

Direct nitration and azidation of aliphatic carbons by an iron-dependent halogenase

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Iron-dependent halogenases employ *cis*-halo-Fe(IV)-oxo (haloferryl) complexes to functionalize unactivated aliphatic carbon centers, a capability elusive to synthetic chemists. Halogenation requires (i) coordination of a halide anion (Cl⁻ or Br⁻) to the enzyme's Fe(II) cofactor, (ii) coupled activation of O₂ and decarboxylation of α -ketoglutarate to generate the haloferryl intermediate, (iii) abstraction of hydrogen (H \bullet) from the substrate by the ferryl and (iv) transfer of the *cis* halogen as Cl \bullet or Br \bullet to the substrate radical. This enzymatic solution to an unsolved chemical challenge is potentially generalizable to installation of other functional groups, provided that the corresponding anions can support the four requisite steps. We show here that the wild-type halogenase SyrB2 can indeed direct aliphatic nitration and azidation reactions by the same chemical logic. The discovery and enhancement by mutagenesis of these previously unknown reaction types suggest unrecognized or untapped versatility in ferryl-mediated enzymatic C-H bond activation.

The capacity to install functional groups onto aliphatic carbon centers late in synthetic sequences is considered a holy grail of synthetic chemistry. Despite much effort and major progress, it remains an unsolved problem^{1,2}. Biosynthetic sequences, by contrast, provide many spectacular examples of late aliphatic C-H bond functionalization³⁻⁹. Enzymes that employ (i) a free-radical or high-valent metal-oxo intermediate to abstract hydrogen (H \bullet) from an aliphatic carbon; (ii) a substrate binding site that permits only the C-H target and not other, more reactive functional groups to approach the reactive intermediate; and (iii) allosterically controlled timing of intermediate formation achieve a degree of specificity and control not yet replicated in chemical systems^{6,7}. However, biological C-H functionalization would, in light of current knowledge, seem to have its own limitations. Whereas enzymatic couplings of carbon¹⁰, oxygen^{3-5,11}, phosphorus¹², sulfur¹³ and halogen atoms¹⁴⁻¹⁶ to aliphatic carbon centers are known, examples of the installation of nitrogenous functional groups have not been forthcoming. Indeed, although there have been recent reports of an intramolecular amination of a benzylic carbon in an elegantly engineered, unnatural substrate and the nitration of an aromatic carbon by cytochrome P450 enzymes¹⁷, enzymatic C-N coupling at a completely unactivated aliphatic carbon center is unknown. In this reaction type, the development of synthetic methods has, ironically, outstripped the discovery or development of enzymatic activities^{18,19}. It remains unclear whether such reactions do not exist in nature or simply remain to be discovered.

The iron(II)- and α -ketoglutarate-dependent (Fe/ α KG) hydroxylases and halogenases have similar structures and share a common initial mechanism for aliphatic C-H activation^{3-5,11}. In each reaction, an Fe(IV)-oxo (ferryl) intermediate, which forms from the Fe(II) cofactor, O₂, and two electrons derived from oxidative decarboxylation of α KG to succinate (Fig. 1a), abstracts H \bullet from the substrate to form a carbon-centered radical and Fe(III)-OH form of the cofactor (Fig. 1b)^{3-5,11,20}. The reactions diverge according to the fate of this intermediate (Fig. 1b). In the hydroxylases, the substrate

radical couples with the OH ligand (termed 'oxygen rebound'²¹), thereby returning the cofactor to its resting +2 oxidation state. In the halogenases, the substrate radical instead couples with a halogen ligand coordinated *cis* to the OH ligand, presumably converting the cofactor to an Fe(II)-OH form^{15,16,22-24}. The halogen that is ultimately transferred to the substrate initially coordinates to the Fe(II) cofactor as free chloride or bromide at a position that is occupied in the hydroxylases by a carboxylate ligand from an aspartate residue^{22,25} (Fig. 1a). The controlling structural distinction is thus the replacement of the protein-donated carboxylate ligand in the hydroxylases by a noncoordinating alanine in the halogenases to open the site for coordination of halide²² in preparation for O₂-driven ferryl formation^{15,16,23} and ferryl-mediated radical group transfer^{24,25}.

For the halogenase SyrB2 from the syringomycin biosynthetic pathway of *Pseudomonas syringae* B301D^{25,26}, the basis for selective coupling of the substrate carbon radical with the *cis* halogen rather than the hydroxyl ligand is the positioning of the C-H target of the substrate away from the oxo or hydroxo ligand and closer to the halogen, which effectively sacrifices proficiency in the initial H \bullet abstraction for selectivity in the subsequent group transfer step²⁴. Modifications to the native substrate (the C₄ amino acid, L-threonine, appended to the carrier protein, SyrB1(Thr-S-SyrB1)²⁵; Fig. 1c) anticipated to permit closer approach of the side chain methyl target to the oxygen ligand (for example, lengthening of the alkyl side chain) are associated with marked acceleration of an otherwise sluggish H \bullet abstraction step and a marked increase in competing hydroxylation. The substrate with the C₅ amino acid, L-norvaline (Nva), appended to SyrB1 (Nva-S-SyrB1) supports 130-fold faster H \bullet abstraction and undergoes primarily hydroxylation at C5. Computational studies have provided structural rationales for the positioning effect²⁷ and suggested that the different target positions result in the engagement of different frontier orbitals in the H \bullet abstraction step, which in turn sets the stage for different group transfer outcomes^{28,29}. SyrB2 thus has two crucial features that enable its halogenation outcome: substrate and cofactor binding

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Figure 1 | Related mechanistic logic of the Fe/ α KG halogenases and hydroxylases.

(a) Steps leading to formation of the H \bullet -abstracting ferryl or haloferryl complexes. (b) Divergence of the mechanisms of ferryl- or haloferryl-mediated hydroxylation (top, magenta arrows) and halogenation (bottom, green arrows). (c) Structure of the native SyrB2 substrate, Thr-S-SyrB1. The substrate used most extensively in this study had Aba appended to SyrB1 (Aba-S-SyrB1), which effectively replaced the side chain hydroxyl group of Thr-S-SyrB1 by hydrogen.

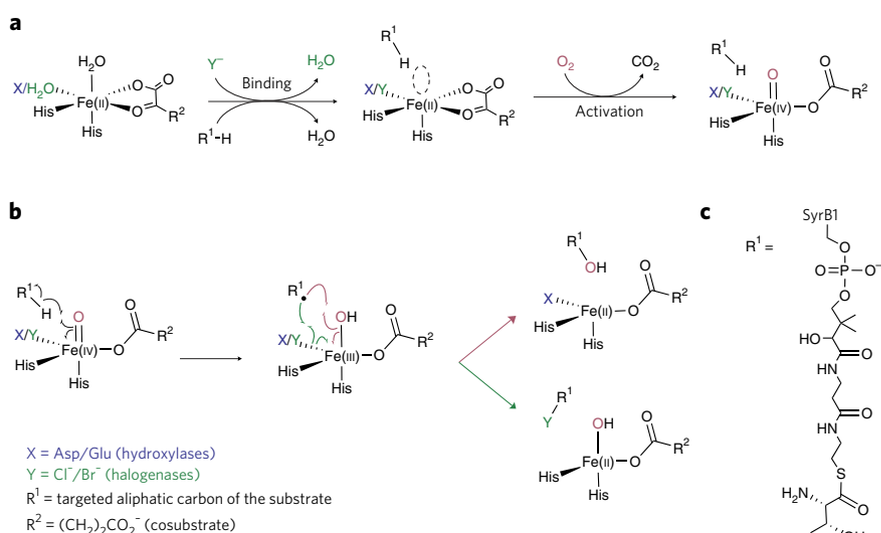
sites arranged so as to disfavor HO \bullet rebound to the substrate radical after abstraction of H \bullet by the ferryl oxygen and an open coordination site *cis* to the oxygen that is more favorably disposed for transfer. We considered that these features might confer capabilities for ferryl-mediated radical group transfer reactions beyond the known halogenation and hydroxylation outcomes, including, perhaps, installation of nitrogenous functional groups.

RESULTS

Demonstration of azidation and nitration products

Analysis by LC/MS of SyrB2 reactions carried out in the presence of the anions N $_3^-$ and NO $_2^-$ revealed that the wild-type halogenase can indeed mediate unprecedented azidation and nitration reactions upon a completely unactivated aliphatic carbon (Fig. 2 and Supplementary Results, Supplementary Fig. 1). We selected L-2-aminobutyrate (Aba; Fig. 2) appended to SyrB1 (Aba-S-SyrB1) as the ideal substrate for this analysis for two reasons. First, we anticipated that it would provide the optimal balance between proficiency in the H \bullet abstraction step, which varies in the order Thr-S-SyrB1 < Aba-S-SyrB1 < Nva-S-SyrB1, and propensity of the substrate radical to couple with the *cis* ligand rather than the HO \bullet , which varies in the opposite order²⁴ (Supplementary Fig. 2). Second, we could more easily obtain standards for the expected 4-NO $_2$ -Aba (1) and 4-N $_3$ -Aba (2) products (Online Methods) than for the corresponding derivatives of the native Thr-S-SyrB1 substrate. We used remotely deuterium-labeled Aba (3,3-[2 H]₂-Aba or 3-*d*₂-Aba) to prepare the Aba-S-SyrB1 substrate (giving 3-*d*₂-Aba-S-SyrB1) so that MS signatures of the enzymatic products would be shifted by +2 units and the standard compounds of natural isotopic abundance could be used as LC/MS internal standards. We also analyzed control reactions lacking α KG ($-\alpha$ KG) to distinguish peaks associated with authentic products of the SyrB2 reaction from those associated with other reaction components and contaminants. The addition of the unlabeled Aba standard permitted quantification of the 3-*d*₂-Aba-S-SyrB1 substrate that was consumed in the complete reactions (+ α KG) (Fig. 2a,b). Single-ion monitoring (SIM) LC/MS chromatograms revealed a diminished intensity of the peak for the 3-*d*₂-Aba released from the substrate relative to that of the unlabeled Aba standard, indicating that the substrate was indeed consumed in the complete reactions. This substrate depletion was associated with formation of products having the *m/z* values expected for the 4-NO $_2$ -3-*d*₂-Aba ([M+H]⁺ = 151.1) and 4-N $_3$ -3-*d*₂-Aba ([M+H]⁺ = 147.1) products (Fig. 2a-d). The peaks for these products had the same LC elution times as the authentic 4-NO $_2$ -Aba and 4-N $_3$ -Aba standards detected in the *m/z* = 149.1 and *m/z* = 145.1 SIM chromatograms. Their proper *m/z* values, co-elution with authentic standards and absence from the chromatograms of the $-\alpha$ KG control samples strongly suggested that the peaks arose from the desired nitration and azidation products.

These assignments were confirmed by both SIM chromatograms (Fig. 2c,d) and MS/MS fragmentation chromatograms



(Supplementary Fig. 1a,b) of samples prepared with isotopically labeled anions. In the reaction with NO $_2^-$, the use of Aba-S-SyrB1 substrate prepared with unlabeled Aba resulted in a product with the same *m/z* as the standard, and, consequently, no peak was seen at *m/z* = 151.1 corresponding to the 4-NO $_2$ -3-*d*₂-Aba product (Fig. 2c). The chromatograms from this reaction exposed a peak from a contaminant eluting just ahead of 4-NO $_2$ -Aba and having the same *m/z* (152.1) as 4- 15 NO $_2$ -3-*d*₂-Aba. The contaminant peak could thus be properly accounted for in the actual experiment. The corresponding MS/MS chromatograms (Supplementary Fig. 1a) were free from this contaminant, owing to the specificity of the fragmentation used for detection. Use of the 3-*d*₂-Aba-S-SyrB1 substrate gave a peak at the proper mass and elution time for 4-NO $_2$ -3-*d*₂-Aba (Fig. 2c), as also shown in Figure 2a. The abolition of this peak upon use of 15 NO $_2^-$ (98% isotopic enrichment) in place of the natural isotopic abundance anion and the appearance of the expected +1 peak (*m/z* = 152.1) eluting just after and largely resolved from the peak of the contaminant (Fig. 2c) confirmed the attachment of the nitrogenous anion to the Aba-S-SyrB1 substrate, an unprecedented C-N coupling reaction. Analogously, in the N $_3^-$ reaction, the use of unlabeled Aba-S-SyrB1 substrate and the anion with natural isotopic abundance resulted in 4-N $_3$ -Aba product (2) indistinguishable from the standard, and no peak for 4-N $_3$ -3-*d*₂-Aba was seen (Fig. 2d). Use of the 3-*d*₂-Aba-S-SyrB1 substrate and natural abundance N $_3^-$ resulted in the expected peak at *m/z* = 147.1 (Fig. 2b). Replacement of the anion by 15 N 14 N $_2^-$ (98% isotopic enrichment) eliminated this peak and gave rise to the expected +1 peak, which was absent from the reaction with natural isotopic abundance N $_3^-$ (Fig. 2d). These data confirmed the enzymatic coupling also of N $_3^-$ to Aba-S-SyrB1. Nitration and azidation at the C4 methyl (rather than C3 methylene) of the appended Aba was implied both by co-elution of the enzymatic NO $_2^-$ and N $_3^-$ -Aba products with the 4-modified standards and, more definitively, by the *m/z* value of the product: attachment at C3 of 3-*d*₂-Aba-S-SyrB1 would have given a product with *m/z* = 150.1, rather than 151.1, as a result of the loss of 2 H from C3, rather than 1 H from C4, in the C-H activation step.

Yields of the nitration and azidation products

We estimated the yields of the C-N coupling products from the LC/MS SIM and LC/MS/MS fragmentation chromatograms (Online Methods). The relative areas of the peaks for deuterium-labeled products and unlabeled standards in Figure 2a,b and Supplementary Figure 1 afforded estimates of the absolute concentrations of the 4-NO $_2$ -3-*d*₂-Aba and 4-N $_3$ -3-*d*₂-Aba products recovered in the small-molecule fractions after extended incubation to

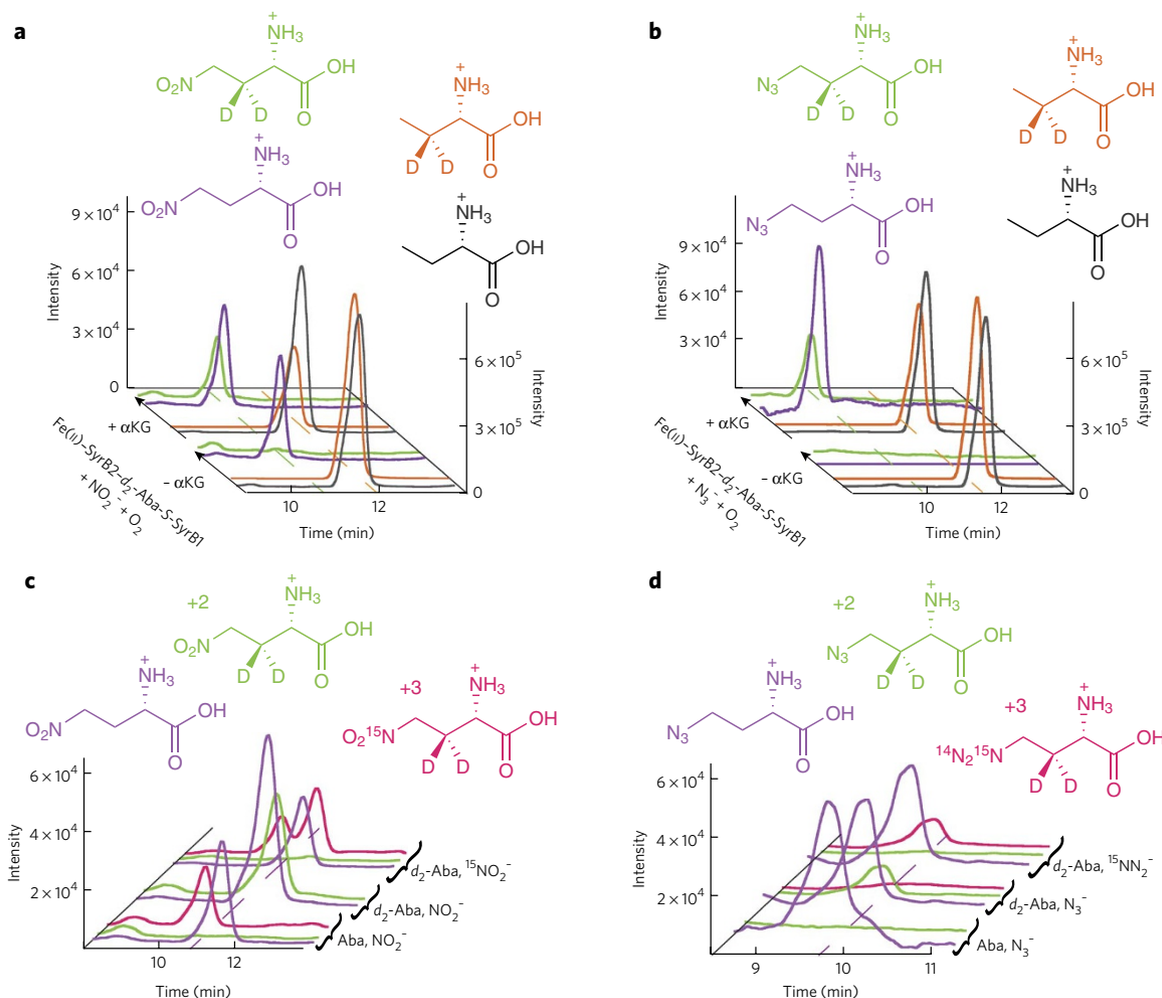


Figure 2 | LC/MS analysis demonstrating the new C-N coupling reactions catalyzed by SyrB2. Panels **a** and **c** illustrate detection of the nitration product (**1**) and **b** and **d** the azidation product (**2**). (**a,b**) Single-ion chromatograms at the m/z values for deuterium-labeled substrate (orange) and unlabeled standard (gray) and for the deuterium-labeled enzymatic nitration/azidation product (green) and unlabeled standard (purple) are shown. The scaling applied to match the intensities of the peaks for the labeled *Aba* substrate and its unlabeled standard in the $-\alpha\text{KG}$ control reactions reveals consumption of the labeled substrate in the complete ($+\alpha\text{KG}$) reaction by diminution of the orange peaks at back. The 4- NO_2 -3- d_2 -*Aba* (**a**, green) and 4- N_3 -3- d_2 -*Aba* (**b**, green) products co-eluting with the unlabeled synthetic standards (purple) are formed only in the presence of αKG . The unlabeled 4- N_3 -*Aba* standard was omitted from the $-\alpha\text{KG}$ control sample (**b**, purple at front) to permit baseline correction of the corresponding chromatogram from the $+\alpha\text{KG}$ reaction sample. The single-ion chromatograms in **c** and **d** definitively show the attachment of the anions in the detected products by the elimination of the '+2' peaks for 4- $^{14}\text{NO}_2$ -3- d_2 -*Aba* and 4- $^{14}\text{N}_3$ -3- d_2 -*Aba* (green) and appearance of the '+3' peaks for 4- $^{15}\text{NO}_2$ -3- d_2 -*Aba* and 4- $^{15}\text{NN}_2$ -3- d_2 -*Aba* (magenta) upon use of the ^{15}N -labeled anion. Reaction conditions, sample preparation and LC/MS methods are provided in Online Methods.

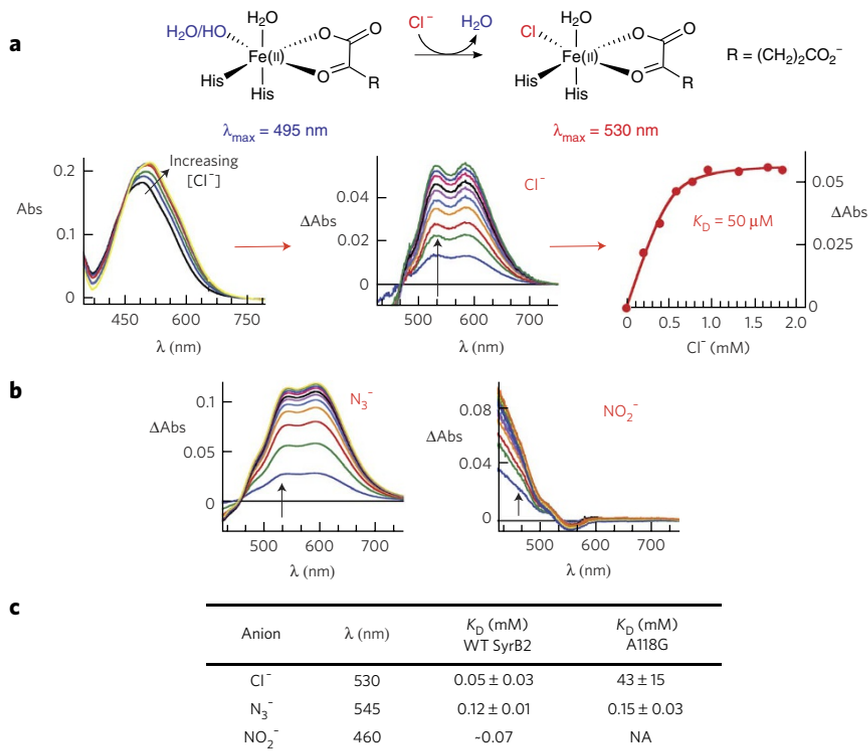
allow for their release from SyrB1 by hydrolysis of the thioester linkage. Comparison of the areas of the peaks for the 3- d_2 -*Aba* recovered from the carrier protein in the complete reaction and $-\alpha\text{KG}$ control samples to the area of the peak for the unlabeled *Aba* internal standard afforded estimates of both the quantity of 3- d_2 -*Aba* recovered in the workup in the absence of any substrate consumption ($-\alpha\text{KG}$) and the quantity consumed in the complete reaction ($+\alpha\text{KG}$). From these values, we obtained yields of $5 \pm 2\%$ of the total *Aba* and $8 \pm 2\%$ of the *Aba* consumed (values represent mean and s.d. from three trials) for the nitration reaction; the corresponding azidation yields were $1 \pm 0.4\%$ and $4 \pm 2\%$ (four trials), respectively. We consider the yields relative to the consumed substrate to be the more relevant metric because unprocessed substrate presumably remains available for conversion in additional turnovers. Consistent with the reasoning from our previous study²⁴, *Aba*-SyrB1 seemed to be more efficiently nitrated than either Thr-SyrB1 or Nva-SyrB1, although these other two substrates did also yield detectable quantities of C-N coupling products (Supplementary Fig. 3).

Anion triggering of halo-, nitrito- and azido-ferryl formation

To establish that the new aliphatic nitration and azidation reactions proceed by the expected ferryl-mediated radical group transfer mechanism, we interrogated the requisite steps in the mechanism (steps i-iv in Abstract) individually. After observing a visible color change upon addition of Cl^- to a solution of the SyrB2-Fe(II)- αKG complex, we discovered that binding of the native anion could be monitored directly by a shift in the Fe(II) \rightarrow αKG metal-to-ligand charge transfer (MLCT) band at ~ 500 nm (Fig. 3a), which is a well-established hallmark of enzymes in this class^{3,30}. Similar perturbations caused by addition of various non-native anions (Fig. 3b and Supplementary Fig. 4) demonstrated both the versatility of the anion-binding site and the robustness of this previously unknown spectral perturbation for assessing anion binding. Binding of the nitrogenous anions N_3^- and NO_2^- could also readily be monitored by this assay (Fig. 3b), although the spectral change associated with addition of NO_2^- was quite different from those associated with the other anions, owing in part to its intrinsic UV absorption³¹.

Figure 3 | Binding of chloride, azide and nitrite to the SyrB2-Fe(II)- α KG complex monitored by perturbations to the absorption spectrum from the Fe(II) \rightarrow α KG MLCT transition.

(a) Summary of the basis for the perturbation (top) and the progression in the analysis from experimental spectra (left) to difference spectra (middle) to titration curve (right) for the native Cl^- . Spectra were collected after each addition of an aliquot of the concentrated, buffered Cl^- solution. For each experimental spectrum, correction was made for dilution by the Cl^- solution, the spectrum of the starting solution was subtracted from the dilution-corrected spectrum, and the absorbance (Abs) at 800 nm was set to zero. (b) The corresponding difference spectra for the titrations with N_3^- (left) and NO_2^- (right) are shown. (c) K_D values determined by plotting the change in absorbance (Δ Abs) at an appropriate wavelength (indicated by the arrows in a and b) versus [anion] and fitting the equation for a hyperbolic or quadratic curve to the data, as appropriate. Difference spectra and titration curves used to determine K_D values reported in c for the A118G variant are shown in **Supplementary Figures 10** and **11**. WT, wild type.



Plots of the extent of spectral change as a function of the anion (Y^-) concentration (Fig. 3a and **Supplementary Fig. 5**) provided estimates of the equilibrium dissociation constants (K_D values) of the anions from the SyrB2-Fe(II)- α KG- Y^- complex. The apparent rank order of affinities for the anions that caused a measurable shift in the MLCT band was $\text{Cl}^- > \text{NO}_2^- > \text{CN}^- \sim \text{N}_3^- \sim \text{OCN}^- > \text{Br}^- \sim \text{HS}^- > \text{HCO}_2^-$ (there was no evidence for binding of SCN^- , CH_3S^- , I^- , NO_3^- or F^- ; data not shown), but the K_D values for the competent anions spanned only a factor of approximately ten (Fig. 3c and **Supplementary Table 1**). The anion site was thus found to be quite tolerant of differences in shape, steric bulk and electronic structure.

By interrogating the O_2 reactivity of the SyrB2 complex with its aminoacyl-SyrB1 substrate in the absence or presence of an added anion, we discovered an important but previously unrecognized functional feature of the halogenase: binding of the anion profoundly activates the enzyme and its Fe(II) cofactor

for reaction with O_2 , just as binding of the aminoacyl carrier protein substrate was previously shown to ‘trigger’ formation of the key ferryl intermediate¹⁶. In the reaction with either the native, chlorination-predominant Thr-S-SyrB1 substrate or the hydroxylation-predominant Nva-S-SyrB1 substrate, development of the UV absorption and Mössbauer quadrupole doublet features of the chloroferryl complex was impeded or eliminated in reactions from which Cl^- was omitted and was restored upon addition of the native anion (Fig. 4a and **Supplementary Figs. 6–8**). The small, faster phase in the Abs_{320} kinetic trace of the Thr-S-SyrB1 reaction lacking added anion (Fig. 4a) suggested the presence of contaminating Cl^- despite extensive dialysis intended to remove it, a deduction consistent with our detection by LC/MS of chlorinated products even in reactions to which Cl^- was not intentionally added (described below). Difficulty in removing Cl^- has been reported in other studies on aliphatic halogenases²⁵. The non-native halide Br^- and the nitrogenous anions N_3^- and NO_2^- all also triggered activation of O_2 (Fig. 4b and **Supplementary Fig. 9d–i**),

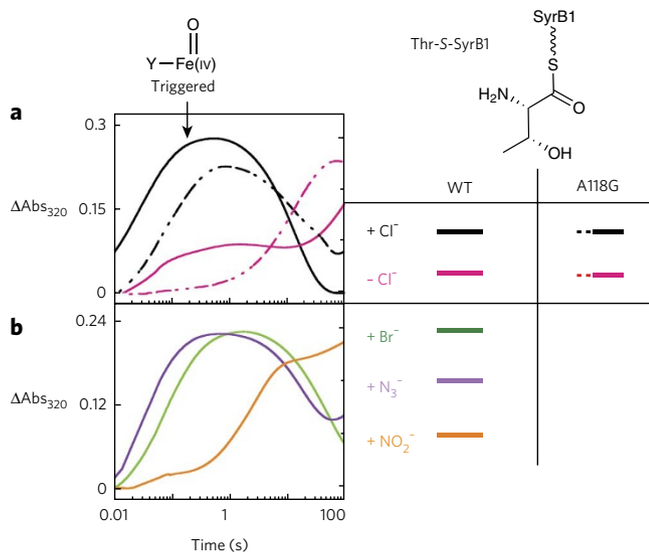
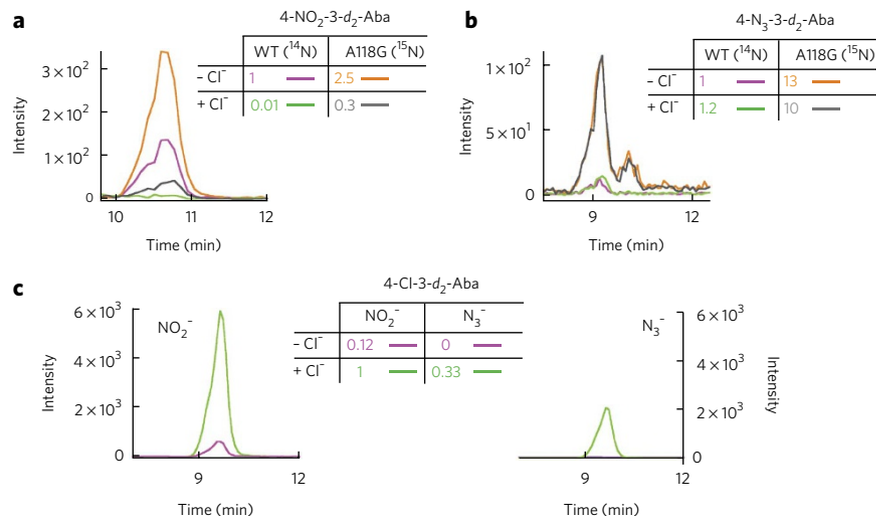


Figure 4 | Triggering of O_2 addition and ferryl formation in SyrB2 by binding of the anion.

(a,b) Development of absorption at 320 nm, previously shown to reflect formation of the ferryl complex, was monitored after an O_2 -saturated buffer solution was mixed in the stopped-flow apparatus with an O_2 -free solution of either the SyrB2-Fe(II)- α KG-Thr-S-SyrB1 complex (solid traces) or the SyrB2^{A118G}-Fe(II)- α KG-Thr-S-SyrB1 complex (dashed traces) containing no added anion (magenta) or 50 mM added Cl^- (black) (a) or 50 mM Br^- (green), 0.20 mM N_3^- (purple) or 1.0 mM NO_2^- (orange) (b). In a, comparisons of the solid black trace to the solid magenta trace and the dashed black trace to the dashed magenta trace reveal the triggering effect of Cl^- in wild-type (WT) SyrB2 and the A118G variant, respectively. The minor fast phase in the solid magenta trace reflects the presence of contaminating Cl^- . In b, the major rise phase in each of the three traces is faster than the slower rise phase in the solid magenta trace in a, illustrating the triggering effects of the non-native anions. Mössbauer spectra of samples freeze-quenched at the reaction times of maximum ΔA_{320} are shown in **Supplementary Figure 6** and demonstrate development of quadrupole-doublet features with parameters characteristic of ferryl complexes.

Figure 5 | Enhanced efficiency of C-N coupling in the A118G variant of SyrB2. (a-c)

The LC/MS/MS chromatograms monitor collision-induced transitions associated with the nitration (a), azidation (b) and chlorination (c) products formed in the reactions of the wild-type (WT) and A118G variant SyrB2. In a and b, reactions of the wild-type protein in the presence of the natural-abundance anion and the A118G variant protein in the presence of $^{15}\text{NO}_2^-$ or $^{15}\text{N}^{14}\text{N}_2^-$ were combined and co-injected. Purple and orange show reactions containing no added Cl^- ; green and gray show reactions containing 1 equivalent per SyrB2 (or SyrB2^{A118G}) added Cl^- . Comparison of the orange and purple traces in a illustrates that the nitration yield of the A118G variant is ~2.5-fold greater than the yield of the wild-type enzyme in the absence of added Cl^- ; comparison of the gray and green traces implies a ~30-fold greater nitration yield for the A118G variant in the presence of 1 equivalent of added Cl^- . Cross comparisons of the orange and gray or purple and green traces illustrate the ability of added Cl^- to suppress the nitration reaction in both wild-type SyrB2 and the A118G variant. Comparison of the orange and purple traces in b illustrates that the azidation yield of the A118G variant is ~13-fold greater than yield of the wild-type enzyme in the absence of added Cl^- . Comparisons of the orange and gray or purple and green traces illustrate the inability of 1 equivalent of added Cl^- to suppress the azidation reaction in either wild-type or A118G variant SyrB2. Shown in c are chromatograms for the chlorination product showing that addition of Cl^- (green) enhances the chlorination yields in both the reaction with NO_2^- (left) and the reaction with N_3^- (right) compared to the corresponding reactions without added Cl^- (purple).



albeit with varying efficacy. Among the non-native anions, N_3^- was the most effective, and NO_2^- was the least effective. For each of the non-native anions, Mössbauer quadrupole doublet features consistent with the accumulation of one or more Fe(IV) complexes could be observed in freeze-quenched samples of the reaction triggered by the anion (Supplementary Fig. 6)²³. These observations established that installation of the nitrogenous anions proceeds by the same mechanism as the native halogenation reaction. They further established anion triggering of O_2 activation as a control mechanism employed by SyrB2 in its native function. The differential triggering of O_2 activation by different anions hints at the possibility of engineering the enzyme not only for improved binding selectivity for a given non-native anion but also for improved triggering efficacy by that anion, which could potentially provide an additional tool for enhancing efficiency of new group transfer outcomes.

Rational improvement of C-N coupling efficiency

The ability to install nitrogenous functional groups, in particular the azido group, onto aliphatic carbon centers could enable new methods in the field of chemical biology. Indeed, the so-called 'click' and Staudinger ligation reactions, already widely used in the field, rely on the unique and 'bioorthogonal' reactivity of the azido group³²⁻³⁴. Development of an azidation catalyst useful for *in vivo* applications would be facilitated by improvement upon the modest yields observed here and might require modification of the enzyme to cope with the relatively high concentrations of Cl^- in biological milieu³⁵. Unlike the native Cl^- , neither N_3^- nor NO_2^- is spherical, and each could potentially experience steric clashes within a pocket evolved to bind Cl^- selectively. We reasoned that substitutions opening space in the anion-binding site of SyrB2 might, therefore, improve the enzyme as a catalyst for C-N coupling reactions by diminishing affinity for the native Cl^- , attenuating the thermodynamic discrimination against binding of the complex anions, perturbing the relative triggering efficacies of the native and non-native anions, permitting binding in a geometry more favorable for radical group transfer or some combination of these four effects. The A118G substitution effectively replaces a methyl group in this pocket by hydrogen. This substitution resulted in ~1,000-fold diminution in the affinity for the native Cl^- without noticeably affecting the affinity for N_3^- (Fig. 3c

and Supplementary Figs. 10 and 11). In addition, anion triggering of O_2 addition was retained in the A118G variant protein (Fig. 4a).

Trials comparing intensities of LC/MS/MS peaks from reaction products generated by the wild-type and variant enzymes under identical reaction conditions showed that, indeed, the quantities of both C-N coupling products were invariably greater with the A118G variant (Supplementary Fig. 1a,b), and so the extent of enhancement associated with the substitution and impact of the substitution on the ability of added Cl^- to suppress the C-N coupling outcomes were accurately assessed in an isotopic ratio experiment (Fig. 5). In this experiment, the products from the wild-type and A118G variant enzymes were distinguished by use of the isotopologous anions $^{14}\text{NO}_2^-$ or $^{14}\text{N}_3^-$ with the wild-type enzyme and $^{15}\text{NO}_2^-$ or $^{15}\text{N}^{14}\text{N}_2^-$ with the A118G variant. After being allowed to proceed to completion, the reaction samples were combined and analyzed by LC/MS/MS. Under the conditions examined (in a single trial), the A118G substitution was found to enhance the yield of nitration by ~2.5-fold (Fig. 5a) and the yield of azidation by ~13-fold (Fig. 5b) in the absence of added Cl^- . An azidation yield of greater than 20% of the 3-d₂-Aba consumed was estimated for this experiment. For the more efficient nitration reaction, yields as high as 52 ± 8% of the 3-d₂-Aba consumed (22 ± 4% of the initial 3-d₂-Aba present) were observed (values represent the mean and range of two experiments in Supplementary Fig. 1). Moreover, nitration by the A118G variant protein was more robust to suppression by Cl^- than nitration by the wild-type enzyme. In the isotopic ratio experiment, the level of contaminating Cl^- was sufficient to support chlorination in competition with nitration (Fig. 5c), and the peak intensities implied similar yields of the two products under these conditions (Fig. 5a,c). In the presence of one equivalent (with respect to SyrB2) of additional Cl^- , nitration by the wild-type enzyme was almost completely suppressed, whereas nitration by the A118G variant still proceeded to a detectable extent (Fig. 5a). Thus, under these conditions, the variant SyrB2 was more efficient than the wild-type enzyme by ~30-fold. The azidation reaction was observed to be intrinsically more robust to Cl^- suppression than the nitration reaction, even in the wild-type enzyme. With just the contaminating Cl^- present, chlorination was undetectable in the presence of the 2.5 equivalents (per SyrB2) of N_3^- that was added in the reaction (Fig. 5c). In the presence of one equivalent of additional Cl^- along with the N_3^- , chlorination was detected (Fig. 5c), but the yield of

the azidation product was not detectably diminished for either the wild-type or variant SyrB2 (Fig. 5b).

DISCUSSION

To assess the virtues and limitations of the newly demonstrated reactions, it is relevant first to compare the mechanistic strategy used by SyrB2 to those of other known enzymatic and chemical C-N coupling reactions. A recently reported nitration of an aromatic carbon mediated by a cytochrome P450 bears some resemblance to the SyrB2-mediated nitration reaction³⁶, but the aromatic nitration almost certainly does not proceed by initial C-H cleavage, as group transfers to aliphatic carbons must. Similarly, C-N coupling by a streptavidin-based artificial Pd-containing metalloenzyme targets an olefinic carbon³⁷, a chemically less challenging transformation. A recently reported intramolecular amination of a benzylic carbon by an engineered cytochrome P450 on an unnatural substrate having an adjacent photo-reactive aryl azide functionality to generate an iron-nitrenoid intermediate is perhaps more closely related, but it also does not target a completely unactivated aliphatic C-H bond in the presence of inherently more reactive targets, as the SyrB2 reactions do¹⁷. Likewise, the several new chemical strategies for coupling nitrogen to aliphatic carbon centers (for example, aminations) that have recently been developed^{38–41} have certain inherent limitations. In some cases, the outcomes are intramolecular couplings directed by proximity engineered into the substrates^{39–41}. In others, the C-H target is partially activated^{38–40}. Most importantly, the intimate involvement of the nitrogen atom in the C-H-cleaving iron-nitrenoid intermediates initiating all of these reactions limits their potential scope.

The strategy used by SyrB2 in its native halogenation and now nitration and azidation reactions has different limitations. First, all of the halogenases discovered to date require a carrier-protein appended substrate (Fig. 1c and Supplementary Fig. 2a). This requirement limits the substrate scope to amino acids that the corresponding charging ('adenylation') function, which in SyrB1 resides in the N terminus of the same polypeptide, will append to the phosphopantetheine cofactor of the carrier protein or domain. However, our previous work showed that SyrB1 is, fortuitously, relatively promiscuous and will self-charge multiple amino acids with diverse side chains¹⁶. Moreover, the two other aminoacyl-SyrB1 substrates (Thr-S-SyrB1 and Nva-S-SyrB1) in the series of three that was used to establish the dominant role of substrate positioning in controlling the reaction outcome²⁴ are also both detectably nitrated. This observation implies that, whereas proper substrate positioning is important also for the nitration outcome, the requirement is not so stringent as to limit the scope of the reaction to a single substrate. Moreover, modification of adenylation modules for altered amino acid specificity has been achieved^{42,43} and could further expand the substrate scope. Finally, there is no *a priori* reason that the mechanistic strategy could not be extended to small-molecule substrates (in either natural or engineered enzymes), a possibility that we continue to explore. The second limitation of the reactions in their current forms is the modest yields. Here, the ability of a single, rationally chosen substitution to enhance the binding specificity of the desired anion, augment the yields of both reactions and confer robustness to suppression by the native anion bodes well for the potential to evolve Fe/αKG oxygenases for such alternative outcomes.

Despite their present limitations, the new C-N coupling reactions extend an enzymatic strategy for functionalization of unactivated carbon centers that is fundamentally distinct from the other chemical and enzymatic precedents and is, at least potentially, more versatile. Given that the requisite initiating H• abstraction is delegated to the common oxo ligand of the ferryl intermediates, the scope of *cis*-coordinated anionic ligands that might conceivably be coupled is broad. Although we have not yet detected products resulting from the transfer of other nonhalide ligands,

the promiscuous binding and triggering of O₂ activation by different anions and the robustness of ferryl formation in their presence (Supplementary Fig. 6) suggest that other reactions will be possible. Moreover, all of the demonstrated and envisaged couplings are intermolecular and expected to be exquisitely controlled by the active site architecture. Indeed, it is precisely this absolute chemo-, regio- and stereospecificity that can be accomplished by enzymes by virtue of their complex substrate binding sites that has made their use in synthetic chemistry seductive and, more recently, practically important^{44,45}. These considerations provide the impetus for ongoing efforts to develop and discover enzymatic catalysts for aliphatic C-N coupling reactions.

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METHODS

Methods and any associated references are available in the [online version of the paper](#).

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Author contributions

M.L.M. prepared reagents, designed and carried out experiments, analyzed data and composed and edited the manuscript. W.-c.C. prepared reagents, designed and carried out experiments, analyzed data and edited the manuscript. A.P.L. prepared reagents, carried out experiments and analyzed data. L.A.M. prepared reagents, carried out experiments and analyzed data. C.K. designed experiments, analyzed data and edited the manuscript. J.M.B. Jr. designed experiments and composed and edited the manuscript.

Competing financial interests

The authors declare no competing financial interests.

Additional information

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ONLINE METHODS

Materials. Sodium bromide (NaBr), sodium sulfide nonahydrate ($\text{Na}_2\text{S}\cdot 9\text{H}_2\text{O}$), sodium cyanide (NaCN), sodium nitrite (NaNO_2), sodium methyl thiolate (NaSCH_3), sodium thiocyanate (NaSCN), sodium fluoride (NaF), L-glycine and ammonium formate (NH_4HCO_2) were purchased from Sigma-Aldrich (St. Louis, MO). Sodium azide (NaN_3) was purchased from EMD Chemicals, Inc. (Gibbstown, NJ). Potassium cyanate (KOCN) was purchased from Alfa Aesar (Ward Hill, MA). Sodium nitrate (NaNO_3) was purchased from J.T. Baker (Phillipsburg, NJ). (2S)-2-Amino-4-azido-butanoic acid (4- N_3 -Aba) was obtained from Toronto Research Chemicals Product List (North York, Canada). All of the other chemicals (except for 4- NO_2 -Aba, which was synthesized as summarized below) were obtained from sources previously noted^{16,24}. Purchased reagents were used as received without purification.

General synthetic methods. NMR spectra were recorded on Bruker 360 MHz spectrometers. Chemical shifts are reported in p.p.m. using a residual solvent peak as an internal standard. *J* values are reported in Hz.

Synthesis of 4- NO_2 -Aba. 4- NO_2 -Aba was synthesized as previously reported⁴⁶. Lithium diisopropylamide (3.5 mL of 0.5 M solution in THF, 1.75 mmol) was added dropwise to a solution of *N*-diphenylmethyleneglycine *t*-butyl ester (500 mg, 1.7 mmol) in THF (5 mL) at -78°C , and the solution was stirred for 1.5 h. Nitroethylene (3.4 mL of 0.5 M solution in THF, 1.7 mmol) was then added to the reaction solution at the same temperature. After stirring for 1 h, the reaction mixture was gradually brought to room temperature and quenched by addition of water (20 mL) followed by ethyl acetate (20 mL). The combined organic layer was washed with brine and concentrated under vacuum. The product was purified by silica gel chromatography using hexanes/ethyl acetate (4:1) as an eluent to obtain the pure product in 40% yield. The $^1\text{H-NMR}$ spectrum of the *t*-butyl 2-diphenylmethyleneimino-4-nitrobutanoate was identical to that reported in the literature⁴⁶. Global deprotection was carried out by adding 1 N HCl (5 mL) to the *t*-butyl 2-diphenylmethyleneimino-4-nitrobutanoate (37 mg, 0.1 mmol) and stirring at room temperature overnight. The reaction mixture was extracted with ethyl ether (3 \times 5 mL), and the aqueous layer was subjected to lyophilization to obtain the final product. $^1\text{H-NMR}$: (360 MHz, D_2O) δ 4.03 (t, *J* = 7.0 Hz, 2H), 2.39–2.49 (m, 3H) (Supplementary Fig. 12).

Protein purification and substrate assembly. These steps were performed as previously described¹⁶.

Anion titrations and analysis to extract dissociation constants. Analogously to published titrations of SyrB2 with Fe(II) ⁴⁷, a concentrated, buffered, O_2 -free stock of the relevant anion was added in small aliquots to an O_2 -free solution of the SyrB2- Fe(II) - αKG complex or SyrB2^{A118G}- Fe(II) - αKG complex containing 0.75 mM Fe(II) , 5 mM αKG and 1 mM SyrB2 or SyrB2^{A118G}. Spectra were acquired after each addition until the absorption spectrum stopped changing (after several min).

Stopped-flow absorption experiments. The kinetics of formation and decay of the ferryl intermediate at 5°C were monitored by its absorbance at 320 nm (ΔAbs_{320}). An O_2 -saturated buffer solution (20 mM Na-HEPES, pH 7.5) was mixed in the stopped-flow apparatus with an equal volume of an O_2 -free solution containing 0.4 mM SyrB2 (or SyrB2^{A118G}), 0.3 mM Fe(II) , 10 mM αKG and 0.9 mM of the aminoacyl-SyrB1 substrate. The anion concentrations in the experiment of Figure 4 were 50 mM Cl^- (Fig. 4a) and 0.2 mM N_3^- or 1 mM NO_2^- (Fig. 4b). The reactant concentrations in the experiments depicted in Supplementary Figures 7 and 9 were equivalent. Solely to clarify presentation of the stopped-flow data, the Steineman smoothing function supplied as a standard feature of the Kaleidagraph (Synergy Software, Reading, PA) analysis software was applied to all of the kinetic traces.

Preparation of Mössbauer samples and spectral analysis. Unless otherwise noted, the final concentrations for all Mössbauer samples in Supplementary Figures 6 and 8 were: [^{57}Fe] \sim 0.4 mM, [SyrB2] \sim 0.5 mM, [aminoacyl-SyrB1] \sim 0.8 mM, [αKG] = 5 mM, [NaCl] = 50 mM, [NaCN/ $\text{NaNO}_2/\text{Na}_2\text{S}$] = 1 mM and [N_3^-] = 0.4 mM. Using regression analysis, 4,2-K/zero-field spectra of the reactant SyrB2- Fe(II) - αKG -Y⁻-aminoacyl-SyrB1 complexes were analyzed assuming one or two symmetrical quadrupole doublets. The fits returned parameters typical of high-spin Fe(II) complexes. Spectra of samples that were reacted with O_2 exhibited features with parameters typical of high-spin Fe(IV) ,

which were assigned to the Y- $\text{Fe(IV)}=\text{O}$ intermediates, and features from an unreacted high-spin Fe(II) complex (or complexes). The high-energy lines of the quadrupole doublets assigned to Fe(IV) and Fe(II) were well resolved, but the associated low-energy lines were unresolved. To overcome this lack of resolution and determine the Mössbauer parameters of the Fe(IV) complexes, we analyzed experimental spectra of freeze-quenched samples in one of three ways. First, we subtracted the appropriate contribution of the experimental spectrum of the corresponding Fe(II) -containing reactant complex, as judged by comparison of the intensity of the resolved high-energy lines. This subtraction produced 'reference spectra' for the Fe(IV) complexes that were of variable quality but yielded acceptable results when the shape of the high-energy line of the Fe(II) component matched well with that from the Fe(II) reactant complex or if the contribution from Fe(II) in the spectrum of the freeze-quenched sample was small. The second method for generation of Fe(IV) reference spectra involved modeling the spectral contribution of the Fe(II) complexes as quadrupole doublets with parameters (isomer shift (δ), quadrupole splitting (ΔE_Q) and line width (Γ)) obtained from fitting of the spectra of the reactant complexes, but in varying proportions. The third method for generation of the Fe(IV) reference spectra was identical to the second method, except that the line widths of the Fe(II) quadrupole doublets were allowed to vary within 10% of the value obtained from fitting of the Fe(II) reference spectrum. The Fe(IV) reference spectra were then simulated as a single quadrupole doublet, except for that of the chloroferryl intermediate in the presence of either Thr-SyrB1 (ref. 16) or d_5 -Nva-S-SyrB1, for which the high-energy line of the Fe(IV) reveals two partially resolved peaks, indicating the presence of two ferryl complexes, as previously reported^{15,16}. The spectra of the presumed nitroferryl ($\text{O}_2\text{N-Fe(IV)}=\text{O}$) and (hydro)sulfidoferryl ($(\text{H})\text{S-Fe(IV)}=\text{O}$) complexes with d_5 -Thr-SyrB1 were rather sharp, and the positions of the low-energy lines of the various reference spectra were well determined. The azidoferryl ($\text{N}_3\text{-Fe(IV)}=\text{O}$) intermediate formed with d_5 -Thr-SyrB1 accumulated in high yield, and the lines were nearly symmetrical but broad. The spectrum of the cyanidoferryl ($\text{CN-Fe(IV)}=\text{O}$) complex was the least well resolved, owing to its lesser accumulation, differences between the Fe(II) components in the reactant and freeze-quenched samples and the breadth of the lines in the Fe(IV) reference spectrum. From these analyses, we estimated the uncertainties in the values for the Mössbauer parameters to be 0.05 mm/s for the isomer shifts and 0.10 mm/s for the quadrupole splitting parameters. These error estimates are conservative: they represent the upper limits of the observed variations in the extracted parameters for the different methods of analysis.

C-N coupling reactions and analysis by LC/MS. Reactions associated with Figure 2 were performed as follows. Aba-S-SyrB1 and 3- d_2 -Aba-S-SyrB1 were prepared by incubation of 3.0 mM HS-SyrB1, 5.0 mM Aba or 3- d_2 -Aba, 30 mM ATP and 30 mM MgSO_4 for 30 min at room temperature. Prior to their use in the SyrB2 reactions, assembled substrates were purified from small molecules as previously described²⁴. The final concentrations in the SyrB2 reactions were 0.2 mM Fe(II) , 0.2 mM SyrB2, 0.8 mM Y^- , 0.12 mM αKG and 0.1 mM desalted Aba-S-SyrB1 or 3- d_2 -Aba-S-SyrB1. L-Glycine, an amino acid that SyrB1 cannot append to its carrier domain, was added to 10 μM as an internal standard. Overnight incubation at ambient temperature allowed the SyrB2 reaction to proceed to completion and allowed the thioester-linked amino acid substrates and products to be hydrolyzed from the phosphopantetheine cofactor. The amino acids were then purified from the free SyrB1 and SyrB2 proteins via passage through a Microcon centrifugal filter unit with a 10-kDa molecular mass cutoff (Millipore). In the control reactions lacking αKG , the typical recovery of 3- d_2 -Aba in this procedure was \sim 50%. Prior to LC/MS analysis, 25 μM Aba, 2.5 μM 4- NO_2 -Aba and/or 2.5 μM 4- N_3 -Aba were added to the filtrate, as appropriate. The concentrations of standards were chosen such that their measured peak areas were comparable to those for corresponding reaction products.

Reactions associated with Figure 5 were performed essentially as described above, with the following exceptions. After the reaction of the wild-type and A118G variant SyrB2 were allowed to reach completion, equal volumes of the two solutions were mixed. C-N coupling products generated by each enzyme could be distinguished by using isotopologic anions. Unlabeled $^{14}\text{NO}_2^-$ or $^{14}\text{N}_3^-$ was used with the wild-type enzyme and isotopically enriched $^{15}\text{NO}_2^-$ or $^{15}\text{N}_3^-$ was used with the A118G variant, necessarily associating 4- N_3^- and 4- NO_2^- - d_2 -Aba products with the reaction of the wild-type enzyme and 4- $^{15}\text{N}_3^-$ - d_2 -Aba and 4- $^{15}\text{NO}_2^-$ - d_2 -Aba products with the reaction of the A118G variant. The concentrations of reactants were 0.25 mM Fe(II) , 0.25 mM SyrB2,

1.3 mM NO₂⁻, 0.25 mM αKG and 0.13 mM 3-*d*₂-Aba-S-SyrB1 for the nitration reaction and 0.25 mM Fe(II), 0.25 mM SyrB2, 0.63 mM N₃⁻, 0.15 mM αKG and 0.13 mM 3-*d*₂-Aba-S-SyrB1 for the azidation reaction. In this case, the 3-*d*₂-Aba-S-SyrB1 substrate was prepared by incubation of 0.19 mM HS-SyrB1, 0.15 mM 3-*d*₂-Aba, 2.3 mM ATP and 2.3 mM MgSO₄ for 30 min at ambient temperature and was used in the SyrB2 reactions without desalting.

The LC was performed on a ZIC-HILIC column (Merck-Millipore). The mobile phase was a mixture of 95:5 (v/v) H₂O:CH₃CN (A) and 95:5 (v/v) CH₃CN:H₂O (B), of which both were supplemented with 0.1% formic acid to facilitate ionization. Following sample injection (10 μL) under 95% B with a flow rate of 0.4 mL/min, the mobile phase was varied linearly to 80% B over the first 10 min, held constant for 5 min, changed to 0% B over the next 20 min, increased linearly to 95% B over the following 20 min and held constant at 95% B in preparation for the next injection. In the SIM chromatograms of **Figure 2**, parent amino acid cations were monitored. In the MS/MS fragmentation chromatograms of **Figure 5**, the daughter iminium ions formed by neutral loss of CO₂ and H₂ (a combined mass of 46 a.m.u.) from the parent cations were selectively monitored (*m/z* values of the relevant transitions are in **Supplementary Table 4**). The fragmentation was detected by application of a collision energy of 3 V to the parent ion. The drying gas temperature was 350 °C with a nebulizer pressure of 40 psi and flow rate of 9 L/min. The capillary voltage was set to 4,000 V. The fragmentor voltage was set at 70 V to achieve maximal electrospray ionization for detection. Dwell times of 50 ms and 400 ms were dedicated to detection of each parent or parent-to-daughter cation transition, respectively.

Estimation of the yields of the 4-NO₂-Aba and 4-N₃-Aba products. The fact that the initial amino acid products of the SyrB2 reaction are linked as thioester to SyrB1 creates the challenge of determining the combined yield of hydrolysis off the carrier protein and recovery in the workup. The immediate 4-NO₂-3-*d*₂-Aba-S-SyrB1 or 4-N₃-3-*d*₂-Aba-S-SyrB1 products were not readily accessible either by synthesis or by enzymatic attachment of the synthetic amino acid products to the phosphopantetheine cofactor of the carrier domain by the adenylation domain (they seem not to be accepted for charging). Recovery of the 3-*d*₂-Aba substrate off the SyrB1 carrier protein was assessed in a control reaction from which αKG was omitted. In the control, comparison of

the peak for the 3-*d*₂-Aba released from SyrB1 to that for the unlabeled free Aba amended to the sample afforded absolute quantification of the released 3-*d*₂-Aba and its fractional recovery (40–60% overall). Likewise, this same comparison for the reaction sample containing αKG permitted quantification of unreacted 3-*d*₂-Aba recovered in the complete reaction. The difference between these concentrations gave the concentration of 3-*d*₂-Aba consumed in the reaction. Comparison of the peak for the 4-NO₂-3-*d*₂-Aba or 4-N₃-3-*d*₂-Aba amino acid to that for the synthetic, unlabeled 4-NO₂-Aba and 4-N₃-Aba standard amended to the reaction afforded absolute quantification of the recovered enzymatic product. By invoking the necessary assumption that the fractional recovery of the 4-NO₂-3-*d*₂-Aba or 4-N₃-3-*d*₂-Aba amino acid from the 4-NO₂-3-*d*₂-Aba-S-SyrB1 or 4-N₃-3-*d*₂-Aba-S-SyrB1 primary product was identical to that determined for the unreacted 3-*d*₂-Aba, the yield of the product with respect to either the 3-*d*₂-Aba consumed or the total 3-*d*₂-Aba initially present in the reaction was calculated. For example, in two typical nitration reactions under conditions optimized for this reaction with the A118G variant, we detected 11 ± 1 μM 4-NO₂-3-*d*₂-Aba (reaction volume of 0.10 mL) when the recovered 3-*d*₂-Aba in the -αKG-control reaction was 49 μM and the amount of 3-*d*₂-Aba recovered in the complete reaction was 28 ± 1 μM (values represent mean and range of two measurements). The 3-*d*₂-Aba concentrations imply that ~21 μM substrate was consumed in the complete reaction. Dividing the 11 μM 4-NO₂-3-*d*₂-Aba produced by the 21 μM 3-*d*₂-Aba consumed gives a nitration yield of ~52% of the 3-*d*₂-Aba consumed. Alternatively, the ratio of the 11 μM 4-NO₂-3-*d*₂-Aba produced to the 49 μM recovered in the absence of any conversion gives a nitration yield of ~22% of the 3-*d*₂-Aba initially present in the reaction.

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