct Substrate Oxidation/Product Release^a



^a Please note, copper domain residues were omitted for clarity. Refer to text for more detail.

oxidation and hydroxylated product release to be directly coupled. In the ‘copper-oxo pathway B’, the Cu^{II} -oxyl species is in a triplet ground state as the most thermodynamically favorable species for H-abstraction. Spin inversion yields an antiferromagnetically coupled singlet state to drive concerted H-abstraction with substrate oxidation/product release. Simulations by Crespo et al. favor the singlet state oxidant relative to either quartet or triplet state oxidants.³⁷

Analysis of the four mechanisms outlined in Scheme 2 shows commonality between the ‘side-on/ η^2 ’,^{35,41} and both ‘copper-oxo’,^{38,39} mechanisms. Product release and substrate oxidation are simultaneous, lacking covalent interaction between H-donor and Cu/O acceptor (‘free-product’). The *direct* hydroxyl transfer reactions of the ‘side-on/ η^2 ’ and ‘copper-oxo pathway B’ mechanisms are electronically similar as hydroxyl transfer begins with an antiferromagnetically coupled singlet state. Therefore, as the reaction coordinate proceeds toward substrate oxidation/product release, the transition states of both ‘copper-oxo pathway B’ and ‘side-on/ η^2 ’ species become superimposable.

Conversely, the ‘end-on/ η^1 ’ mechanism is defined by an inner-sphere alcohol intermediate produced through a radical recombination reaction.^{7,20} The Cu^{II} -oxyl radical species in the ‘end-on/ η^1 ’ mechanism is unique in that the reductive cleavage of O–O bond in the Cu^{II} -hydroperoxo moiety results from an intramolecular reduction followed by homolysis to yield the active species. Unlike the ferryl ($\text{Fe}^{\text{IV}}=\text{O}$; CpI) complex of P450, the Cu^{II} -oxyl lacks the strong electron delocalization into the *d*-orbitals of the metal, allowing full radical stabilization in the oxygen $2\pi^*$ orbital.⁴¹ Formation of the Cu^{II} -oxyl species by reduction of the Cu^{II} -hydroperoxo followed by homolytic O–O bond cleavage suggests that this cupryl species must exist as a quartet, with well-defined radical character on the oxygen.

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The hypothesis of the present study was that the PHM catalyzed reaction proceeds through an ‘inner-sphere alcohol’ intermediate. To test this hypothesis we designed a novel PHM substrate, benzaldehyde imino-oxy acetic acid (BIAA), which would undergo PAL-independent dealkylation. Compared to other known PHM substrates that release a hydroxylated product,¹ the unique ability of BIAA to decompose during oxygen insertion allows isolation of the substrate oxidation step. Characterization of BIAA dealkylation demonstrated that this chemistry is exclusively dependent on PHM-catalyzed oxidation (and independent of product release), allowing hybrid QM/MM simulations of substrate radical oxidation to be accurately modeled. Our data provides compelling evidence that the PHM reaction does, in fact, proceed via an inner-sphere alcohol intermediate.

Materials and Methods

Materials. Benzamide, α -hydroxyhippurate, benzaldoxime (BOX), bromoacetic acid, [α -²H₂]-bromoacetic acid, hydroxylamine hydrochloride, benzaldehyde, and rabbit muscle lactate dehydrogenase were purchased from Sigma-Aldrich. Carboxymethylhydroxylamine was purchased from TCI. Bovine liver catalase was purchased from Worthington. *N*-dansyl-Tyr-Val-Gly was purchased from Fluka Biochimika. Recombinant type A rat medullary thyroid carcinoma bifunctional PAM was produced and purified as described⁴² and was a gift from Unigene Laboratories, Inc. (Fairfield, NJ, see www.unigene.com). All other reagents were of the highest quality available from commercial suppliers.

Synthesis of Benzaldehyde Imino-oxy Acetic Acid (BIAA). Benzaldehyde was converted to the corresponding benzaldoxime using a published procedure.⁴³ Oxime (4.1 mmol) was dissolved in 20 mL of H₂O containing 5 equiv of NaOH, and the reaction stirred at room temperature for 45 min, after which 1.5 equiv of [α -²H]- or [α -²H₂]-bromoacetic acid was slowly added in small increments with continued stirring. Upon completion (as determined by TLC using a 1:3 hexanes/ethyl acetate as the mobile phase) the reaction was acidified with dilute HCl, the BIAA was collected in a Büchner funnel, and solvent was removed by washing the solid with petroleum ether. BIAA was recrystallized twice with benzene/petroleum ether, yielding a white crystalline solid. Compound purity was analyzed by RP-HPLC and ¹H and ¹³C NMR.

[α -H₂]-BIAA: ¹H NMR analysis (250 MHz, MeOD-*d*₄) δ 4.96 (singlet, 2H, CH₂, α -methylene), 7.66–7.87 (m, 5H, ArH), 8.48 (singlet, 1H, H–C=N). ¹³C NMR analysis (62.5 MHz, MeOD-*d*₄) δ 171.841 (C=O, carboxylic acid), 149.486 (C=N, imine), 130.911 (Ar, C-1), 129.188 (Ar, C-4), 127.673 (Ar, C-3, C-5), δ 126.136 (Ar, C-2, C-6), δ 69.325 (CH₂, α -methylene), mp 93–94 °C.

[α -²H₂]-BIAA: ¹H NMR analysis (400 MHz, Me₂SO-*d*₆) δ 7.397–7.589 (m, 5H, ArH), 8.302 (singlet, 1H, H–C=N). ¹³C NMR analysis (100 MHz, Me₂SO-*d*₆) δ 171.660 (C=O, carboxylic acid), 150.455 (C=N, imine), 130.865 (Ar, C-1), δ 129.502 (Ar, C-4), 127.688 (Ar, C-3, C-5), 125.644 (Ar, C-2, C-6), mp 96–97 °C.

Standard PAM Reaction Conditions. All reactions were performed at 37.0 \pm 0.1 °C in the following solution: 100 mM MES/NaOH (pH 6.0), 30 mM NaCl, 1% (v/v) EtOH, 0.001% (v/v) Triton X-100, 1.0 μ M Cu(NO₃)₂, 5.0 mM sodium ascorbate, and 5.75 μ g/mL bovine liver catalase, 217 μ M O₂ and the desired oxidizable substrate. Morrison and Billett⁴⁴ showed that the ambient O₂ concentration at 1 atm and 37 °C is 217 μ M.

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Reverse-Phase HPLC (RP-HPLC) Analysis of PAM-Catalyzed Reactions. A RP-HPLC separation for *N*-benzoylglycine, α -hydroxy-*N*-benzoylglycine, benzamide, BIAA, and BOX was performed at 50 °C using a 65% (v/v) 50 mM potassium phosphate pH 6.0, 30% (v/v) methanol, and 5% (v/v) acetonitrile with a flow rate of 1.0 mL/min. Analytes were detected at 248 nm (Figure A, Supporting Information). A Hewlett-Packard HP-1100 HPLC equipped with a Phenomenex C₁₈-Hypersil column (100 mm \times 4.6 mm) attached to a HP-variable wavelength detector was employed for all HPLC separations. Standard curves for *N*-benzoylglycine, α -hydroxy-*N*-benzoylglycine, benzamide, BIAA, and BOX were prepared from the corresponding peak area values.

Characterization of Oxidation Product from BIAA. PAM (360 nM) was added to initiate the oxidation of 4.0 mM BIAA under standard reaction conditions, reaction volume = 0.5 mL. After a 20 h incubation, the reaction was diluted by addition of 8 mL of H₂O, extracted with ethyl acetate (20 mL \times 3), rinsed with brine, and dried over MgSO₄. The product was isolated by flash column chromatography and concentrated under reduced pressure. A small fraction of the BIAA oxidation product was first evaluated by RP-HPLC by retention time comparison to commercially available benzaldoxime. The remainder of the BIAA oxidation product was analyzed by ¹³C NMR using a Bruker 250 MHz NMR.

Determination of Glyoxylate Stoichiometry. The stoichiometry of BIAA oxidation to glyoxylate formation was determined under standard reaction conditions using 5.0 mM BIAA as the oxidizable substrate. BIAA oxidation was initiated by the addition of 80 nM PAM. At the desired time, a 40 μ L reaction aliquot was removed and the reaction terminated by the addition of trifluoroacetic acid (TFA) yielding a final concentration of 1% (v/v) TFA. Conversion of BIAA to benzaldoxime was monitored by RP-HPLC. All the RP-HPLC samples were diluted to a constant BOX concentration of 48 μ M (midpoint of the BOX standard curve) for glyoxylate analysis. The concentration of glyoxylate in the acid-quenched samples was determined by the lactate dehydrogenase-mediated reduction of glyoxylate to glycolate coupled to the oxidation of NADH to NAD⁺ ($\Delta\epsilon_{340} = 6.22 \times 10^3 \text{ M}^{-1} \text{ cm}^{-1}$). Glyoxylate-containing samples were combined with 7.6 units/mL lactate dehydrogenase in 100 mM potassium phosphate pH 7.0 and 220 μ M NADH and incubated for 1 h. at 37 \pm 0.1 °C. The increase in absorbance at 340 nm was measured using a JASCO V-530 UV–vis spectrophotometer.

Determination of Oxygen Stoichiometry during BIAA Oxidation. Amperometric analysis was used to correlate total [O₂] consumed during PAM catalysis under conditions of limiting [BIAA]. The PAM-dependent consumption of O₂ was followed using a Yellow Springs Instrument model 53 oxygen monitor equipped with a polarographic oxygen electrode interfaced with a personal computer using a Dataq Instruments analogue/digital converter (model DI-154RS) as previously described in McIntyre et al.⁴⁵

Background levels of [O₂] consumed, obtained in the absence of enzyme, were subtracted from the total [O₂] after the addition of PAM.^{46,47} Standard reactions were initiated with 2.0 nM PAM with the BIAA concentrations being 20, 35, 75, or 100 μ M. For each initial [BIAA], the concentration of O₂ (217 μ M) was in excess, and thus, BIAA was the limiting reagent. The total [O₂] consumed at each initial [BIAA] was determined from the asymptote of the O₂ consumption progress curve using SigmaPlot 8.0.

Measurement of Noncompetitive Kinetic Isotope Effects. Initial rates of [α -H₂]- and [α -²H₂]-BIAA were compared independently as a function of initial [O₂]. Working stocks of BIAA were calibrated by correlating the O₂ consumption to 20 μ M BIAA for

both substrates under standard reaction conditions. Variation in initial [O₂] was accomplished by mixing different proportions of an O₂ and N₂ with the final [O₂] was calibrated versus 217 μ M [O₂] at 37.0 \pm 0.1 °C after a 4 min equilibration.⁴⁸ Enzyme-dependent rates of O₂ consumption were obtained after initiation of the reaction with a \sim 110 nM (4–5 μ L) of PAM. Background rates of O₂ consumption obtained in the absence of PAM were subtracted from the enzyme-dependent rates. Values for both ^D(V_{MAX}/K_M)_{APPARENT, BIAA} and ^D(V_{MAX}/K_M)_{APPARENT, OXYGEN} were attained from the ratio of protio/deuterio kinetic parameters (eq 1) using Kaleidagraph.

$$\text{rate} = \frac{V_{\text{MAX,APPARENT}}^* [S]}{K_{\text{M,APPARENT}} + [S]} \quad (1)$$

Values for the ^D(V_{MAX}/K_M)_{APPARENT, BIAA} were also determined directly at a single BIAA concentration below the K_{M,APPARENT} values for [O₂], as the observed rate/[substrate] at low substrate concentration is expressed in the units of a second order rate constant, mM⁻¹ s⁻¹ (eq 2). The ^D(V_{MAX}/K_M)_{APPARENT, BIAA} was calculated by dividing the (V_{MAX}/K_M)_{APPARENT, BIAA} values obtained for protiated substrate by the deuterated substrate. The reported error is the standard error.

$$\begin{aligned} \text{rate} &= \frac{V_{\text{max,APPARENT}}^* [S]}{K_{\text{M,APPARENT}} + [S]} \rightarrow \text{rate} = \\ &\frac{V_{\text{max,APPARENT}}^* [S]}{K_{\text{M,APPARENT}}}, \text{ as } [S] \ll K_{\text{M,APPARENT}} \\ \therefore \frac{\text{rate}}{[S]} &\approx \frac{V_{\text{MAX,APPARENT}}}{K_{\text{M,APPARENT}}} \equiv \text{s}^{-1} \text{mM}^{-1} \quad (2) \end{aligned}$$

The plot of ^D(V_{MAX}/K_M)_{APPARENT, BIAA} versus [O₂] was fit to a hyperbolic curve and analyzed by the least-squares method, while the ^D(V_{MAX}/K_M)_{APPARENT, OXYGEN} versus [BIAA] was described as the mean isotope effect \pm standard deviation. The full initial rate data matrix for both α -diprotiated and α -dideuterated substrates versus [O₂] concentration were analyzed separately and fit to both eq 3 (steady-state preferred) and eq 4 (equilibrium preferred) using the programs of Cleland.⁴⁹ Single point ^D(V_{MAX}/K_M)_{APPARENT, BIAA} determinations were excluded from this analysis

$$\text{rate} = \frac{V_{\text{max}} [\text{BIAA}] [\text{O}_2]}{K_{\text{D,BIAA}} K_{\text{M,O}_2} + K_{\text{M,BIAA}} [\text{O}_2] + K_{\text{M,O}_2} [\text{BIAA}] + [\text{BIAA}] [\text{O}_2]} \quad (3)$$

$$\text{rate} = \frac{V_{\text{MAX}} [\text{BIAA}] [\text{O}_2]}{K_{\text{D,BIAA}} K_{\text{M,O}_2} + K_{\text{M,O}_2} [\text{BIAA}] + [\text{BIAA}] [\text{O}_2]} \quad (4)$$

Rates from eqs 3 and 4 are composed of the substrate concentrations; BIAA and O₂, the maximal initial velocity; V_{MAX}, Michaelis constants; K_{M,BIAA} and K_{M,O₂}, and the dissociation constant for BIAA, K_{D,BIAA}. Further analysis of ^D(V_{MAX}/K_M)_{APPARENT, BIAA} vs [O₂] and ^D(V_{MAX}/K_M)_{APPARENT, OXYGEN} vs [BIAA] data were performed to extrapolate values for ^D(V_{MAX}/K_M)_{APPARENT, OXYGEN}, as [O₂] \rightarrow 0 μ M and the value of ^D(V_{MAX}/K_M)_{APPARENT, OXYGEN}, as [O₂] \rightarrow ∞ μ M. The ^D(V_{MAX}/K_M)_{APPARENT, OXYGEN} as [O₂] \rightarrow 0 μ M values were determined through nonlinear regression analysis fitting replot data to eq 5. The values for ^D(V_{MAX}/K_M)_{APPARENT, OXYGEN} as [O₂] \rightarrow ∞ μ M were calculated from the reciprocal expression of eq 5, shown with the expression in eq 6.⁵⁰

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$$D\left(\frac{V_{\max}}{K_M}\right)_{\text{APPARENT.BIAA}} = \frac{k_{5H}}{k_{5D}} + \frac{k_{5H}\left(1 + \frac{k_3[\text{O}_2]}{k_2}\right)}{k_4 + k_{5H}\left(1 + \frac{k_3[\text{O}_2]}{k_2}\right)} \frac{1}{1 + \frac{k_{5H}\left(1 + \frac{k_3[\text{O}_2]}{k_2}\right)}{k_4 + k_{5H}\left(1 + \frac{k_3[\text{O}_2]}{k_2}\right)}} \quad (5)$$

$$\frac{1}{D\left(\frac{V_{\text{MAX}}}{K_M}\right) - D\left(\frac{V_{\text{MAX}}}{K_M}\right)_{[\text{O}_2] \rightarrow 0}} = \frac{1}{D\left(\frac{V_{\text{MAX}}}{K_M}\right)_{[\text{O}_2] \rightarrow \infty} - D\left(\frac{V_{\text{MAX}}}{K_M}\right)_{[\text{O}_2] \rightarrow 0}} \left(\frac{K}{\text{O}_2} + 1\right) K = \frac{k_2(k_4 + k_5 + k_{5H})}{k_3(k_5 + k_{5H})} \quad (6)$$

Quantum Mechanical/Molecular Mechanics Reaction Coordinate. The crystal structure of reduced peptidylglycine α -hydroxylating monooxygenase (PHM) was acquired from the protein databank (1SDW, 1.85 Å). All waters were removed from the structure along with all occurrences of the cocrystallized peptide ligand (IYT) outside of the active site.^{11,12,51} The bond orders and charges were then corrected on the cocrystallized protein structure. The copper atoms “atom types” were designated as ‘Cu’, hydrogens were added, and a restrained minimization was performed. The *bounding box* containing all ligand atoms was set to 14 Å³. Molecular oxygen was removed to prepare the structure using the *ProteinPrep* wizard for the quantum polarized ligand docking of BIAA (QPLD; Schrödinger First Discovery Suites, www.schrodinger.com). Initially, Glide⁵² was used to select five top poses using standard precision (SP) mode. These ligand–receptor complexes were analyzed using the Q-site⁵³ module where the bound ligand for each selected pose was treated by *ab initio* methods to calculate partial atomic charges utilizing electrostatic potential fitting within the receptor. All poses exceeding root mean squared (rms) and maximum atomic displacement of value 0.5 and 1.3 Å were rejected. Single-point energy determination was treated by the QPLD algorithm to determine the most energetically favorable ligand pose with respect to the receptor. Glide was then used to redock the ligand using each of the ligand charge sets previously calculated in Q-site.⁵³ To the QPLD output pose with the BIAA substrate docked into the PHM crystal structure, 3 Na⁺ ions were added to neutralize (zero net charge) the system along with the molecular oxygen using the coordinates from the original pdb. The system was explicitly solvated using the TIP3P water model to 10 Å from the receptor and equilibrated.³⁷ It should be noted that the O₂ coordinates were frozen during the entire system equilibration because it was not formally bound to the Cu_M domain and would, therefore, traverse the active site during equilibration if its coordinates were not restrained. The final minimized structure of the PHM–ligand central complex was attained through quantum mechanical treatment of the Cu_M pocket while treating the remainder of the protein structure classically to yield the correct final geometry. The resulting structure was used with Q-Site⁵³ for reaction

coordinate calculations. For the QM portion of the calculation a spin unrestricted hybrid density functional theory, with B3LYP hybrid-exchange functional using an LAVCP* basis set was used to more accurately define copper atoms using effective core potentials.^{54–57} The quantum mechanical region was treated with spin unrestricted open-shell calculations (UDFT) due to the presence of a radical in the system. The molecular mechanics portion of the protein was treated with the OPLS_2001 force field of Jorgensen.^{58,59}

There were two species of interest for study in the PHM oxidation reaction: The copper-hydroxyl (Cu^{II}-OH) and the copper-oxyl radical (Cu^{II}-O•). The Cu^{II}-superoxo species and the bound ligand were set to Cu^{II}-OH or Cu^{II}-O• and BIAA α -carbon radical in the input file. The quantum mechanical (QM) region was calculated using the Cu_M atom and domain residues (His₂₄₀, His₂₄₂, Met₃₁₄), the metal-bound oxygen species (either OH or O•), and BIAA C α radical substrate. On this selection, a Q-site job was run without coordinate constraints to determine the optimized starting point geometry. The calculation was repeated with the oxygen species frozen and the C α of BIAA frozen to determine the energetic minima starting point. Analysis of the reaction coordinate for substrate oxidation was determined through the difference calculated from incremental increase in the distance between the copper bound oxygen bond and the corresponding decrease in the C α -O(H) bond. The coordinates determined for each change in reaction coordinate were refrozen for those two atoms and the energy calculated quantum mechanically. Results were visualized using the *Maestro*.⁶⁰

Results

Reaction Stoichiometry and Product Identification for the PHM-Catalyzed Oxidation of BIAA. The ¹³C NMR of the PHM-generated product generated from BIAA exhibited a spectrum with chemical shifts similar to those of the BIAA starting material, containing signature aromatic and imine carbon shifts (represented A–E, Figure B, Supporting Information). However, the two carbon shifts found at 171.8 and 69.3 ppm from the *O*-acetyl moiety in BIAA were not present in the product spectrum. The RP-HPLC retention time of the isolated product was 12.4 min, identical within experimental error to that of commercially available benzaldoxime, and differed significantly from that of the BIAA starting material (6.0 min).

Using BIAA as the limiting substrate, the stoichiometry for the PHM-dependent O₂ consumption was 0.97 ± 0.06 O₂ consumed/BIAA oxidized. (Figure C and Table C Supporting Information). The LDH-catalyzed reduction of glyoxylate to glycolate with the concomitant oxidation of NADH to NAD⁺ provided a convenient spectroscopic method (a) to demonstrate that glyoxylate is a product of the PHM-catalyzed oxidation of BIAA and (b) to measure the glyoxylate concentration generated from BIAA upon PHM treatment. Percent conversion of BIAA to BOX was measured by RP-HPLC. Using the individual methods to measure [BOX] and [glyoxylate], we determined that the [glyoxylate]/[BOX] stoichiometry for the PHM-catalyzed oxidation of BIAA was 1.1 ± 0.1 (Figure 1).

Stability of the BIAA Oxidation Product. Following the PHM-dependent oxidation of BIAA, there are two possible routes for the generation of the observed products, BOX and

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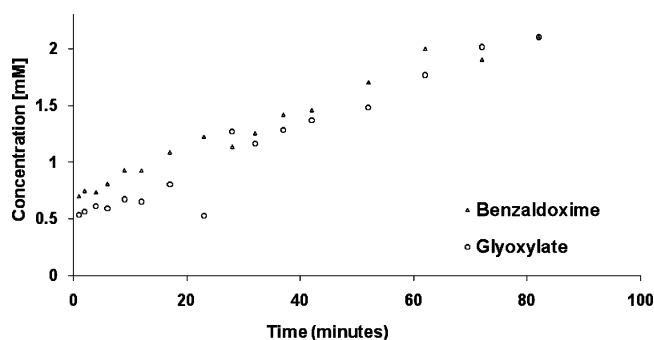


Figure 1. Determination of the PAM-dependent ratio of formed products (oxime and glyoxylate) using the imino-oxy acetic acid substrate. The concentrations of oxime and glyoxylate products were independently determined by RP-HPLC and spectrophotometry.

glyoxylate (Scheme 4a). Path A in Scheme 4 represents a stable α -hydroxylated BIAA which then requires PAL to catalyze BOX and glyoxylate formation while path B requires only PHM for BOX and glyoxylate formation from BIAA. Here, PHM-catalyzed oxidation of the α -carbon would accompany a PAL-independent, chemical dealkylation to yield BOX and glyoxylate.

Our approach to differentiating between paths A and B in Scheme 4 was to measure BOX formation in the presence of α -hydroxyhippurate, a known PAL substrate.^{17,48} If the putative α -hydroxylated BIAA product generated by PHM requires PAL for dealkylation, BOX formation would be inhibited by α -hydroxyhippurate. As a control, benzamide formation via the PAL-catalyzed dealkylation of α -hydroxyhippurate would also be inhibited by BIAA. Of course, if the BIAA product generated by PHM is not a PAL substrate, neither BOX formation nor benzamide formation would be affected.

The progress curves for BOX formation from BIAA measured in the presence of either 6 mM or 10 mM α -hydroxyhippurate were unaffected at all BIAA concentrations employed (Figure 2). Similarly, the progress curves for benzamide formation from α -hydroxyhippurate were unaffected by the presence of 0.5, 2, or 3 mM BIAA (Figure 3).

Kinetic Isotope Effects. The $D(V/K)_{\text{APPARENT,BIAA}}$ decreased as the concentration of O_2 decreased, while the $D(V/K)_{\text{APPARENT},O_2}$ remained constant over the range of BIAA concentrations used (Tables A and B, Supporting Information). Data fit to the steady-state preferred mechanism (eq 3) had root square values (σ) of 0.237 and 0.140 for α -protiated and α -dideuterated BIAA, respectively. Conversely, data fit to the equilibrium preferred mechanism (eq 4) displayed root square values (σ) of 0.236 and 0.156 for the protiated and α -dideuterated substrates, respectively. An approximate 10-fold increase in variance was observed for the fit to equations for the equilibrium preferred mechanism, as compared with its steady-state preferred counterpart.⁴⁹ A large increase in the average residual least-squares (variance) term was observed, showing a greater deviation in error between experimental rates (VEXP) and the predicted/optimized rates (VOPT) for the linear regression model. The variance for [α - 1H_2]-BIAA and [α - 2H_2]-BIAA increased from 0.0561(H) and 0.0230(D) from for steady-state preferred to 0.555(H) and 0.242 (D) to for equilibrium preferred. Kinetic data fit to the steady-state preferred model (eq 3) displayed an increased statistical significance based on their variance terms, though the square root (σ) values were statistically very similar.

The graphs of $1/\text{rate}$ vs $1/[\text{BIAA}]$ and $1/\text{rate}$ vs $1/[O_2]$ both exhibit intersections in the same quadrant, with a negative abscissa and positive ordinate when fit to the steady-state preferred equation.

The extrapolated value of $D(V/K)_{\text{APPARENT,BIAA}}$ as $[O_2] \rightarrow 0$ was determined to be 14.7 ± 1.0 (Figure D, Supporting Information), and the $D(V/K)_{\text{APPARENT,BIAA}}$ as $[O_2] \rightarrow \infty$ was calculated to be 1.01 ± 0.03 (Figure E, Supporting Information). The value for the $D(V/K)_{\text{APPARENT},O_2}$ based on a steady-state preferred mechanism (eq 3) was calculated to be 5.8 ± 0.7 (Table 1). This value is consistent with experimental determinations of $D(V/K)_{\text{APPARENT},O_2}$ obtained at different initial O_2 concentrations at one fixed initial BIAA concentration, yielding an average $D(V/K)_{\text{APPARENT},O_2}$ of 4.7 ± 1.1 (Figure 4, blue circles, O). $D(V_{\text{MAX}})$ was also calculated to be near unity at 0.8 ± 0.1 , while $D(K_{D,BIAA})$ was found to be 2.1 ± 0.5 . The $K_{M,BIAA}$ parameters for both α -protiated and α -dideuterated BIAA were determined to be negative with large standard errors. This suggests that the $K_{M,BIAA}$ was not statistically significant, consistent with a steady-state preferred kinetic mechanism.^{48,49}

QM/MM Reaction Coordinate. The two species evaluated computationally were the $Cu^{II}\text{-OH}$ (singlet) and the $Cu^{II}\text{-O}\bullet$ (quartet). The potential energy of the reaction coordinate for the $Cu^{II}\text{-OH}$ species, calculated as the difference in bond distance between the $C_\alpha\text{-O}$ and $Cu\text{-OH}$ bonds, displayed a nearly linear increase in relative energy from the starting point to a ‘saddle point’ value of +45.93 kcal/mol. Following this point, the calculated energy of the reaction coordinate decreased to -25.98 kcal/mol (Figure 5). The $C_\alpha\text{-O}$ bond lengths decreased from 4.60 to 2.80 Å at the energetic ‘saddle point’, resting at a final distance of 1.30 Å. The $Cu^{II}\text{-OH}$ bond lengths increased over the reaction coordinate from 1.82 through 2.81 Å at the energetic maxima, to 4.08 Å at the energetic minima. The imino-oxy \leftrightarrow α -carbon bond lengths for BIAA increased 0.16 Å over the reaction coordinate, from 1.36 to 1.52 Å (Figure 5). The sequence shown in Scheme B and movie A (Supporting Information) represent the quantum mechanical aspects of the performed QM/MM study. The displayed poses correspond directly with their respective steps in Figure 5. The orientation of the carboxyl of BIAA was unchanged until the step prior to the saddle point (+37.45 kcal/mol). Following this point, a coupled rotation in the hydroxyl moiety and the carboxyl of BIAA was shown which oriented the α -carbon essentially perpendicular to the approaching hydroxyl group (Scheme B, Supporting Information).

The reaction coordinate for the copper-alkoxide ($Cu^{II}\text{-O}\bullet$) species (Figure 6) displayed a more complex relationship with potential energy trend, relative to that of $Cu^{II}\text{-OH}$, as an energetic plateau was observed midway to the saddle point at ~ 28 kcal/mol. Following this, potential energy at the ‘saddle point’ was +54.75 kcal/mol, +12.82 kcal/mol greater than that observed for the reaction coordinate for the $Cu^{II}\text{-OH}$ species. Although a higher potential energy was observed for the $Cu^{II}\text{-O}\bullet$ species, a rapid decline in the potential energy was observed beyond the ‘saddle point’ on the reaction coordinate (Figure 6), in contrast to slow decline observed for the $Cu^{II}\text{-OH}$ species (Figure 5). The ‘saddle points’ for the $Cu^{II}\text{-OH}$ species occurred with nearly equal bond lengths for the OH between the C_α and Cu^{II} (~ 2.8 Å at RC = 0.014). Conversely, the ‘saddle point’ of the $Cu^{II}\text{-O}\bullet$ reaction coordinate was much less symmetrical, reaching its maxima when the $C_\alpha\text{-O}$ and $Cu^{II}\text{-O}$ bond lengths were 1.9 and 3.7 Å, respectively. The $Cu^{II}\text{-OH}$ potential energy decreased over the reaction coordinate (RC) from +45.93 kcal/mol at RC = 0.014 to -25.97 kcal/mol at RC = 2.78 while the $Cu^{II}\text{-O}\bullet$ decreased from +58.75 kcal/mol at RC = 1.78 to -9.68 kcal/mol at RC = 3.05. Therefore, the total change in potential energy for each species was similar for each species (within

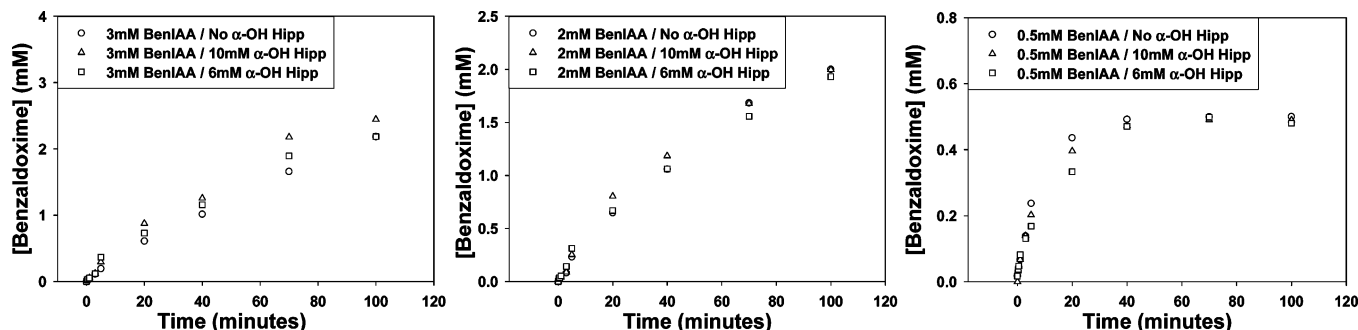


Figure 2. Time course for the PAM-dependent conversion of 3, 2, and 0.5 mM benzaldehyde imino-oxy acetic acid to benzaldoxime in the presence and absence of α -hydroxy-*N*-benzoylglycine. Results were calculated from RP-HPLC analysis (percent conversion).

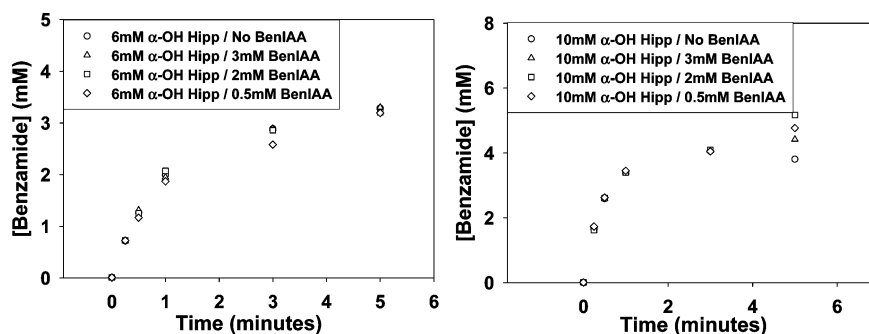
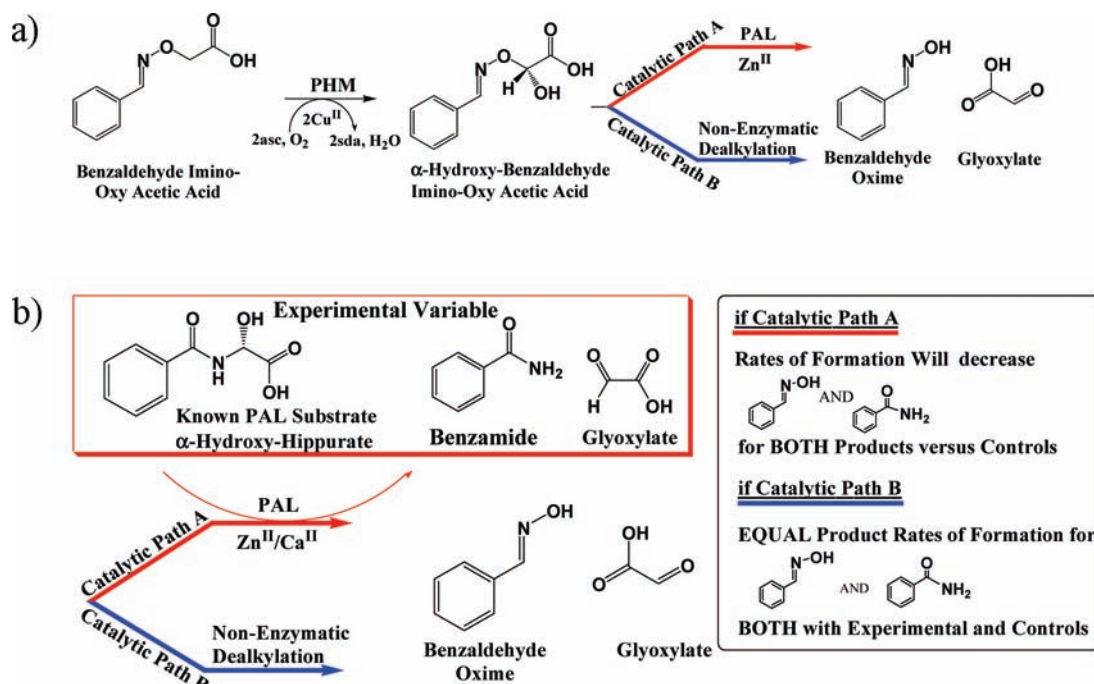


Figure 3. Resolved time course for the conversion of 6 mM and 10 mM α -hydroxy-*N*-benzoylglycine to benzamide in the presence and absence of 0.5, 2, and 3 mM BIAA (BenIAA). Please note, time point data were truncated for clarity.

Scheme 4. (a) Representation of the Two Possible Pathways for BOX and Glyoxylate Formation Following BIAA Oxidation in the PHM-Domain. Please Note That ‘asc’ and ‘sda’ Represent Single Electron Contributions from Exogenous Ascorbate and Semi-dehydroascorbate; (b) Experimental Design to Kinetically Differentiate the Two Possible Degradation Pathways for BIAA to Its Corresponding BOX Product Following PHM-Dependent Oxidation



experimental error), 68.43 kcal/mol for $Cu^{II}\text{-O}^\bullet$ and 71.90 kcal/mol for $Cu^{II}\text{-OH}$.

Substrate Dealkylation. The increase in imino-oxy \leftrightarrow α -carbon bond length for the $Cu^{II}\text{-O}^\bullet$ species (Figure 6) was observed to be constant over the reaction coordinate until the energetic ‘saddle point’ was reached. It was here that the substrate bond

length was dramatically increased as compared to the effect observed studying the $Cu^{II}\text{-OH}$ species, ranging from 1.42 to 2.91 Å. The corresponding poses for the quantum mechanically treated region of the $Cu^{II}\text{-OH}$ simulation showed very little change in substrate orientation, with major functionalities of the of the BIAA substrates (phenyl ring, imino-oxy, and carboxy

Table 1. Kinetic Parameters Determined from the Steady-State Preferred Bi-Substrate Michaelis–Menten Eq 3

	$V_{\text{MAX}}(\text{s}^{-1})$	$K_{\text{D,BIAA}}(\text{mM})$	$K_{\text{M,O}_2}(\text{mM})$	$(V_{\text{MAX}}/K_{\text{M}})_{\text{O}_2}(\text{M}^{-1}\text{s}^{-1})$	$^{\text{D}}K_{\text{D,BIAA}}$	$^{\text{D}}(V_{\text{MAX}})$	$^{\text{D}}(V_{\text{MAX}}/K_{\text{M}})_{\text{O}_2}$
protiated	4.75 (0.19)	758.18 (133.42)	95.37 (12.18)	$4.98 \times 10^4 (5.03 \times 10^3)$	2.06 (0.52)	0.84 (0.08)	5.78 (0.71)
deuterated	5.68 (0.46)	367.95 (66.58)	658.08 (95.02)	$8.63 \times 10^3 (6.02 \times 10^2)$			

termini moieties) remaining constant with respect to starting points. At the ‘saddle point’ of the $\text{Cu}^{\text{II}}\text{-O}^{\bullet}$ simulation, an increase in imino-oxy \leftrightarrow α -carbon bond length was observed during a small RC window with a 1.47 Å change between RC = 1.78 and 2.09 yielding a final imino-oxy \leftrightarrow α -carbon bond of 2.91 Å (Scheme C and movie B, Supporting Information).

Discussion

Dealkylation is Non-enzymatic. Oxidation of BIAA by PAM yields two products in a 1:1 molar ratio: BOX and glyoxylate. BOX was confirmed by ^{13}C NMR Figure B, (Supporting Information). In addition, the stoichiometry of $[\text{O}_2]_{\text{consumed}}: [\text{BOX}]_{\text{produced}}$ was also 1:1 Figure C and Table C (Supporting Information), consistent with evidence for tight coupling between substrate oxidation and O_2 consumption for PHM⁶¹ and the related enzyme, D β M.²⁰ PAM-catalyzed oxidation/dealkylation of BIAA is unaffected by α -hydroxyhippurate (a PAL substrate) and the PAL-catalyzed dealkylation of α -hydroxyhippurate is unaffected by BIAA (Figures 2 and 3). This suggests that the putative α -hydroxylated product generated by PHM, $\text{C}_6\text{H}_5\text{-CH-N-O-CH(OH)-COOH}$, is unstable and non-enzymatically decomposes into observed oxime and glyoxylate products. These data provide strong evidence that the oxidative dealkylation of BIAA to BOX and glyoxylate requires only the PHM domain of bifunctional PAM.

Kinetic Mechanism. Previous work with small-molecule substrates, *N*-benzoylglycine⁴⁸ and *N*-acetylglycine (N. R. McIntyre and Merkler, D. J., manuscript in preparation), found that $^{\text{D}}(V_{\text{MAX}}/K_{\text{M}})$ values for both were equal and constant as the O_2 concentration was varied. The minimal kinetic mechanism of these small-molecule substrates is equilibrium-ordered with O_2 binding after *N*-benzoylglycine or *N*-acetylglycine. Comparison of eqs 3 and 4 shows that K_{M} for the first bound substrate ($K_{\text{M,A}}[\text{B}]$) is absent from the denominator for an equilibrium-preferred mechanism. In this case, binding of the first substrate to the free enzyme (E) is in

equilibrium. The dissociation constant ($K_{\text{D,A}}[\text{B}]$) represents the affinity of the enzyme for the first bound substrate. Analysis of deuterium KIEs for a steady-state preferred mechanism, will include the K_{M} for the first bound substrate ($K_{\text{M,A}}[\text{B}]$) in the denominator, yielding a symmetrical mechanism. Therefore, the $^{\text{D}}(V_{\text{MAX}}/K_{\text{M}})_{\text{APPARENT}}$ value cleavage will be dependent on the concentration of the second substrate for a steady-state preferred mechanism, as we observed for $\text{C}_{\alpha}\text{-D/H}$ bond cleavage using BIAA as a substrate. The difference between a steady-state *ordered* and a *random* mechanisms is that the $^{\text{D}}(V_{\text{MAX}}/K_{\text{M}})_{\text{APPARENT}}$ values approach unity as concentration of the second substrate becomes infinite for ordered substrate addition vs a nonunity limit for the random substrate addition.⁶² The values of $^{\text{D}}(V_{\text{MAX}}/K_{\text{M}})_{\text{APPARENT, BIAA}}$ showed a dramatic decrease as the concentration of O_2 increased. Extrapolation of the $^{\text{D}}(V_{\text{MAX}}/K_{\text{M}})_{\text{APPARENT, BIAA}}$ to zero $[\text{O}_2]$ yields a value ~ 15 (well beyond the semiclassical limit), while extrapolation of infinite $[\text{O}_2]$ yields of 1.0 (Figures C and D, Supporting Information). The $^{\text{D}}(V_{\text{MAX}}/K_{\text{M}})_{\text{APPARENT, O}_2}$ was constant over the range [BIAA] used. This is a clear example of a steady-state ordered mechanism with BIAA binding to reduced PHM prior to O_2 (Scheme 5). Using hippurate as the oxidizable substrate, it has been established that PHM exhibits an equilibrium-ordered mechanism with O_2 binding after hippurate to reduced enzyme,⁴⁸ while the kinetic mechanism for the Y318F mutant is steady-state random with either hippurate or O_2 binding to reduced enzyme.⁶³ A steady-state random mechanism was observed for the Y318F mutant of PHM as the a nonunity values of $^{\text{D}}(V_{\text{MAX}}/K_{\text{M}})_{\text{APPARENT, HIPPURATE}}$ were obtained as $[\text{O}_2] \rightarrow \infty$.⁶³ For the related enzyme, D β M, the absence of the anionic activator (fumarate) at pH 6.0 alters the steady-state mechanism from *ordered* to *random*.^{50,64}

The steady-state mechanism is attained for BIAA through a decrease in the rate constant responsible for dissociation of the binary complex (E–S), k_2 .⁴⁸ Unlike an equilibrium preferred mechanism, the binary complex (E–S) becomes the predominant enzyme state. The reduction in $^{\text{D}}(V_{\text{MAX}}/K_{\text{M}})_{\text{APPARENT, BIAA}}$ with increasing $[\text{O}_2]$ was due to an increase in the ‘‘commitment to catalysis’’.^{65–67} Under steady-state ordered conditions, as $[\text{O}_2] \rightarrow \infty$, dissociation of the reactant ($\text{R}_{\text{C}\alpha\text{-H}}$) from the central complex does not occur as the commitment to the forward reaction also becomes infinite ($^{\text{D}}(V_{\text{MAX}}/K_{\text{M}})_{\text{APPARENT, BIAA}} = \sim 1$). The magnitude of $^{\text{D}}(V_{\text{MAX}}/K_{\text{M}})_{\text{BIAA}}$ as $[\text{O}_2] \rightarrow 0$, has a commitment to the forward reaction which is dependent on the ratio of the chemical step vs the dissociation of O_2 from the central complex (k_5/k_4 , Scheme 5). Therefore, the predominant enzyme form would be E– $\text{R}_{\text{C}\alpha\text{-H}}$ due to the decrease in the rate constant for O_2 insertion to achieve the central complex, $k_3[\text{O}_2]$.

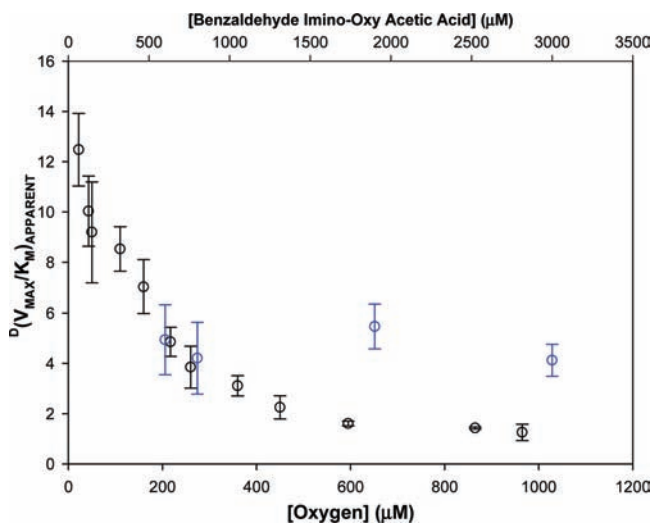


Figure 4. Dependence of [oxygen] on $^{\text{D}}(V_{\text{MAX}}/K_{\text{M}})_{\text{APPARENT, BIAA}}$ (black symbols) and the [BIAA] dependence on $^{\text{D}}(V_{\text{MAX}}/K_{\text{M}})_{\text{APPARENT, O}_2}$ (blue symbols).

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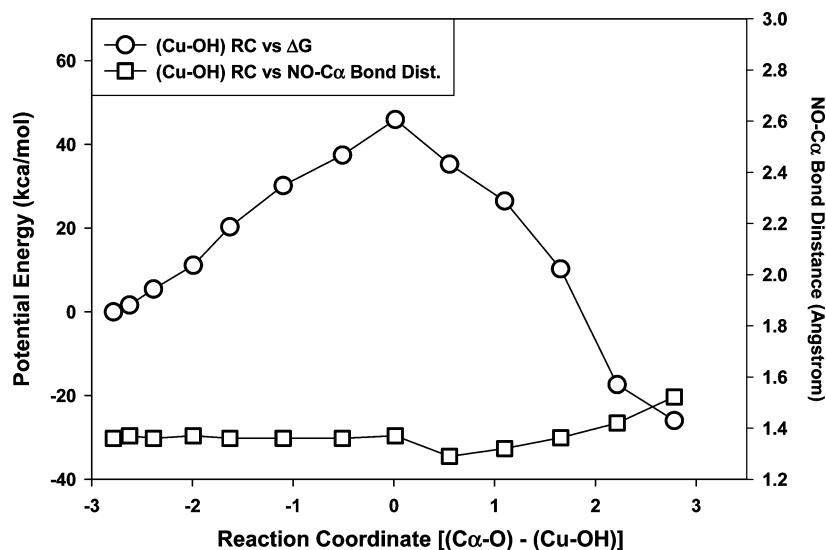


Figure 5. QM/MM simulated reaction coordinate for the oxidation chemistry observed for the BIAA C_α-radical oxidation with a singlet Cu^{II}-OH species (circles). Squares represent dealkylation distances observed for imino-oxy ↔ C_α bond lengths versus reaction coordinate.

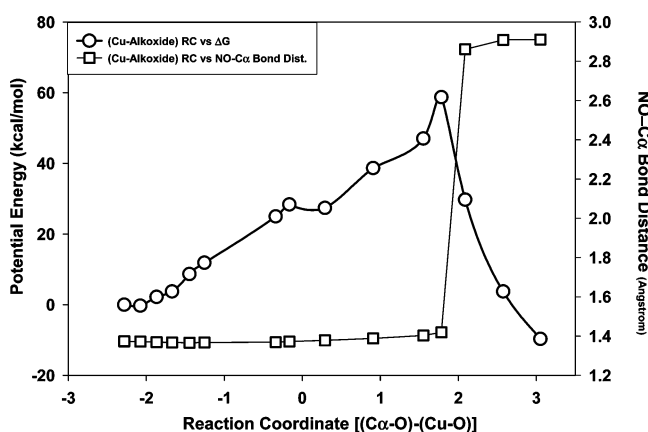


Figure 6. QM/MM simulated reaction coordinate for the oxidation chemistry observed for the BIAA C_α-radical oxidation with a quartet Cu^{II}-O[•] species (circles). Squares represent dealkylation distances observed for imino-oxy ↔ C_α bond lengths versus reaction coordinate.

The dissociation constant for BIAA showed a normal isotope effect of $^D K_{D,BIAA} = 2.06 \pm 0.52$ (Table 1). Conversely, the binding isotope effect for hippurate is inverse for both PHM and PAM, $^D K_{D,hippurate} < 1.0$.⁴³ Often the effect of isotopic labeling on binding is dismissed as negligible, but binding isotope effects have been reported for a number of enzymes, including lactate dehydrogenase,⁶⁸ hexokinase,⁶⁹ thymidine phosphorylase,⁷⁰ and purine nucleoside phosphorylase.⁷¹ The $^D K_{D,BIAA}$ suggests that deuteration of the BIAA α -carbon reduces stringent vibrational contributions associated with ground-state destabilization during formation of the E•BIAA complex formation, potentially decreasing k_2 . A decrease in k_2 increases the forward commitment to catalysis, k_3/k_2 , and may be a contributing factor in the kinetic mechanism for BIAA being

steady-state ordered vs equilibrium ordered. The forward commitment for an equilibrium order mechanism will be relatively low.

While neither hippurate nor BIAA showed an appreciable $^D V_{MAX}$, the turnover number decreased approximately 6-fold for BIAA relative to hippurate. This difference in product release (V_{MAX}) for BIAA cannot be attributed to an uncoupling of O₂ consumption from BOX and glyoxylate formation (Figures 1, B and Table B), as the reaction stoichiometries for BIAA oxidation match those reported for glycine-extended substrates.⁷² One possible reason for the relatively low V_{MAX} for BIAA is that the imino-oxy moiety is complicating C_α-O insertion into its respective C_α-radical species. This conclusion would mean that BIAA dealkylation would be included in the minimal kinetic mechanism and would require no other rate constant for product release other than k_7 . This rate constant describes the slow step responsible for the irreversible conversion of the E•P complex to free enzyme (E) and product (P) for all PHM substrates. Therefore, the absence of $^D V_{MAX}$ may best be explained as a requisite rate-limiting conformational change necessary for product release with glycine-extended substrates, though an additional energetic penalty associated with BIAA oxidation (dealkylation) appears to further decrease the magnitude of k_7 for this substrate.

Substrate Oxidation and Product Release are Uncoupled. QM/MM simulation of both the oxygen insertion into the BIAA C_α-radical and the resulting dealkylation process provided an experimental constraint used to probe the nature of the Cu/O oxidant species. The Cu^{II}-OH (singlet) species was postulated to affect the direct OH transfer and product release steps of the ‘side-on/ η^2 ’ and ‘copper-oxo pathway B’ mechanisms. The effect of direct product formation on the bond length of our dealkylation probe was negligible, suggesting that imino-oxy ↔ C_α bond dissociation would likely not occur through this mechanism. Formation of the copper^{II}-alkoxide species (quartet) had a dramatic effect on the structure of the dealkylation probe, suggesting that the process for substrate oxidation and product release were completely uncoupled, and likely to pass through

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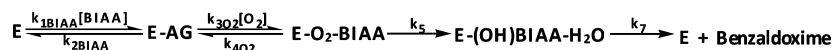
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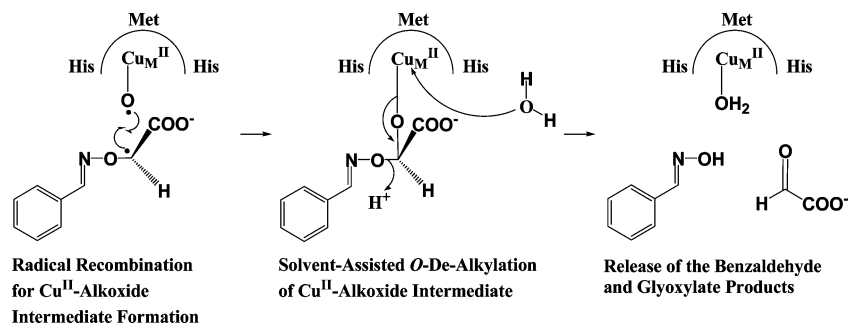
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Scheme 5. Minimal Kinetic Mechanism Determined for BIAA Oxidation As Determined by Primary Deuterium KIEs



Scheme 6. Dealkylation Reaction Predicted for the Cu^{II}-Alkoxide Intermediate Formation with BIAA As the Substrate



a copper-alkoxide intermediate (red coordinates, Figure 7). The doublet copper-oxo species ('copper-oxo pathway A') could not be used for this analysis as H-abstraction and oxidation steps were concerted resulting in an absence of a substrate intermediate (radical).³⁷ As the separation of H-abstraction from substrate oxidation was a necessary caveat for our theoretical treatment of dealkylation, calculations using the double species could not be performed.

Conclusion

Due to the complexities of PHM, theoretical models lacking experimental constraint often lead to ambiguous conclusions in thermodynamic preference between chemical models. Here, the catalytic power of the enzyme is determined at its upper limit and does not take into account that the reaction regime only requires the *necessary* energy for catalysis. The work of Karlin (*supra vide*) infers that the nature of the Cu/O oxidant cannot definitively be characterized in PHM, as the reaction coordinate may proceed through several possible catalytic paths. Therefore, the dealkylation event was designed to describe the nature of substrate oxidation chemistry in PHM. BIAA was a superb probe for the glycine-extended substrate functionalization as it retained all reaction stoichiometries important to PHM. When docked to PHM, non-Lewis hyperconjugative stabilization

of the oxime moiety is electronically analogous to an amide (*manuscript in preparation*). For the QM/MM study, C_α-H bond cleavage was not part of the reaction coordinate; instead the dealkylation of BIAA was used to provide the system with an independent variable to constrain the theoretical model. The mixture of both *in vitro* and *in silico* data suggests that *direct* hydroxyl insertion into the C_α-radical of BIAA is unlikely as the NO ↔ C_α distance did not significantly increase. Interestingly, the singlet species reaction coordinate was thermodynamically favored as the oxidant with high symmetry over the simulation. This mechanism would likely lead to a complete uncoupling of the BIAA radical and a reduced oxygen species from the PHM active site, as the substrate radical would resist conversion into a thermodynamically unstable hemiacetal derivative. PHM would not be expected to stabilize the putative HO-C_α BIAA intermediate as the active site is highly solvated during catalysis.⁷³ Thus, if this hemiacetal formed, its instability would not be suppressed through protein docking.

Using the natural glycine-extended substrate, hydrolysis of the 'copper-alkoxide/substrate' intermediate complex yields a stable α-hydroxylated glycine product. The BIAA model displayed changes in the bond lengths for the oxygen within the Cu^{II} ↔ O• complex approaching the carbon radical (a decrease) as NO ↔ C_α (an increase) over the reaction coordinate (Figure 7). The steady-state data suggest that this process is irreversible and tightly coupled with respect to both oxygen and BIAA, similar to the PHM natural substrate.²⁰ Taken together, the reason for the *O*-dealkylation of BIAA occurring independently of PAL is likely that the oxidation step becomes increasingly rate limiting versus the subsequent hydrolysis step in BIAA compared to a glycine-extended substrate. This irreversible event was calculated to be an endothermic process (Figure 7) in both singlet and quartet copper-oxo complexes, though *O*-dealkylation of BIAA was only observed for the singlet-state species calculation. By comparison, for glycine-extended substrates, this event has been calculated to be an exothermic process.^{19,35}

Our proposal that the 'Cu^{II}-O•' species was the oxidant in PHM is based on the dealkylation characteristics unique to BIAA. Formation of the 'inner-sphere alcohol' complex during BIAA oxidation was the only mechanism which could account for the tightly coupled reaction stoichiometries while still displaying the ability to dealkylate independently of PAL. The copper^{II}-alkoxide was thermodynamically disfavored compared

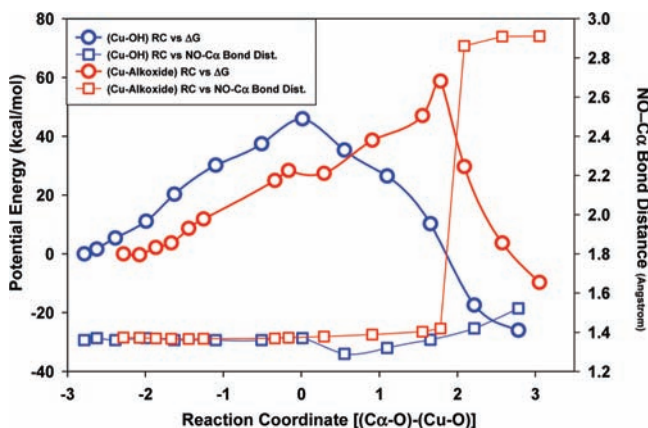


Figure 7. Combined plot of QM/MM reaction coordinate simulation comparing the Cu^{II}-OH (singlet) and the Cu^{II}-O• (quartet) species against bond distances for the BIAA dealkylation event. Please note, squares represent the distance change between the NO ↔ C_α bond distance, while the circles signify Cu—O bond distance changes over the reaction coordinate. For more details, please see the Materials and Methods section.

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to the singlet $\text{Cu}^{\text{II}}\text{-OH}$ species, though it was the increase in $\text{NO} \leftrightarrow \text{C}\alpha$ bond length that pointed strongly to an 'inner-sphere alcohol' intermediate. Therefore, production of an oxidized copper^{II}-alkoxide intermediate complex would be favored to facilitate the reaction characteristics observed during the BIAA *O*-dealkylation process (Scheme 6). In conclusion, this study provides compelling evidence that PHM-dependent oxidation of glycine-extended substrates occurs through a covalently linked copper^{II}-alkoxide substrate complex.

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Supporting Information Available: Experimental figures, tables, schemes, and movies. This material is available free of charge via the Internet at <http://pubs.acs.org>.

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