

NIH Public Access

Author Manuscript

Biochemistry. Author manuscript; available in PMC 2012 May 31.

Published in final edited form as: *Biochemistry*. 2011 May 31; 50(21): 4733–4740. doi:10.1021/bi102042t.

The second coordination sphere of FIH controls hydroxylation[†]

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Abstract

The factor inhibiting HIF (FIH) is a proximate oxygen sensor for human cells, hydroxylating Asn⁸⁰³ within the α subunit of the hypoxia inducible factor (HIF). FIH is an α -ketoglutatrate (aKG) dependent, non-heme Fe(II) dioxygenase, in which Fe(II) is coordinated by a (His2Asp) facial triad, αKG , and H₂O. Hydrogen bonding between the facial triad, the HIF-Asn⁸⁰³ sidechain, and various second-sphere residues suggests a functional role for the second coordination sphere in tuning the chemistry of the Fe(II) center. Point mutants of FIH were prepared to test the functional role of the αKG-centered (Asn²⁰⁵, Asn²⁹⁴) or HIF-Asn⁸⁰³ centered (Arg²³⁸, Gln²³⁹) second-sphere residues. The second sphere was tested for local effects on priming Fe(II) to react with O₂, oxidative decarboxylation, and substrate positioning. Steady-sate kinetics were used to test for overall catalytic effects, autohydroxylation rates were used to test for priming and positioning, and electronic spectroscopy was used to assess the primary coordination sphere and the electrophilicity of α KG. Asn²⁰⁵ \rightarrow Ala and Asn²⁹⁴ \rightarrow Ala exhibited diminished rates of steadystate turnover, while minimally affecting autohydroxylation, consistent with impaired oxidative decarboxylation. Blue shifted MLCT transitions for (Fe+ α KG)FIH indicated that these point mutations destabilized the π^* orbitals of α KG, further supporting a slowed rate of oxidative decarboxylation. The Arg²³⁸ → Met mutant exhibited steady-state rates too low to measure and diminished product yields, suggesting impaired substrate positioning or priming; Arg²³⁸ → Met was capable of O₂-activation for the autohydroxylation reaction. The Gln²³⁹ \rightarrow Asn mutant exhibited significantly slowed steady-state kinetics and diminished product yields, suggesting impaired substrate positioning or priming. As HIF binding to $Gln^{239} \rightarrow Asn$ stimulated autohydroxylation, it is more likely that this point mutant simply mis-positions the HIF-Asn⁸⁰³ sidechain. The present work combines kinetics and spectroscopy to show that these second sphere hydrogen bonds play roles in promoting oxidative decarboxylation, priming Fe(II) to bind O_2 , and positioning HIF-Asn⁸⁰³.

A small number of Fe(II), α -ketoglutarate (α KG)-dependent dioxygenases directly control cellular oxygen sensing in humans.(1–3) These enzymes hydroxylate specific residues within the α -subunit of the hypoxia inducible factor (HIF), thereby affecting the transcriptional level of hundreds of genes.(4) The best characterized of these enzymes is 'factor inhibiting HIF-1' (FIH), which hydroxylates the β -carbon of HIF-Asn⁸⁰³;(5) HIF-Asn⁸⁰³ lies within the C-terminal transactivation domain (CTAD) of HIF α . Hydroxylating HIF-Asn⁸⁰³ turns off HIF-dependent gene expression and is a key step in down-regulating angiogenesis. Understanding the chemical basis of how FIH reacts with HIF α and O₂ is

[†]Supported by the NIH (R01-GM077413)

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Supporting Information Available: Tabulated results for thermal melting (DSC), and Co(II) binding titrations for FIH variants. This material is available free of charge via the Internet at http://pubs.acs.org.

crucial to understanding tissue O_2 homeostasis, and holds potential for treating disease states such as cancer or stroke.(6–8)

According to the consensus chemical mechanism (Scheme 1), O_2 binds to (Fe^{II}+aKG)FIH only after HIF α binds.(9, 10) In this mechanism, the active site Fe(II) of FIH is proposed to change its coordination geometry upon binding HIF α , thereby creating a binding site for O_2 . A close correlation between pO_2 and HIF hydroxylation would then result, as the rate would be proportional to pO_2 . X-ray crystallography of FIH in various substrate-bound states revealed a structural linkage between HIF-binding and O_2 activation through changes in multiple hydrogen-bonding interactions.(11–13) Controlling oxygenation chemistry through varied hydrogen bonding to atoms coordinated to the metal has been found in other enzymes and models,(14–16) suggesting that the structural changes in FIH might also have functional significance.

The chemistry within the FIH active site occurs at the Fe(II) cofactor, making the extended coordination environment of this metal the focus of our investigation. FIH provides a primary coordination sphere of His¹⁹⁹, Asp²⁰¹, and His²⁷⁹, forming a facial triad,(17) with other coordination sites occupied by H₂O or α KG (Figure 1). The secondary coordination sphere is comprised of those residues that form hydrogen bonds to the ligands. Spectroscopic and structural data from FIH and related enzymes suggest that coordination changes at Fe are crucial to turnover.(9, 18, 19) Based on evidence from related enzymes, these key coordination geometries are proposed to be a six-coordinate (6C) center for the (Fe^{II}+ α KG)FIH, and a five-coordinate (5C) center for (Fe^{II}+ α KG+CTAD)FIH.

Hydrogen bonds between surrounding residues, substrates, and iron ligands may comprise a functionally significant second-coordination sphere for FIH, both for priming as well as for oxidative decarboxylation. X-ray crystal structures of FIH imply that the single H₂O-ligand is lost from the 6C (Fe^{II}+ α KG)FIH upon binding CTAD, which would prime the Fe(II) to react with O₂.(11–13) Concomitant alterations in nearby hydrogen bonds between Arg238, Asp201, and the H₂O ligand suggest that these second-sphere interactions may be necessary for priming. We prepared and characterized the isosteric mutant Arg²³⁸ \rightarrow Met in order to determine how loss of this hydrogen bond linkage affected priming of the Fe(II) to react with O₂.

Decarboxylating α -ketoacids generally requires stabilization of negative charge within the transition state.(18–20) Notably, a Lys or Arg is positioned as a hydrogen bond donor to the C1-carboxylate of α KG in most structures of the α KG dioxygenase superfamily.(9) In the case of FIH, Asn²⁰⁵ and Asn²⁹⁴ appear to fulfill this role via hydrogen-bond donation to the C-1 carboxylate. We prepared and characterized the single-point Asn \rightarrow Ala mutants to test how loss of each hydrogen bond affected O₂-activation in normal turnover as well as autohydroxylation.

Subsequent to O₂-activation, CTAD hydroxylation likely proceeds via a hydrogen transfer/ rebound steps involving a putative [FeO]²⁺ oxidant observed in related enzymes.(21, 22) Precise substrate positioning is necessary to ensure that the putative [FeO]²⁺ oxidant attacks the proper C-H bond. The side chain of the HIF α -Asn⁸⁰³ residue is located directly above the open coordination site of the Fe(II) in FIH, and forms a hydrogen bond pair to FIH-Gln²³⁹. This interaction may be crucial for substrate positioning, such that the [FeO]²⁺ oxidant reacts selectively with the β -carbon of HIF-Asn⁸⁰³. Although Gln²³⁹ is not strictly a second-sphere residue in FIH, the Gln239 \rightarrow Asn mutant was included in the present study as a way to investigate the role of CTAD positioning on catalysis.

Testing the functional role of the second-sphere is crucial to understand the intricate relationship between the overall structure and function of FIH. This manuscript reports the

activity, metal binding, and catalytic precision of second-sphere point mutants of FIH. Our results indicate that several second-sphere residues are key to tuning the O_2 -reactivity of FIH.

Materials and methods

Protein expression and purification

FIH and its mutants were expressed from *E. coli* with an N-terminal His₆ tag and purified as previously described.(23) Following purification via Ni-NTA column hromatography, the His₆ tag was removed by thrombin digestion. Exogenous metal was removed by prolonged incubation with EDTA, and then FIH was further purified by size-exclusion chromatography to yield the FIH dimer.

Differential Scanning Calorimetry

DSC experiments were performed using a MicroCal VP-DSC microcalorimeter.(24) 50 mM Hepes buffer pH 7.50 was used and 50 μ M samples were heated in the calorimeter over a 25–75 °C range at a scan rate of 60°C/hour. A buffer scan was subtracted from each dataset to correct for base-line drift. Data analysis was performed using Origin Microcal software, (25) the results of which are provided in the supplementary material.

Metal binding

The experimental protocol was modified from Marletta et al.(26) Citrate was used as a chelator to buffer the concentration of free Co(II) ion, $[Co(H_2O)_6]2+$, for which stability constants are reported.(27) Co(II) binding to FIH was monitored by fluorescence quenching in a deoxygenated solution at room temperature. FIH was present in a fluorescence cuvette as a 200 µL solution of 1.00 mM sodium citrate, 20 µM FIH, 100 µM α -ketoglutarate, and 50 mM HEPES pH 7.50. Small volumes of CoCl₂ were added from a solution of 1.00 mM CoCl₂ and 1.00 mM sodium citrate, 50 mM HEPES pH 7.50. Following addition of Co(II), the cuvette was gently rocked by hand and the fluorescence measured over several minutes until a steady reading was obtained. Samples were excited at 280 nm and the fluorescence intensity was measured at 340 nm. Data fitting and results are provided in the supplementary material.

EPR

X-band EPR spectra were recorded on a Bruker Elexsys E-500 ESR Spectrometer equipped with a Bruker ER 4118CF-O LHe/LN₂ cryostat. EPR samples were prepared by reconstituting enzymes with CuSO₄ in a FIH:Cu²⁺ ratio of 1:0.9; the CuSO₄ solution was slowly added to the enzyme solution to prevent precipitation. Similarly, FIH(Cu+ α KG) was prepared by adding α KG slowly as the final step. Samples totaling 300 µL of 1 mM enzyme, 0.9 mM CuSO₄ and 1 mM α KG in 50 mM Hepes, pH 7.50, were frozen in quartz tubes with liquid nitrogen. The spectra were obtained by averaging 4 scans at 9.438 GHz frequency, 20 mW power, 20 G modulation amplitude, 100 GHz modulation frequency and a 327 ms time constant. The microwave power was variatied to ensure the samples were not saturated under reported conditions.

UV-Vis spectroscopy

Enzyme stocks, FeSO₄, and α KG were made anaerobic under an Argon flush. 50 mM HEPES pH 7.50 was also made anaerobic by a repeating cycle of vacuum and nitrogen flush. (Fe^{II}+ α KG)FIH spectra were obtained on anaerobic samples containing FIH (250 μ M), FeSO₄ (230 μ M) and α KG (500 μ M) in buffer. Apo FIH spectra were recorded for similar samples by omitting α KG and FeSO₄, and used as background spectra.

Activity assays

Initial rate measurements were performed in 50 mM HEPES pH 7.50 and incubated at 37 °C in 50 µL reaction volume. Reaction buffer included 2.00 mM ascorbate, 100 µM DTT, 5 u/ µL catalase, 600 µM αKG, 25 µM FeSO₄, and 0–600 µM CTAD. The reaction was initiated by adding the enzyme (0.5–5 µM), and at certain time points 5 µL aliquots were taken and quenched in 45 µL 0.1% formic acid. For each reaction 10 time points were collected and quenched, then analyzed by LC-ESI-MS. Samples were first loaded onto a C8 column for desalting, and CTAD and hydroxylated CTAD (CTAD^{OH}) were detected by ESI-MS to determine the mole fraction of peptide that had been converted to product, $\chi_{CTAD-OH}$. Product concentrations were calculated as [CTAD^{OH}] = $\chi_{(CTAD-OH)} \times [CTAD]_0$, and used to determined initial rates, which were then used for Michaelis-Menten fits.

Autohydroxylation

Autohydroxylation was measured as described previously, with minor changes.(28) FIH (100 μ M), FeSO₄ (500 μ M), and α KG (500 μ M) were anaerobically incubated in 50 mM HEPES pH 7.50 for 20 min. Autohydroxylation was initiated by adding an equal volume of buffer that had been equilibrated under air, and the reaction monitored at 583 nm. Competitive autohydroxylation assays included 50 μ M CTAD that was pre-incubated with the FIH.

Coupling

A Hamilton PRP-X300 anion exclusion column and UV detection at 210 nm was used to separate and detect succinate yield from quenched steady-state reactions of FIH. The concentration of hydroxylated CTAD, CTAD^{OH}, was determined as per the activity assays. The coupling ratio was defined as C = [succinate]/[CTAD^{OH}].

Results

Inspection of the X-ray crystal structure of FIH revealed hydrogen-bonding networks surrounding the Fe, which were centered on either α KG or CTAD. Asn²⁰⁵ and Asn²⁹⁴ residues donate hydrogen-bonds to the C-1 carboxylate of α KG, suggesting that these second-sphere residues may stabilize charge buildup during decarboxylation. Arg²³⁸ and Gln²³⁹ make hydrogen bonds with CTAD, suggesting that these residues may position the HIF-Asn⁸⁰³, or may play a role in priming the Fe for oxygenation chemistry (Figure 1). The role of these hydrogen bonds was tested by functional assays and electronic spectroscopy of point mutants.

The effect of hydrogen bonding network on turnover was investigated by steady-state kinetics (Table 1). Saturating conditions were determined for ascorbate, α KG and Fe(II) and then initial rates of WT-FIH and its mutants were acquired by varying the CTAD concentration. DTT and catalase were used in the steady-state assays to prevent ferroxidase chemistry by Fe(II) in solution. WT-FIH showed k_{cat} of 31 min⁻¹ with a $k_{cat}/K_{M(CTAD)}$ of 0.41 μ M⁻¹ min⁻¹, similar to previously reported values (Figure 2).(29–31) Each of the mutants had a modest effect on $K_{M(CTAD)}$, as both k_{cat} and k_{cat}/K_M were diminished by a similar factor for each mutant: N205A (3-fold), N294A (15-fold), and Q239N (150-fold) (Figure 2 and 3). R238M exhibited very low activity, which we could only detect through endpoint assays.

Endpoint assays showed that CTAD-centered mutants were impaired in terms of product yield, relative to WT-FIH and the α KG-centered mutants (Table 2). WT-FIH hydroxylated 94% of CTAD during prolonged incubations; the α KG-centered mutants, N205A and N294A, were nearly as thorough as WT-FIH, converting over 79% of CTAD to product. In

The ratio of succinate production to CTAD hydroxylation was measured, to check for coupling between O₂-activation and hydroxylation (Table 2). As O2-activation produces succinate, but might not lead to hydroxylated CTAD, uncoupling between these two enzymatic steps would lead to elevated coupling ratios, $C = [succinate]/[CTAD^{OH}]$. WT-FIH was tightly coupled,(43) with *C* equal to unity within experimental uncertainty (*C* = 0.98 ± 0.03). The Q205A and Q294A point mutants exhibited slightly elevated *C* values, however Q239N exhibited a coupling ratio was appreciably greater than unity (*C* = 3.1 ± 0.4).

FIH will slowly activate O₂ in the absence of CTAD, autohydroxylating Trp²⁹⁶ to form an Fe(III)-O-Trp²⁹⁶ chromophore with $\lambda_{max} = 583$ nm.(23, 28) While the rate-limiting step in autohydroxylation is not known, priming of the (Fe^{II}+ α KG)FIH is a likely requirement to permit O₂-activation. WT-FIH and the point mutants were tested for autohydroxylation rates in the absence of CTAD in order to determine how the priming of Fe(II) changed upon mutation. The autohydroxylation rates for the point mutants were only moderately altered from WT-FIH (Table 3), suggesting that the second coordination sphere had minor effects on steps directly involved in autohydroxylation. It is notable that both α KG-centered mutants exhibited 2-fold increases in the autohydroxylation rates relative to that for WT-FIH.

We felt that a competition assay, in which autohydroxylation was monitored in the presence of CTAD, would be an interesting way to further test the effect of second-sphere mutations on positioning. In the competition assay, HIF-Asn⁸⁰³ and Trp²⁹⁶ were both present as hydroxylation targets (Table 3). The rate of autohydroxylation for WT-FIH, and most of the point mutants, decreased in the presence of CTAD. The lone exception to this was the CTAD-centered mutant Q239N, which underwent autohydroxylation 60% faster in the presence of CTAD than in its absence. This increased autohydroxylation rate for Q239N strongly suggested that CTAD binding primed Fe(II) to react with O₂, but that HIF-Asn⁸⁰³ was improperly oriented to receive the oxidant in this point mutant.

The ability of WT-FIH and its mutants to stabilize negative charge on C-1 of α KG was analyzed spectroscopically by the energy of the metal to ligand charge transfer (MLCT) transition. UV-Vis absorption spectra of (Fe^{II}+ α KG)FIH form of each enzyme was measured, with the spectrum of the apo enzyme subtracted as a background. Each of the (Fe^{II}+ α KG)FIH samples exhibited an MLCT peak near 500 nm (Figure 4). This absorption band has been attributed to three overlapping transitions between populated Fe $T_{2g}(\pi)$ orbitals and empty π^* orbitals delocalized over the C1 carboxylate and C2 keto group of α KG.(18) WT-FIH displayed an MLCT maximum at 500 nm, similar to that of other α KG hydroxylases such as TauD (530 nm)(32) and CS2 (476 nm).(33) In α KG-centered mutants this MLCT was blue shifted (Table 4), with the MLCT peaks for N205A and N294A appearing at 485 and 490 nm respectively. This blue shift indicated that the α KG π^* orbitals in these point mutants were destabilized from WT-FIH, consistent with the hydrogen-bonds donated by Asn²⁰⁵ and Asn²⁹⁴ pulling electron density away from the keto-group of α KG.

Substrate centered mutations also showed blue-shifted MLCT bands (Table 4). Charge transfer complexes of Q239N and R238M appeared at 495 and 488 nm respectively. The slight shift in the MLCT energy for Q239N is consistent with the short distance between residue 239 and α KG, suggesting only minor structural perturbations. The larger shift observed for R238M is similar in magnitude to that observed for N294A, and may be

tentatively attributed to polarization effects from the loss of the positive charge at residue 238.

Cu(II) was used as a spectroscopic probe for the electronic fine structure in FIH. EPR spectra of both (Cu)FIH and (Cu+ α KG)FIH were measured for WT-FIH and each point mutant (Table 5). The spectra of the point mutants were similar to that of WT-FIH, with some heterogeneity evident in the (Cu)FIH samples that diminished significantly in the (Cu + α KG)FIH samples. As FIH only provided three protein derived ligands, heterogeneity in the (Cu)FIH samples likely arose due to the presence of two or three solvent-derived ligands with variable bond lengths to the Cu(II). The WT (Cu)FIH sample exhibited an axial spectrum with apparent g_{eff} and A_{\parallel} values ($g_{\perp} = 2.06$; $g_{\parallel} = 2.3$; $A_{\parallel} = 148.1$ G) appropriate for a "Type 2" Cu(II) site with mixed N and O-donor ligands (Figure 5).(34) Spectra of the WT (Cu+ α KG)FIH sample exhibited increased anisotropy in g_{eff} ($g_{\perp} = 2.06$, $g_{\parallel} = 2.35$) and reduced hyperfine coupling ($A_{\parallel} = 136.4$ G) as compared to (Cu)FIH (Figure 6). The small A_{\parallel} is similar to that reported for other α KG hydroxylases,(35–37) where it was attributed to an O-rich anionic ligand set and distorted planarity.(37) The spectra for the (Cu+ α KG)FIH point mutants were nearly identical to those of WT-FIH, showing that the point mutations did not alter the metal coordination geometry in the (Cu+ α KG)FIH enzyme form.

Global enzyme stability was not altered by point mutation, as shown by the irreversible melting temperatures ($T_{M(app)}$) determined by DSC (Table S1). The $T_{M(app)}$ of wild type FIH was 54.5 °C, whereas the $T_{M(app)}$ for the mutants were, in fact, slightly higher. Substrate centered mutants R238M and Q239N had slightly higher $T_{M(app)}$ than wild type, 56.2 and 56.7 °C respectively. The $T_{M(app)}$ of α KG centered mutants N205A and N294A were 58.6 and 58.9 °C respectively, again indicating that the second sphere mutations did not alter overall protein stability.

The metal binding of FIH was slightly affected by point mutation, as shown by the Co(II) titrations (Table S2). The equilibrium dissociation constant for $(\text{Co}^{2+}+\alpha \text{KG})$ FIH was 1.38(6) $\times 10^{-7}$, indicating a relatively strong affinity for Co²⁺. Each of the point mutants exhibited a similar dissociation constant, with K_{D} ranging from $1.0 - 1.9 \times 10^{-7}$. This indicated that the primary coordination sphere of the active site was not altered by mutagenesis in the secondary sphere.

Discussion

Hydrogen bonding networks have been shown to be crucial to the function of enzymes reacting with O_2 -derived species, such as lipoxygenase and SOD,(15, 16) as well as for O_2 -activating models.(14) In the case of FIH, hydrogen bonding was observed between the facial triad ligands of Fe(II), the α KG, and various second-sphere residues, suggesting a functional role in tuning the chemistry of the Fe(II) center. In addition Gln²³⁹ hydrogen bonds to the CTAD substrate, and it's role in substrate positioning has been included. The present work combines kinetics and spectroscopy to show that these second sphere hydrogen bonds play roles in promoting oxidative decarboxylation, priming Fe(II) to bind O_2 , and positioning HIF-Asn⁸⁰³.

The steady-state rate constants, k_{cat} and $k_{cat}/K_{M(CTAD)}$, were diminished for each point mutant of FIH (Table 1), indicating a reduction in the rate of one or more steps during turnover. These rate constants can be interpreted within the context of an ordered sequential mechanism, which is the consensus chemical mechanism for α KG hydroxylases.(9, 10) A minimal kinetic model (Scheme 2) for FIH in the presence of saturating α KG involves separate microscopic steps for binding S =CTAD (k_1) and O₂ (k_2), with an irreversible chemical step (k_3) followed by product release (k_4).

The steady-state rate constants are composites of the above microscopic steps. All steps after substrate binding contribute to k_{cat} , while $k_{cat}/K_{M(CTAD)}$ is a function of all steps up through the first irreversible step. The reduced values to both k_{cat} and $k_{cat}/K_{M(CTAD)}$ for each point mutant suggests that the second coordination sphere plays a significant role in determining k_3 , the common microscopic step. This first irreversible step may be oxidative decarboxylation of α KG to form the [FeO]²⁺ intermediate, as seen for TauD,(38, 39) or a another step forming an Fe(II) center, as implied for PHD.(40) In the absence of additional mechanistic probes such as kinetic isotope effects, we turned to the use of electronic spectroscopy and inactivation kinetics to test the effect of the second sphere residues.

Point mutations to second sphere residues were non-perturbing to the protein structure and primary coordination of the metal center in mutated FIH, as shown by the thermodynamic stability and electronic spectroscopy of each mutant. EPR spectra of the point mutants showed that the metal center in (Cu+ α KG)FIH retained a virtually identical ground-state electronic environment to WT-FIH (Figure 6). The MLCT transitions for (Fe^{II}+ α KG)FIH variants revealed that the α KG was a bidentate ligand to Fe(II), while also providing a view of the changing Lewis acidity of the C2-position of α KG (Figure 4). As nucleophilic attack of superoxide at C2 is thought to be a key step in O₂-activation by α KG-dependent oxygenases,(41) the Lewis acidity of the keto group is essential for oxidative decarboxylation. The blue shifts in the MLCT energies for point mutants indicated that the α KG π^* orbitals were destabilized in the N294A and N205A mutants, suggesting that the hydrogen bonds found in WT-FIH serve to pull electron density away from the C2 position of α KG.

Oxidative decarboxylation is promoted by stabilizing negative charge buildup in the transition state, as shown for enzymes(20) and model complexes.(41) Structural analysis and the MLCT shifts in FIH indicate that both Asn²⁰⁵ and Asn²⁹⁴ pull electron density away from α KG, although the Fe(II) likely plays the predominant role in activating α KG for decarboxylation. Asn²⁹⁴ plays a more significant role than Asn²⁰⁵ in stabilizing charge, as shown by the relative values for k_{cat} : WT > N205A > N294A (Table 1). This may be due to Asn²⁹⁴ being the sole residue pulling electron density away from the distal O of the C1 carboxylate of α KG, while both Asn²⁰⁵ and Fe(II) pull electron density from the proximal O of the C1 carboxylate (Figure 1).

The significant decrease in the steady-state rate constants for CTAD-centered mutants may arise from changes in Fe(II) priming or CTAD positioning. The consensus chemical mechanism suggests that mutations affecting priming should principally reduce $k_{cat}/K_{M(CTAD)}$, as steps up through O₂-activation determine this rate constant. Conversely, mutations affecting positioning should reduce k_{cat} , as steps following O₂-binding determine this rate constant. Although the structure of FIH with CTAD bound suggests that there may be hydrogen bonds with specific roles, such as those between Gln²³⁹ and HIF-Asn⁸⁰³ (positioning), and Arg²³⁸ and Asp²⁰¹ (priming), the kinetics data is less clear as each point mutation reduced k_{cat} as well as $k_{cat}/K_{M(CTAD)}$ when compared to WT FIH. This may arise from an effect on a chemical step common to both rate constants, such as O₂-activation, or from the extended hydrogen bond network linking Asp²⁰¹ to HIF-Asn⁸⁰³ thorugh Arg²³⁸ and Gln²³⁹ (Figure 1). It appears that positioning the sidechain of HIF-Asn⁸⁰³ is integral to the priming of Fe(II), and may even be a design feature to ensure tight coupling between O₂-activation.

While priming could not be separated from substrate positioning by steady-state kinetics, the two roles could be distinguished by the coupling ratio, and by analysis of hydroxylation products, as shown for TauD.(42) A functional view of priming is the ability of FIH to activate O_2 only once CTAD binds, leading to close coupling between the production of

succinate and CTAD^{OH}. In the case of WT-FIH, breakdown in priming leads solely to autohydroxylation as FIH does not release reactive oxygen species.(23, 28, 43) This was supported by our observation of a coupling value of unity ($C = 0.98 \pm 0.03$) for WT-FIH, and a near-unity coupling ratio for N205A and N294A.

Positioning HIF-Asn⁸⁰³ in the proper registry within the active site is crucial to ensure that the $[FeO]^{2+}$ oxidant attacks the correct residue and forms the CTAD^{OH} product. The Q239N mutant exhibited notable effects on the coupling ratio, indicating that Gln²³⁹ was crucial for local positioning (Table 2). Q239N was the only mutant in which the coupling ratio differed significantly from unity (3.1 ± 0.4), and the only mutant that autohydroxylated faster in the presence of CTAD. These observations indicate that, while Q239N was able to activate O₂, the HIF-Asn⁸⁰³ sidechain was improperly positioned to form normal product.

FIH serves a crucial role in sensing hypoxia within human cells. As an O₂-sensing enzyme, it must maintain tight coupling between O₂-activation and substrate hydroxylation, which it achieves by hydrogen bonding between ligands to the Fe and the second coordination sphere. These second sphere hydrogen bonds play roles in promoting O₂-activation as well as maintaining the structural registry between CTAD and Fe. Hydrogen bonds from Asn²⁰⁵ and Asn^{294} pulled electron density aweay from αKG , as shown by the LMCT transitions; removal of eityher hydrogen bond in the Asn→Ala point mutants slighly accelerated autohydroxylation, but made normal turnover much slower in the case of $Asn^{294} \rightarrow Ala$. It is likely that autohydroxylation rates reflect the intrinsic reactivity of aKG, whereas the rate of normal turnover is more dependent on precise positioning of CTAD. For this reason, the rate constants for normal turnover are greatly reduced in $Gln^{239} \rightarrow Asn$ and $Arg^{238} \rightarrow Met$, both CTAD-centered mutants. By virtue of their respective locations relative to Asp²⁰¹ and HIF-Asn⁸⁰³, it is likely that Arg^{238} is necessary for priming of the Fe(II) in response to CTAD binding, whereas Gln²³⁹ is more involved in positioning HIF-Asn⁸⁰³. The second sphere interactions play a significant role in O₂-sensing function of FIH by tuning the chemistry at the Fe(II) center.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

We thank the NIH for funding (R01-GM077413); the NSF for support of the EPR facility (CHE-0443180 NSF-CRIF); and Prof. Stephen Eyles for technical assistance.

Abbreviations

αKG	alpha-ketoglutarate
CTAD	C-terminal transactivation domain of $HIF\alpha$
DSC	differential scanning calorimetry
EPR	electron paramagnetic resonance
FIH	factor inhibiting HIF-1
HIF	hypoxia inducible factor
LC-ESI-MS	liquid chromatography - electrospray ionization mass spectrometry
MLCT	metal-to-ligand charge transfer

NTA	nitrilotriacetate	
SOD	superoxide dismutase	

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Figure 1.

Hydrogen bonding network in the active site of wild type FIH (PDB 1H2K). FIH (gray) and CTAD (cyan) shown as strands; hydrogen bond distances (Å), FIH residues, and HIF-Asn⁸⁰³ are noted.



Figure 2.

Steady state kinetics of WT-FIH and N205A. WT-FIH (0.5 μ M), N205A (1 μ M), ascorbate (2 mM), DTT (100 μ M), catalase (5 u/mL), α KG (500 μ M), FeSO₄ (25 μ M), CTAD (0–250 μ M) in 50 mM HEPES pH 7.50.



Figure 3.

Steady-state kinetics of N294A and Q239N. N294A (5 μ M), Q239N (5 μ M), ascorbate (2 mM), DTT (100 μ M), catalase (5 u/mL), α KG (500 μ M), FeSO₄ (50 μ M), CTAD (0–600 μ M) in 50 mM HEPES pH 7.50.



Figure 4.

UV-Vis spectra of (Fe^{II}+ α KG)FIH under anaerobic conditions after subtraction of (Fe^{II})FIH spectra. FIH & mutants (250 μ M), FeSO₄ (230 μ M), α KG (250 μ M) in 50 mM HEPES pH 7.50.



Figure 5.

X-band EPR spectra of (Cu^{II}) FIH variants. FIH and point mutants (1 mM), $CuSO_4$ (0.9 mM) in 50 mM HEPES pH 7.50. 9.438 GHz, 20 mW power, 20 G modulation amplitude, 100 GHz modulation frequency, 327 ms time constant, 77 K.



Figure 6.

X-band EPR spectra of (Cu^{II}+ α KG)FIH variants. FIH and mutants (1 mM), CuSO₄ (0.9 mM), α KG (1 mM) in 50 mM HEPES pH 7.50. 9.438 GHz frequency, 20 mW power, 20G modulation amplitude, 100 GHz modulation frequency, 327 ms time constant at 77 K.



Scheme 1. Consensus chemical mechanism



Scheme 2.

Kinetic parameters for WT-FIH and its mutants for aKG and the synthetic peptide CTAD (39 residues).^{*a,b*}

	$k_{\rm cat} ({\rm min}^{-1})^a$	$K_{{ m M}(lpha{ m KG})}(\mu{ m M})^b$	$K_{\mathrm{M(CTAD)}} (\mu \mathrm{M})^{a}$	$k_{\text{cat}}/K_{\text{M(CTAD)}} (\mu \text{M}^{-1} \text{min}^{-1})^a$
WT-FIH	31 ± 4	22 ± 6	77 ± 22	0.41 ± 0.1
N205A	10.2 ± 0.5	7.5 ± 2	43 ± 8	0.24 ± 0.07
N294A	2.2 ± 0.1	14 ± 2	137 ± 26	0.016 ± 0.004
Q239N	0.123 ± 0.002	24 ± 4	38 ± 2	0.0032 ± 0.0002
R238M ^C	-	-	-	-

^{*a*}Assays in which CTAD was the varied substrate were performed with FIH and mutants (0.5–5 μM), ascorbate (2 mM), DTT (100 μM), catalase (5 u/mL), α KG (500 μM), FeSO4 (25–50 μM), CTAD (0–600 μM) in 50 mM HEPES pH 7.50 at 37 °C. CTAD corresponds to a Cys→Ala point mutant of HIF-1 α ^{788–826}

 b Assays in which α KG was the varied substrate were as above, with these exceptions: CTAD (200 μ M), and α KG varied (3 – 160 μ M).

^cR238M was too slow to assay initial rates.

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Table 2

Coupling and yield of hydroxylated CTAD by FIH variants.

	C ^a	% CTAD ^{OHb}
WT-FIH	0.98 ± 0.03	94%
N205A	1.08 ± 0.04	90
N294A	1.3 ± 0.1	79
Q239N	3.1 ± 0.4	23
R238M	n.d.	5

 ${}^{a}C = [succinate]/[CTAD^{OH}]$ as determined under steady-state turnover conditions; see text for details. Not determined (n.d.).

 b Qualitative endpoints: FIH (100 μM) FeSO4 (500 μM), αKG (500 μM), CTAD (50 μM) 50 mM HEPES, pH 7.50.

Autohydroxylation rates for FIH variants.

	auto-hydroxylation initial rates (μ M/min)		
	0 CTAD	50 µM CTAD	
WT FIH	0.0442 ± 0.0007	0.0265 ± 0.0008	
N205A	0.100 ± 0.002	0.078 ± 0.001	
N294A	0.0816 ± 0.0008	0.068 ± 0.002	
Q239N	0.034 ± 0.001	0.0564 ± 0.0008	
R238M	0.0262 ± 0.0008	0.0183 ± 0.0013	

100 μM FIH, 500 μM FeSO4, 500 μM $\alpha KG,$ 0 or 50 μM CTAD, 50 mM HEPES, pH 7.50.

MLCT transitions in FIH variants.

λ_{\max} (nm)		
WT FIH	500	
N205A	485	
N294A	490	
Q239N	495	
R238M	488	

FIH (250 μM), FeSO4 (230 μM), αKG (250 μM) in 50 mM HEPES pH 7.50.

EPR spectral parameters for (Cu)FIH and (Cu+ α KG)FIH.

	EPR parameters		
	g_{\parallel}	g⊥	$A_{\parallel}({\rm G})$
(Cu)FIH	2.30	2.06	146.2
(Cu)N205A	2.31	2.06	131.0
(Cu)N294A	2.31	2.06	134.6
(Cu)Q239N	2.29	2.06	135.0
(Cu)R238M	2.31	2.06	144.8
(Cu+aKG)FIH	2.35	2.06	136.4
(Cu+aKG)N205A	2.35	2.06	136.4
(Cu+aKG)N294A	2.35	2.07	136.4
(Cu+aKG)Q239N	2.35	2.07	136.8
(Cu+aKG)R238M	2.35	2.07	136.1

Parameters observed using Spincount(44)