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An Iron(IV)—Oxo Intermediate Initiating L-Arginine Oxidation but Not Ethylene Production by the 2-Oxoglutarate-Dependent Oxygenase, Ethylene-Forming Enzyme

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converts carbons 1, 2, and 5 of 2OG to CO_2 and carbons 3 and 4 to ethylene, a fourelectron oxidation drastically different from the simpler decarboxylation of 2OG to succinate mediated by all other Fe/2OG enzymes. EFE also catalyzes a minor reaction, in which the normal decarboxylation is coupled to oxidation of L-arginine (a required activator for the EF pathway), resulting in its conversion to L-glutamate semialdehyde and guanidine. Here we show that, consistent with precedent, the L-Arg-oxidation (RO) pathway proceeds via an iron(IV)—oxo (ferryl) intermediate. Use of 5,5-[²H₂]-L-Arg slows decay of the ferryl complex by >16-fold, implying that RO is initiated by hydrogen-atom transfer (HAT) from C5. That this large substrate deuterium kinetic isotope effect has no impact on the EF:RO partition ratio implies



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that the same ferryl intermediate cannot be on the EF pathway; the pathways must diverge earlier. Consistent with this conclusion, the variant enzyme bearing the Asp191Glu ligand substitution accumulates \sim 4 times as much of the ferryl complex as the wild-type enzyme and exhibits a \sim 40-fold diminished EF:RO partition ratio. The selective detriment of this nearly conservative substitution to the EF pathway implies that it has unusually stringent stereoelectronic requirements. An active-site, like-charge guanidinium pair, which involves the L-Arg substrate/activator and is unique to EFE among four crystallographically characterized L-Arg-modifying Fe/ 2OG oxygenases, may serve to selectively stabilize the transition state leading to the unique EF branch.

INTRODUCTION

Iron(II)- and 2-oxoglutarate-dependent (Fe/2OG) oxygenases use a common cosubstrate (2OG) as the source of the two electrons needed to balance the four-electron reduction of dioxygen with the two-electron oxidation of an Fe(II) cofactor to an Fe(IV)-oxo (ferryl) intermediate.¹⁻⁴ The cosubstrate chelates the Fe(II) cofactor via its C1 carboxylate and C2 carbonyl oxygens, contributing to a square-pyramidal coordination sphere that is rounded out by (most often) three facially disposed amino acid ligands-two histidines and one carboxylate—provided by the conserved β -sandwich protein architecture that defines the enzyme family.⁵⁻⁷ Dioxygen adds to the open axial site of this five-coordinate complex, and the cosubstrate undergoes oxidative decarboxylation (C1-C2 bond cleavage) to produce succinate. In the process, one of the two oxygen atoms from O_2 is incorporated into the C2derived carboxylate, as the other oxygen remains bonded to iron in the ferryl complex. The ferryl intermediate most often abstracts a hydrogen $(H\bullet)$ from an aliphatic carbon of the "prime" substrate (R−H) to form a fleeting Fe(III)-OH/R● intermediate.⁸ Different fates of this intermediate lead to the varied outcomes-hydroxylation, halogenation, desaturation, cyclization, ring expansion, stereoinversion, endoperoxidation,

and others—of which members of the versatile enzyme family are collectively capable.^{2,3,9,10} Most of these outcomes extract two electrons from the prime substrate, thus resetting the enzyme for another turnover.

Ethylene-forming enzyme (EFE) from pseudomonads and fungi is a unique, ambifunctional Fe/2OG oxygenase that, in its namesake, major reaction, diverges rather dramatically from the chemical logic that characterizes the enzyme family.^{11,12} In this reaction (Scheme 1, lower pathway), EFE confers all four oxidizing equivalents of O_2 to the cosubstrate, globally fragmenting it to 3 equiv of CO_2 (from carbons 1, 2, and 5) and 1 equiv of ethylene (from carbons 3 and 4). The enzyme also catalyzes a minor reaction (upper pathway) that *does* conform to the family logic, in which oxidation of 2OG to succinate is coupled to the oxidative fragmentation of L-

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Scheme 1. Reactions Catalyzed by EFE



arginine to guanidine and L-glutamate semialdehyde (which cyclizes in solution to pyrroline-5-carboxylate, P5C). Intriguingly, although it is transformed only in the minor reaction, L-Arg is still strictly required for the major reaction.^{12,13} In essence, the amino acid has distinct roles as prime substrate for the ordinary, L-Arg-oxidizing (RO) reaction and essential effector for the unique, ethylene-forming (EF) reaction.

To date, experimental investigations of EFE have been limited to structural characterization of its substrate-bound states, examination of the kinetics and ratios of products (ethylene, pyrroline-5-carboxylate, and succinate) in multipleturnover assays of wild-type (wt) and variant enzymes, and evaluation of substrate analogues for the ability to support one or both reactions.^{12–18} By all accounts, wt EFE exhibits an EF:RO partition ratio of $\sim 2-3:1$ when turning over its native substrates. Substitutions to active-site-proximal residues have, in general, proven deleterious to both reactions, but a number of those examined compromise the EF pathway more profoundly than the RO pathway. By contrast, none has been shown to increase the EF:RO partition ratio relative to that of the wt enzyme. Experiments with L-Arg analogues have suggested a remarkable specificity for the allosteric activator/ prime substrate. All analogues that have been tested are, according to these reports, poorer activators of the EF pathway.^{13,16} From the limited available mechanistic data, it has not yet been possible to answer the central mechanistic question posed by the unusual EF outcome: at which point, and by what novel step, does this pathway diverge from the more canonical RO pathway?

The conformity of the RO pathway to the characteristic Fe/ 2OG-oxygenase logic suggests a mechanism involving ferrylmediated abstraction of H• from C5 of L-Arg to initiate either (i) hydroxylation followed by C5–N6 fragmentation^{12,13} or (ii) C5–N6 desaturation followed by hydrolysis.¹⁹ Two recent X-ray crystal structures confirmed the required proximity of C5 to the metal center [*N*-oxalylglycine-coordinated Fe(II) or 2OG-coordinated Mn(II)], but no direct evidence for the targeting of C5 by a ferryl complex has been reported.^{16,17} It is less obvious that the major EF pathway should proceed through a ferryl intermediate; mechanisms with and without such a complex can be proposed. Indeed, two of the aforementioned experimental studies posited *different* ferryl complexes—in one case, a succinate-coordinated ferryl initiating ethylene production¹⁶ and, in the second case, an oxalate-coordinated ferryl forming concomitantly with ethylene¹⁷—whereas a recent computational study proposed an EF pathway not proceeding through a ferryl intermediate.²⁰

In the reactions of other Fe/2OG oxygenases, ferryl intermediates have been demonstrated on the basis of three hallmarks: (i) their increased UV absorption (typically monitored at \sim 320 nm) relative to the enzyme•Fe(II) •20G•substrate and enzyme•Fe(II)•succinate•product complexes; (ii) their characteristic Mössbauer isomer shifts ($\delta \sim$ 0.3 mm/s) and quadrupole splitting ($\Delta E_Q \sim -1 \text{ mm/s}$) parameters;^{21,22} and (iii) the large substrate deuterium kinetic isotope effects (D-KIEs) on their decay by H•/D• transfer, which results in increased intensity and longevity of these spectral signatures.^{3,8,23} The weak absorption feature of the reactant complexes at 500-530 nm ($\varepsilon \sim 200 \text{ M}^{-1} \text{ cm}^{-1}$) arising from a metal-to-ligand [Fe(II)-to-2OG] charge-transfer (MLCT) transition⁵—has also proven useful in transientkinetic monitoring of the reactions. The ferryl complexes are approximately isosbestic in this region, but the enzyme•Fe(II) •succinate•product complexes (referred to in the original work on TauD as a "second intermediate"²¹) are transparent, with the result that this optical feature provides an additional way to monitor either conversion of the reactant complex to the product complex in cases of a nonaccumulating ferryl intermediate or decay of the ferryl intermediate to the product(s) complex when the high-valent species accumulates substantially.^{24,2}

In this study, we used stopped-flow absorption (SF-Abs) and freeze-quench Mössbauer (FQ-Möss) spectroscopies along with substrate deuterium labeling, quantitative product analysis, and site-directed mutagenesis to test for a ferryl complex in either or both pathways of the EFE reaction, and we solved the structure of the authentic EFE•Fe(II)•2OG•L-Arg reactant complex. The kinetic and spectroscopic results not only confirm the expectation that the RO reaction proceeds via the canonical ferryl intermediate but also reveal that the detected complex forms after the branchpoint of the two pathways, meaning that it is not on the EF pathway. The ability of a nearly conservative Asp \rightarrow Glu ligand substitution to almost completely derail the EF pathway while barely affecting the RO pathway is consistent with unusually stringent stereoelectronic requirements for the major reaction, and structural features of the EFE reactant complex that distinguish it from those of several other L-Arg-modifying Fe/2OG

oxygenases characterized previously—most notably, the flipped orientation of L-Arg¹⁶ and resulting like-charge guanidinium pair involving its side-chain and an active-site Arg residue—suggest how the transition state leading to the unique outcome might be selectively stabilized.

RESULTS

Activity and Partition Ratio of Our Preparations of **EFE.** To verify the integrity of our preparations of *Pseudomonas* savastanoi pv phaseolicola EFE, we quantified the products of multiple-turnover reactions by liquid chromatography with mass spectrometric detection (LC-MS) and gas chromatography with flame-ionization detection (GC-FID). We detected pyrroline-5-carboxylate (P5C), a product of the RO pathway, by first reducing it to proline with sodium cyanoborohydride and then appending an Fmoc protecting group prior to LC-MS analysis. Our results corroborate previous reports^{12,13} that, with unlabeled L-Arg as substrate, $\sim 70\%$ of the 2OG consumed by wild-type EFE yields ethylene, while the remaining \sim 30% is converted to succinate (Figure 1). The rates ([product] time⁻¹ [EFE]⁻¹) of succinate and ethylene production that we measured (~15 and ~40 min⁻¹, respectively) are also consistent with published reports.^{13,14,26}



Figure 1. Yields of succinate, Fmoc-proline (P5C), and ethylene after incubation of 10 μ M wt or D191E EFE with 40 μ M (NH₄)₂Fe(SO₄)₂, 200 μ M 2OG, and 500 μ M of unlabeled L-arginine (L-Arg), perdeuterated L-arginine (d_{7} -L-Arg), or 5,5- d_{2} -L-arginine (d_{2} -L-Arg) in air-saturated 40 mM sodium HEPES (pH 7.5) buffer at 21 °C for ~10 min. 2OG was completely consumed during the reaction. P5C was derivatized to Fmoc-proline prior to quantification. Ethylene analysis was performed by GC-FID; succinate and Fmoc-proline were quantified by LC-MS. Error bars correspond to standard errors of the mean values for two experiments. Detailed procedures are provided in the Supporting Information.

Evidence for an L-Arg-Oxidizing Ferryl Intermediate in the EFE Reaction. To test for accumulation of a ferryl intermediate in the EFE reaction, we mixed the anoxic EFE•Fe(II)•2OG•L-Arg complex with O₂-containing buffer at 5 °C in a stopped-flow apparatus. Weak UV absorption developed within ~0.07 s and decayed by more than 90% within 1 s (Figure 2A, solid black circles). The loss of absorbance at 510 nm in this same time regime (Figure 2B, solid black circles) signified conversion of the reactant state to the transparent Fe(II)-containing product state(s) via a barely accumulating (presumptive) ferryl complex. Use of L-Arg bearing deuterium at C5 (here, 2,3,3,4,4,5,5-[²H₇]-L-Arg or d_7 -L-Arg) resulted in 4–5-fold increases in both the upward deflection in the ΔA_{318} -vs-time trace and the time for its return back to the initial absorbance (Figure 2A, open black circles). Article



Figure 2. Kinetic analysis of the wt and D191E EFE reactions with protium- and deuterium-bearing L-Arg substrates by stopped-flow absorption spectroscopy. The change in absorbance (A) at 318 nm and (B) at 510 nm was monitored after mixing (at 5 °C) of an anoxic solution containing 0.7 mM wt EFE (black) or D191E EFE (red), 0.6 mM Fe(II), 8 mM 2OG, and 8 mM L-Arg (filled circles) or d_7 -L-Arg (open circles) in 40 mM sodium HEPES (pH 7.5) buffer with an equal volume of the same buffer containing ~0.48 mM O₂. The solid lines are the results of global fitting of the data to a kinetic model in which two rapidly equilibrating reactant complexes react with O₂, as described in the Supporting Information and shown in Scheme S1.

Use of deuterated L-Arg also resulted in a modest decrease in the downward deflection of the ΔA_{510} -vs-time trace (Figure 2B, open black circles), but importantly, the temporal delay and diminution of the amplitude of this trace were not as profound as one would have expected in view of the markedly greater accumulation of the presumptive ferryl complex, which, as noted, absorbs significantly in the visible regime in other Fe/ 20G enzymes. This observation is explained by the facts that (i) only approximately one-third of the reaction flux (the RO pathway) is sensitive to the presence of deuterium in L-Arg, and (ii) the other two-thirds (the EF pathway) converts the reactant complex to a transparent product complex without substantial accumulation of an intervening, 510 nm-absorbing state (as explained more below). These observations are consistent with a modest but detectable accumulation of the ferryl intermediate with the protium-containing substrate and enhanced accumulation with d_7 -L-Arg as a result of a large, normal D-KIE on HAT to the high-valent complex.

To confirm the expectation that the H• donor to the presumptive ferryl complex is C5, we also examined the reaction with site-specifically C5-deuterated L-Arg $(5,5-[^{2}H_{2}]$ -L-Arg or d_{2} -L-Arg). As anticipated, the increased accumulation and slower decay seen with the d_{7} -L-Arg substrate were also observed with the specifically labeled substrate (Figure S3). The fact that the kinetics of decay with d_{2} -L-Arg and d_{7} -L-Arg were essentially identical implies that the ferryl complex cannot abstract H• from a different site with an appreciable rate constant, a conclusion supported by our inability to detect any other L-Arg-derived product in the reaction with d_{2} -L-Arg.

We next confirmed by freeze-quench Mössbauer spectroscopy that the UV-absorbing intermediate is, in fact, a ferryl complex. The spectrum of the frozen $EFE \bullet {}^{57}Fe(II) \bullet 2OG \bullet d_{7}$ -L-Arg reactant solution acquired at 4.2 K without an applied magnetic field (Figure 3A, i, solid lines) consists of a



Figure 3. Analysis of the reactions of (A) wt and (B) D191 EFE by freeze-quench Mössbauer spectroscopy. Experimental spectra (i, ii, and iv) were collected at 4.2 K with either no applied magnetic field (solid lines) or a field of 53 mT applied parallel to the γ -beam (vertical bars). In preparation of the freeze-quenched samples (spectra ii and iv), an anoxic solution containing 1.8 mM EFE (wt or D191E), 1.4 mM 57 Fe(II), 5 mM 2OG, and 5 mM d_7 -L-Arg in 40 mM sodium HEPES buffer (pH 7.5) containing 5% glycerol was mixed at 5 °C with an equal volume of O_2 -saturated (~1.8 mM O_2) buffer, and the reaction was allowed to proceed for the indicated time before being terminated by rapid freezing in the cryosolvent (2-methylbutane at \sim -150 °C). For samples of the reactant complexes (spectra i), the same anoxic protein reactant solution was manually frozen (wt, panel A) or diluted with an equal volume of O2-free buffer before being manually frozen (D191E, panel B). Spectra iii (vertical bars) result from subtraction of the experimental zero-field spectra (ii - i) to illustrate the changes that occur at early time. These difference spectra can be accounted for as summations (black lines) of either three (wt, panel A) or two (D191E, panel B) quadrupole-doublet components (colored lines) corresponding to features that form (downward) and decay (upward) over short reaction times. The colored theoretical spectra are associated with the ferryl intermediate (red), the Fe(II)containing reactant complex (blue), and the Fe(II)-containing product complex of the EF pathway (green in panel A only). The parameters used to generate these theoretical spectra are given in the text and Table S8.

quadrupole doublet with isomer shift (δ) and quadrupole splitting ($\Delta E_{\rm O}$) parameters characteristic of N/O-coordinated, high-spin Fe(II) ions ($\delta \sim 1.1 \text{ mm/s}$ and $\Delta E_0 \sim 2.6 \text{ mm/s}$).²⁷ This reactant solution was mixed at 5 °C with an equal volume of buffer containing ~ 1.8 mM O₂, and the resulting solution was frozen after 0.34 s, the reaction time at which the UVabsorbing complex was seen to accumulate maximally (Figure S4). The 4.2 K/zero-field spectrum of this sample (Figure 3A, ii, solid lines) exhibits a broadened doublet for the Fe(II) species (indicated by black arrows) and a new peak at ~ 0.8 mm/s (indicated by the red arrow). The latter feature is at the position typical for the high-energy line associated with the high-spin ferryl complexes detected in other Fe/2OG enzymes.⁸ Both sets of features are less sharp in the spectra acquired at the same temperature (4.2 K) with a 53 mT magnetic field oriented parallel to the γ -beam (Figure 3A, i and ii, vertical bars), as has also been seen in studies of other Fe/ 20G enzymes. Subtraction of the zero-field spectrum of the frozen reactant solution from the zero-field spectrum of the 0.34-s reaction sample resulted in a difference spectrum that can be simulated as a superposition of three quadrupole doublets (Figure 3A, iii, vertical bars and black solid line). Of the two downward-pointing quadrupole doublets (which arise from the species that form in the 0.34 s incubation), ohas parameters typical of a high-spin ferryl complex ($\delta = 0.26 \text{ mm}/$ s, ΔE_Q = 0.96 mm/s, Γ = 0.25 mm/s, red line) and contributes 11% of the total intensity. The 4.2 K/53 mT Mössbauer spectrum of a sample reacted with O₂ for 5 s—by which time the 318 nm-absorbing intermediate was observed to decay in SF-Abs experiments-lacks the line at 0.8 mm/s, demonstrating decay of the ferryl complex and the transient UV absorption on the same time scale (Figure 3A, iv). Together, the SF-Abs and FQ-Möss results establish the accumulation of a ferryl intermediate that can initiate L-Arg oxidation by abstracting H• from C5.

Evidence That the Ferryl Intermediate Is Specific to the RO Pathway. The modest (11%) accumulation of the ferryl intermediate arises from two factors. First, despite the retarding effect of the large D-KIE, the decay of the intermediate in the d_7 -L-Arg reaction is still not fully kinetically resolved from its formation, and this competition suppresses accumulation. Second, intermediacy of the ferryl complex in only the minor, RO pathway diminishes the quantity that can accumulate to only ~ 0.3 times (the fraction proceeding down the RO branch) the limit imposed by the former effect. This more mechanistically informative origin of ferryl suppression is established by the quantitative, global simulations of the stopped-flow data (Figure 2, Table 1, and the Supporting Information), but it can also be understood qualitatively by comparing the deflections in the ΔA_{510} and ΔA_{318} -vs-time traces from the reactions with the unlabeled and d_{7} -L-Arg substrates (Figure 2). From anoxic titrations of the enzyme with Fe(II) in the presence of the other substrates (Figure S1), we determined a molar absorption coefficient at 510 nm (ε_{510}) of 110 \pm 10 M⁻¹ cm⁻¹ for the EFE•Fe(II)•2OG•L-Arg complex. This value is somewhat greater than that reported by Martinez and Hausinger ($\sim 80 \text{ M}^{-1} \text{ cm}^{-1}$),²⁶ but problems with determination of enzyme concentration^{13,26} in that study allay our concern over the discrepancy. With the protiumbearing substrate, the ferryl complex accumulates minimally, and the ΔA_{510} -vs-time trace (Figure 2B, solid black circles) reflects (primarily) a cycle of net conversion of the absorbing reactant complex to the transparent product complex(es)

Table 1. Rate Constants for Formation and Decay of the RO Ferryl Complex Obtained by Fitting the ΔA_{318} vs Time Data Shown in Figure 2 to an A + B \rightarrow C \rightarrow D Model Using KinTek Explorer^{*a*}

	substrate	$k_{\rm formation, \ net} \ (10^3 \ { m M}^{-1} \ { m s}^{-1})$	$k_{\text{decay}} (s^{-1})$	$k_{ m H}/k_{ m D}$
wt	L-Arg	54	16	
wt	d7-l-Arg	54	1	16
D191E	L-Arg	31	42	
D191E	d_7 -L-Arg	31	1.6	26

^{*a*}Initial reactant and O₂ concentrations were reduced by a factor that reflects flux through the RO pathway (i.e., divided by 3.5 for wt and 1.05 for D191E). ε_{318} and ε_{510} values for the reactant complexes were ~40 and 90 M⁻¹ cm⁻¹, respectively, and ε_{318} and ε_{510} for the ferryl complexes were ~1450 and 190 M⁻¹ cm⁻¹, respectively.

followed by regeneration of the reactant. The downward deflection of 0.011 in this trace implies a net loss of 0.10 mM of the reactant complex (0.3 iron equiv) at ~0.85 s, the minimum of the trace. This shortfall from the maximum imposed by the $[O_2]$ of 0.24 mM is accounted for by kinetic

Scheme 2. Potential Mechanisms of Ethylene Formation by EFE

competition between decay and regeneration of the absorbing reactant complex. At the approximately equal concentrations of $EFE \bullet Fe(II) \bullet 2OG \bullet L$ -Arg and O₂, the comparable rates of the two steps, combined with the second-order nature of the first step, suppress accumulation of the product complexes (i.e., net loss of the reactant) to \sim 40% the concentration of the limiting reactant. The ferryl complexes that have been characterized in other Fe/2OG enzymes all have molar absorption coefficients (ε_{318}) of ~1500 M⁻¹ cm⁻¹, and our global kinetic model (see the Supporting Information) is consistent with an equivalent ε_{318} for the EFE ferryl complex. The upward deflection of the ΔA_{318} vs time trace from the d_7 -L-Arg reaction of 0.05 at 0.55 s thus implies accumulation of a maximum of only 0.033 mM of the ferryl intermediate (0.11 equiv), consistent with the quantity detected by freeze-quench Mössbauer spectroscopy. The time scales for decay of the ferryl complex in the d_{7} -L-Arg reaction and regeneration of the reactant complex in the reaction with unlabeled substrate differ by no more than a factor of 1.5, and so a kinetic origin of the \sim 3-fold shortfall in ferryl accumulation relative to reactant decay can be ruled out.

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The explanation is that the ferryl intermediate accumulates only in the minor, RO pathway.

The Mössbauer spectra from the freeze-quench experiment provide additional evidence for this conclusion. In addition to the downward features for the ferryl complex that develops (11% of total intensity), the difference spectrum showing the changes that occur in the first 0.34 s of the reaction (Figure 3A, iii, vertical bars) also exhibits upward features associated with loss of the Fe(II)-containing reactant complex ($\delta = 1.10 \text{ mm/s}$, $\Delta E_{\rm O} = 2.47$ mm/s, line width $\Gamma = 0.24$ mm/s, blue line) corresponding to 39% of the total intensity. The quantitative mismatch in these contributions is accounted for by the signature of a new Fe(II) species ($\delta = 1.16 \text{ mm/s}, \Delta E_Q = 3.10$ mm/s, $\Gamma = 0.54$ mm/s, 32% of total intensity, green line) that develops along with the ferryl complex. The combined analysis of the SF-Abs and FQ-Mössbauer data reveals that only approximately one-third of the reactant complex that decays in this short reaction time yields the ferryl intermediate, while approximately two-thirds yields a new Fe(II)-containing species, which is presumably the product complex of the EF pathway.

Implications of the L-Arg C5 D-KIE on Ferryl Decay and Absence of Its Effect on the EF:RO Partition Ratio. Further analysis lends even greater weight to the conclusion that the detected ferryl complex is specific to the RO pathway and not a branchpoint between the pathways. In a scenario involving partitioning of a common ferryl intermediate between HAT from C5 of L-Arg and succinate fragmentation (e.g., by proton-coupled electron transfer or PCET) to ethylene and 2 (additional) equiv of CO_2 (Scheme 2, orange vs purple or blue dotted arrows), a large D-KIE on the C5-HAT step should impact the partition ratio, making the EF pathway even more dominant. We verified that, to the contrary, the presence of deuterium does not measurably impact the EF:RO partition ratio (Figure 1). The observed partition ratio ($F_{\rm EF}/F_{\rm RO}$, where F is "fraction") would reflect the relative magnitudes of the rate constants for the two competing steps, k_{HAT} and k_{PCET} , and the observed rate constant for decay of the ferryl complex (k_{obs}) would be the sum of the rate constants for the two steps (eqs 1-4).

$$F_{\rm EF} = k_{\rm PCET} / (k_{\rm PCET} + k_{\rm HAT}) \tag{1}$$

$$F_{\rm RO} = k_{\rm HAT} / (k_{\rm PCET} + k_{\rm HAT})$$
(2)

$$F_{\rm EF}/F_{\rm RO} = k_{\rm PCET}/k_{\rm HAT} \tag{3}$$

$$k_{\rm obs} = k_{\rm HAT} + k_{\rm PCET} \tag{4}$$

In this case, with succinate oxidation being faster than Larginine oxidation by a factor of ~2.5 ($k_{\rm PCET} = 2.5k_{\rm HAT}$), the intrinsic D-KIE on the HAT step ($k_{\rm HAT}/k_{\rm DAT}$) would be only minimally "expressed" in the observed D-KIE on decay of the ferryl complex ($k_{\rm obs,H}/k_{\rm obs,D}$) as a consequence of the insensitivity of the other step (PCET) to deuterium substitution in L-Arg. Thus, even an extremely large intrinsic D-KIE on the HAT step giving a very small value of $k_{\rm DAT}$ would yield an observed effect of only ≤ 1.4 (eqs 5 and 6).

$$k_{\text{obs,H}}/k_{\text{obs,D}} = (k_{\text{HAT}} + k_{\text{PCET}})/(k_{\text{DAT}} + k_{\text{PCET}})$$
(5)

$$k_{\rm obs,H}/k_{\rm obs,D} = (k_{\rm HAT} + 2.5k_{\rm HAT})/2.5k_{\rm HAT} = 1.4$$
 (6)

By contrast, the global simulations summarized in Table 1 and the Supporting Information imply an observed D-KIE of at least 16 on decay of the intermediate with either the d_{7^-} or the d_{2^-L} -Arg substrate. This D-KIE is simply too large for there to be more than a few percent of flux (ferryl decay) through another step that competes with HAT from C5 to the intermediate complex (at least for the case of the protiumbearing L-Arg substrate). The failure of these predictions of Scheme 2 rules out this latest possible EF/RO branchpoint, with the ferryl complex as a common intermediate.

Demonstration of a Selectively Compromised EF Pathway and Enhanced Ferryl Accumulation in the **D191E Ligand Variant.** The conclusion that the detected ferryl complex is specific to the RO pathway raised the question of whether perturbations impacting the EF:RO partition ratio (e.g., amino acid substitutions or substrate modifications) might also impact the quantity of ferryl complex to accumulate. We were encouraged to address this question by previous studies, in which single substitutions near the cofactor were reported to alter the partition ratio, compromising ethylene production to a greater extent than L-Arg oxidation.¹⁷ However, none of these variant proteins have (to the best of our knowledge) been examined in the transient state for detriments to O₂ activation leading to intermediate formation, which, in past work, has generally been seen for such active-site-proximal substitutions.^{28,29} Seeking a minimal structural perturbation to alter the partition ratio without causing such unwanted, global kinetic effects, we substituted the facial-triad ligand, Asp191, with Glu. LC-MS/GC-FID assays showed that the D191E variant degrades L-Arg and 2OG to P5C (again detected after reduction to proline and Fmoc derivatization) and succinate nearly stoichiometrically, with only ~5% of the cosubstrate being fragmented to ethylene (Figure 1). As with the wt enzyme, mixing of the anoxic EFE-D191E•Fe(II)•2OG•L-Arg complex with O_2 at 5 °C resulted in development of transient absorption at 318 nm in tens of milliseconds and subsequent decay over seconds (Figure 2A, solid red circles), and use of d_7 -L-Arg in place of the protiumbearing substrate increased the amplitude and decay time by a factor of \sim 4 (open red circles). Importantly, the deflections in the ΔA_{318} -vs-time traces were, for both substrates, ~3.5 times as large as for wt EFE under the same reaction conditions, suggesting that the variant protein accumulates more of the ferryl complex.

We confirmed this implication by freeze-quench Mössbauer spectroscopy. Figure 3B shows the difference spectrum (iii, vertical bars) obtained by subtracting the 4.2 K/zero-field spectrum of the frozen D191E-EFE \bullet ⁵⁷Fe(II) \bullet 2OG \bullet d₇-L-Arg reactant complex (i, solid line) from the corresponding spectrum of a sample frozen after this complex was exposed to O2 for 0.14 s (ii, solid line). A developing quadrupole doublet from the high-spin ferryl complex appears as downward features contributing 43% of the total intensity (red line). The upward features arising from the decaying reactant complex have essentially the same relative intensity (40%, blue line). Interestingly, the Mössbauer parameters of the ferryl complexes in the variant ($\delta = 0.32 \text{ mm/s}, \Delta E_0 = 0.88$ mm/s, $\Gamma = 0.27$ mm/s) and wt (see above) enzymes are not identical, and those of their reactant complexes differ to an even greater extent (Table S8). As in the reaction of the wt enzyme, the ferryl complex in the D191E variant was seen to decay within 5 s, as indicated by the absence of a feature at \sim 0.8 mm/s in the 4.2 K/53 mT spectrum of the sample frozen at this reaction time (Figure 3B, iv).

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Figure 4. X-ray crystallographic models of the EFE active site with 2OG and L-Arg bound. (A) Comparison of the 1.83-Å-resolution wild-type EFE \bullet Fe \bullet 2OG \bullet L-Arg structure (PDB accession code 6VP4) to the published 1.43-Å-resolution EFE \bullet Mn \bullet 2OG \bullet L-Arg structure (PDB accession code 5V2Y) reveals no significant differences. (B) One of two conformations of the carboxylate side chain observed in the 1.97-Å-resolution D191E EFE \bullet Fe \bullet 2OG \bullet L-Arg structure (conformation A) closely resembles that observed in the wild-type enzyme. In (C) and (D), $2F_o-F_c$ maps contoured to 1.0 σ are shown in black for wild-type and D191E EFE, respectively. Conformation A of the E191 side chain forms a single hydrogen bond with the L-Arg substrate (E), whereas conformation B forms two hydrogen bonds (F).

The Mössbauer spectra of Figure 3 confirm the implication of the stopped-flow absorption data that 3-4 times as much of the ferryl complex accumulates in the reaction of the D191E variant as in the reaction of wt EFE. This difference does not have a kinetic origin in formation and decay rates, which are similar for the two reactions. Rather, it is explained by the almost complete elimination of flux through the EF pathway and the consequent ~3.5-fold increase in flux through the RO pathway. This explanation is further supported by the observations that, in the Mössbauer difference spectrum associated with the reaction of the D191E variant protein, the contribution from the developing ferryl complex quantitatively matches that from the decaying reactant complex (~40%), and no new features for an Fe(II)containing product complex are seen. This behavior contrasts with that of the wt EFE reaction, in which loss of the reactant

complex is associated with accumulation of both the ferryl intermediate (11%) and a new Fe(II)-containing "product" (32%). With the reaction of the D191E variant proceeding almost homogeneously through the RO pathway, the ferrylstabilizing effect of deuterium at C5 of L-Arg results in a simple net conversion of the reactant complex to the intermediatewithout significant "loss" to the nonabsorbing Fe(II) product states—over short reaction times. This simpler conversion has a noteworthy manifestation also in the ΔA_{520} vs time traces an initial rise phase as a result of the fact that the ferryl complex, which in other Fe/2OG enzymes is nearly isosbestic with its precursor, here has a greater molar absorptivity than the (unusually hypochromic) EFE•Fe(II)•2OG•L-Arg reactant complex. This fact would not have been obvious solely from the data on wild-type EFE, because the net conversion of the reactant complex to the ferryl complex is, in that case,



Figure 5. Comparison of the active sites of EFE and three other L-Arg-modifying Fe/2OG oxygenases, NapI (PDB: 6DAW), OrfP (PDB: 4M27), and VioC (PDB: 6ALM), showing a distinct L-Arg binding mode in EFE that features (i) a stacked guanidinium pair involving the substrate and Arg171 and (ii) projection of C3 and C4 over His189 to shield them from HAT.

accompanied by ~ 2.5 times as much flux directly to a transparent Fe(II)-containing product complex. These observations on the variant protein corroborate that the detected ferryl complex is specific to the RO pathway.

Structural Basis for Effect of the D191E Substitution on the EF:RO Partition Ratio. To address why the EF:RO partition ratio is so markedly perturbed by the nearly conservative ligand substitution, we solved crystal structures of the wt and D191E variant proteins in complex with Fe(II), 2OG, and L-Arg. Crystals were grown under anoxic conditions to prevent turnover, and structures were solved to resolutions of 1.83 Å (wt) and 1.97 Å (D191E). Our structure (the first) of the authentic reactant complex of wt EFE closely matches both the A conformation in the structure of the EFE•Fe(II) •N-oxalylglycine•L-Arg complex reported by Zhang et al. (PDB accession code 5LUN, rmsd ~ 0.31 Å over 329 C α atoms) and the structure of the EFE•Mn(II•2OG•L-Arg complex presented by Martinez et al. (PDB accession code 5V2Y, rmsd ~ 0.40 Å over 329 C α atoms, Figure 4A).^{16,17} Comparison of our structural models of the wt and D191E EFE•Fe(II)•2OG•L-Arg complexes reveals no global differences between the two proteins (root-mean-square deviation of ≤ 0.12 Å over 328 C α atoms). The longer tether of the ligand in the D191E variant causes the only significant deviation in the active site. For all monomers in the asymmetric unit, the new glutamate is modeled best in two different, approximately equally prevalent side chain conformations (Figure 4D), of which one is analogous to that of the Asp ligand in the wild-type protein (Figure 4B). In the other conformation, the carboxylate remains coordinated to the cofactor and hydrogen bonded with the L-Arg substrate, but the projection of both carboxylate oxygen atoms toward the substrate enables the engagement in an additional hydrogen bond with the guanidino group (Figure 4F). The ligand substitution does not, however, detectably perturb the position of either substrate. Moreover, neither of our data sets suggests the presence of the alternative L-Arg binding mode that Zhang et al. posited to enable the EF pathway.¹⁶

DISCUSSION

Regioselectivity of HAT by EFE in Comparison to Other L-Arg-Oxidizing Fe/2OG Oxygenases. The inability of deuterium-bearing isotopologues to generate alternative products of L-Arg oxidation and the identical rates of ferryl decay observed with perdeuterated L-Arg and 5,5-d₂-L-Arg demonstrate site specificity in the HAT step by EFE that contrasts with the behavior of several other ferryl-utilizing non-

heme-iron enzymes. In many of these systems, the ferryl complexes can be redirected to neighboring sites (leading to different outcomes) when challenged by substrates bearing deuterium only at the preferred $H\bullet$ -donor sites.^{19,30-34} The different behavior of EFE can be explained, at least in part, by the fact that the binding orientation of the amino acid is essentially opposite (rotated by $\sim 180^{\circ}$) to that in these other enzymes. Zhang et al. previously noted this flipped L-Arg orientation in the EFE active site relative to its orientation in the L-Arg 3-hydroxylase, VioC, from the viomycin biosynthetic pathway,^{35,36} which processes L-Arg itself and clavaminate synthase, CAS, which acts on a β -lactam derivative of L-Arg.^{16,37,38} Figure 5 illustrates the intriguing fact that the VioC/CAS-like orientation of the amino acid is, in fact, conserved in two additional, structurally characterized L-Arg modifying Fe/2OG enzymes: the 3,4-dihydroxylase, OrfP, from the streptolidine biosynthetic pathway and the 4,5desaturase, NapI, from the naphthyridinomycin pathway.^{19,39,40} Indeed, the flipped binding mode appears, at present, to be unique to EFE. The comparison with NapI is particularly germane because the 4,5-desaturase also targets C5 to initiate its reaction but can, unlike EFE, be redirected to C4 and perhaps even C3 by the presence of deuterium at the preferred HAT ${\rm site(s)}.^{19}$ In the NapI binding mode, the guanidinium functional group of the side chain is poised over His146 of the His2Asp "facial triad", and the propyl tether (C5–C3) to the α -carbon (C2) projects away. Consequently, the propyl moiety projects over the cofactor in space that is not occupied by protein ligands and is thus potentially available for the H•-abstracting ferryl oxo ligand to take up residence. Although it has been posited that, in general, the oxo ligand can be located trans to either the more C-terminal His ligand (the so-called inline position) or the more N-terminal His ligand (offline),² the vector of the C5–C3 chain potentially exposes each of these carbons for HAT to a ferryl complex in either configuration. By contrast, the EFE binding mode centers C5 above the cofactor but projects C4-C2 toward the His2Asp facial triad, making these carbons inaccessible for hydrogen abstraction. Its unique L-Arg binding mode nicely rationalizes the unusually strict regiochemistry of HAT to the ferryl complex in EFE. Might it also hold clues to the mechanism of the novel EF outcome?

Implications for the Mechanism of Ethylene Formation. The failure of the large D-KIE on HAT from L-Arg to alter the EF:RO partition ratio implies that the EFE•Fe(IV)oxo•succinate•L-Arg complex that initiates RO cannot be the branchpoint of the two pathways, but it does not rule out the

possibility that a different, fleeting (nonaccumulating) succinate-ligated ferryl complex-possibly in an "offline" configuration and originating from earlier pathway bifurcation-initiates ethylene production. Indeed, Zhang et al. proposed a ferryl-mediated Grob-like fragmentation of succinate involving migration of electron density over seven atomic centers (this type of concerted fragmentation more typically involves only five centers), in which the Fe(IV) center would function, uncharacteristically, as a nucleofuge by accepting two electrons from the coordinated succinate carboxylate (Scheme 2).¹⁶ An alternative means by which a distinct, undetected ferryl complex could initiate ethylene production would involve single-electron transfer (most likely coupled to proton transfer) to the Fe(IV) center from the succinate carboxylate (purple dotted arrow in Scheme 2) followed by radical β -scission of the C1-C2 bond, a mechanism akin to those of the electrolytic Kolbe reaction and the oxidation of 4-hydroxyphenylacetate initiated by a thivl radical in hydroxyphenylacetate decarboxylase (HPAD).⁴¹

We consider it more likely that an unusual fragmentation of an intermediate that would normally be a precursor to the succinate-coordinated ferryl complex, such as the unstable Fe(IV)-peroxyhemiketal complex or Fe(II) persuccinate intermediate, preempts its formation in the EF pathway (Scheme 2). In a mechanism envisaged by Martinez et al.,¹³ the unstable Fe(IV)-peroxyhemiketal complex would fragment between C2 and C3 rather than between C1 and C2, thus generating an oxalate-coordinated rather than a succinatecoordinated ferryl complex. Oxidative fragmentation of the oxalate to the C1- and C2-derived CO2 molecules would proceed by inner-sphere ET to the ferryl complex, decarboxylation of the oxalyl radical, and reduction of the Fe(III) center by the formate radical, similar to the mechanism of the manganese-dependent oxalate oxidase.^{42,43} In principle, the unique C2-C3 fragmentation could occur by a polar or radical mechanism (Scheme 2), but the heterolytic pathway would require the strained bicyclic complex-which is already thought to be unstable⁴⁴-to adopt a conformation with appropriate orbital alignment across the seven atomic centers involved, and the homolytic pathway would generate an Fe(V)complex, unprecedented for the Fe/2OG-oxygenase class. Interestingly, in proposing the Fe(IV)-peroxyhemiketal complex as the likely branchpoint and thereby implying that pathway bifurcation precedes ferryl formation, Martinez et al. also posited that encroachment of the Phe283 side chain toward the open coordination site of the cofactor upon L-Arg binding could promote ethylene production by impeding rearrangement of an "offline" ferryl complex to an inline configuration required for L-Arg hydroxylation. According to this hypothesis, displacement of the active-site Phe caused by the increased length/bulk of the Glu ligand in the D191E EFE variant might, in principle, cause the change in partition ratio. However, the position of the Phe283 side chain is essentially unchanged by the ligand substitution (Figure S6). Given that we do not discern an obvious structural rationale for the marked shift in the EF:RO partition ratio, it is possible that the basis is more dynamic than structural or, alternatively, arises from structural changes too subtle to detect.

In our view, the crystallographically validated Fe(II)persuccinate complex,⁴⁵ which canonically undergoes O–O bond heterolysis to generate the succinate-coordinated ferryl intermediate, is a more likely branchpoint between the RO and EF pathways than the Fe(IV)-peroxyhemiketal complex pubs.acs.org/JACS

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(Scheme 2). One possible mechanism of ethylene formation from the Fe(II) persuccinate intermediate—a seven-center Grob-like fragmentation, wherein the nucleofugal oxygen is bound to the iron center-would seem likely to require concomitant protonation of this oxygen to avoid formation of an extremely basic Fe(II)-oxo complex. An extensive hydrogen-bonding network, which includes the cosubstrate along with Arg171, Glu84, and the L-Arg substrate, could deliver the requisite proton(s). Alternatively, as has been explored extensively in studies of heme enzymes and model complexes,⁴⁶ homolysis of the iron-coordinated peroxide could compete with the canonical heterolysis, generating the Fe(III)-(hydr)oxo/succin-1-yl state that would break down as previously delineated. Given that neither of these two alternative fragmentations of the Fe(II) peroxysuccinate complex is known to occur in any other Fe/2OG enzyme, unique electrostatic features within the EFE active site would seem to be required to counteract the favored polarity of the O-O cleavage step. In this respect, it seems possible that the flipped L-Arg orientation could have greater import than merely ensuring strict regiochemistry for the ferryl-mediated HAT step in the minor RO pathway. In this orientation, the substrate guanidinium stacks with the side chain guanidinium of Arg171 (Figure 5). This counterintuitive pairing of likecharged Arg side chains in peptides and proteins has received considerable attention over the past decade and has been explained computationally.⁴⁷⁻⁵⁰ Its uniqueness to EFE among the L-Arg processing Fe/2OG oxygenases, the reported sensitivity of the EF pathway to modifications to L-Arg and substitutions of residues that interact with it, and the close proximity of the guanidinium pair to the cofactor (especially to the expected location of the coordinated peroxide of the peroxysuccinate intermediate) all suggest that this feature of the EFE•substrates complex might have a role in lowering the barrier for a noncanonical fragmentation of the intermediate that commits to ethylene production.

CONCLUSIONS

A ferryl complex accumulates in the reaction of EFE, but quantitative analysis of time-dependent absorption and Mössbauer spectra implies that it forms in only approximately one-third of the events. The 16-fold extension of its lifetime by the presence of deuterium at C5 of L-Arg-without impact to the EF:RO partition ratio-proves that the ferryl complex forms only along the RO pathway, beyond the reaction branchpoint. Thus, the D191E ligand substitution, which virtually abolishes the EF reaction, also promotes accumulation of \sim 4 times as much of the ferryl complex. The identity of the branchpoint intermediate and mechanism of the step leading to ethylene production remain to be established, but it is clear from results presented here and elsewhere that the marginal, selective stabilization of the transition state for this step by the wt enzyme is quite fragile. The basis for the differential robustness of the two pathways and whether an even greater degree of relative stabilization of the ethylene-committing transition state is possible are intriguing questions for the future.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacs.0c10923.

Materials and Methods, Figures S1–S7, Tables S1–S8, and Schemes S1 and S2 (PDF)

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Notes

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