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Docking and Migration of Carbon Monoxide in Nitrogenase: The Case for Gated Pockets from IR Spectroscopy and Molecular Dynamics

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Abstract

Evidence for a CO docking site near the FeMo-cofactor in nitrogenase has been obtained by FT-IR monitored low temperature photolysis. We investigated the possible migration paths for CO from this docking site using molecular dynamics calculations. The simulations support the notion of a gas channel with multiple internal pockets from the active site to the protein exterior. Travel between pockets is gated by motion of protein residues. Implications for the mechanism of nitrogenase reactions with CO and N_2 are discussed.

Keywords

nitrogenase; FT-IR; molecular dynamics; myoglobin; carbon monoxide

Introduction

Nitrogenase (N₂ase) is the enzyme responsible for biological nitrogen fixation.^{1–3} For molybdenum-containing N₂ase from *Azotobacter vinelandii*, which this work will focus on, X-ray diffraction has revealed a unique [Mo-7Fe-9S-C]-homocitrate cluster at the active site of the MoFe protein of N₂ase,^{4, 5} with an interstitial carbide at the center of a prismatic 6 Fe cage.^{5–7} This cluster, known as the FeMo-cofactor (FeMo-co), is capable of reducing a wide variety of triply bonded substrates, such as N₂, C₂H₂, HCN, as well as protons.⁸ It has recently been shown that this enzyme can also produce C_xH_y hydrocarbons from CO^{9–11} and even CH₄ from CO₂.^{12, 13} The migration of substrates, such as: N₂ and CO and protons,

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Supporting Information

Contains a detailed description of the methods and figures S1 and S2: the dynamics course for N₂ and an image of the crater region respectively. This material is available free of charge via the Internet at http://pubs.acs.org.

Several different channels have been proposed for access to or from the FeMocofactor. 14-21 In 2003, Igarashi and Seefeldt pointed out a candidate substrate channel starting near surface residues α -K209 and α -W205. This mostly hydrophobic pathway (now called the 'IS channel'²⁰) ultimately passes the α -V70, α -H195, and α -R96 region and terminates at the Fe2,3,6,7 face of the FeMo-cofactor.¹⁴ At the opposite end of the FeMocofactor, a hydrophilic channel extends from a 'water pool'¹⁵ proximal to the homocitrate ligand, through the interface between α and β subunits, and finally to the surface. Durrant offered this 'interstitial channel' as an efficient path for diffusion of both dinitrogen and ammonia,¹⁶ and his proposition was later tested through site-directed mutagenesis work by Barney and coworkers.¹⁷ Dance has noted a similar channel for the egress of ammonia, starting at α -Q191.¹⁸ Smith and coworkers have recently proposed a very different dynamic channel that opens and closes on the tens of nanoseconds time scale, starting near surface residues α -R281 and α -H383 and leading to the Fe2,3,6,7 face,¹⁹ which may have a role in proton transport.^{22, 23} Additional substrate/product pathways have most recently been proposed by Morrison and coworkers, based on Caver calculations combined with analysis of binding sites for Xe and small molecules.²⁰ Apart from these small molecule channels, multiple proton relay chains ('proton bays' and 'proton wires'), leading to S2B, S3B, and S5A were noted by Durrant ¹⁶ and more recently analyzed by Dance.²¹

Pockets and channels for small molecules like CO and O_2 have proven important in the study of myoglobin (Mb) dynamics,²⁴ and they have also been found important for hydrogenases,²⁵ cytochrome oxidase,²⁶ carbon monoxide dehydrogenase/acetyl-CoA synthase,²⁷ and many other 'gas-processing' enzymes.²⁸ There is a flourishing literature on how to deduce and evaluate these conduits by computational methods.^{29–34} On the experimental side, the migration of CO following MbCO photolysis has been followed by infrared spectroscopy, including static,³⁵ picosecond time-resolved,³⁶ and temperature derivative³⁷ methods. Two photolysis product IR bands at 2131 and 2119 cm⁻¹, respectively labeled B₁ and B₂, are observed for wild-type MbCO.³⁵ These have been taken as evidence for two different CO orientations at the Mb 'docking site',³⁸ and the magnitude of the splitting has allowed calculation of the interior electric field.³⁷ Extensive time-resolved x-ray diffraction studies have allowed observation of CO migration between different pockets within Mb³⁹ and the correlation of CO pockets with those observed under high-pressure Xe.⁴⁰

Pockets and channels should certainly be relevant for understanding N₂ase. Analogous to Mb, Xe pockets have been identified by x-ray crystallography in *Klebsiella pneumoniae*⁴¹ and *Azotobacter vinelandii* (Av).⁴² Recently, a detailed comparison of Xe sites in Av and *Clostridium pasteurianum* has been made, along with binding sites for CO and other small molecules.⁴³ In this paper, we present FT-IR cryophotolysis data that supports a docking site for CO near the FeMo-cofactor. We then use molecular dynamics calculations to identify a location for the docking site in *Av* Mo N₂ase as well as a channel allowing for escape of CO and for entry of N₂.

Experimental

Photolysis of CO-Nitrogenase

The N₂ase enzyme was prepared from Av and reacted with CO using the same protocol from our previous studies.^{23, 44} The infrared spectra were collected with a Bruker Vertex 70v FT-IR under cryogenic temperatures. Photolysis was induced by a broadband Sutter Instruments xenon-arc lamp. Samples were held in custom built cells with Teflon spacers to give a pathlength of 70 microns.

Molecular Dynamics

Simulations were performed using a customized forcefield in the Gromacs molecular dynamics package.⁴⁵ The MoFe N₂ase-CO α -subunit was first energy minimized, then followed by dynamics at multiple temperatures.

More detailed information about all experimental methods is available in the supporting information.

Results

Infrared Spectroscopy

As shown in Figure 1, positive product bands appear upon low temperature photolysis of CO-inhibited wild-type N₂ase as well as the α -H195Q variant. The 37 cm⁻¹ downshifts with ¹³CO substitution confirm that these are indeed CO bands. By analogy with the MbCO literature, the absorption bands at 2135 and 2123 cm⁻¹ are consistent with 'free CO' species adopting two orientations at a docking site.³⁸ We label the N₂ase free CO bands respectively NB₁ and NB₂ by analogy to the myoglobin (Mb) CO labels.³⁵ For Mb mutants, additional bands have been observed to range from 2108 to 2152 cm⁻¹, and we cannot rule out the existence of such minor species in N₂ase.

There are differences between wild-type and α -H195Q spectra that suggest that altering the amino acid environment around the FeMo-cofactor has an effect on the photolyzed CO. In the wild type enzyme, the NB₂ band at 2123 cm⁻¹ is more intense than the 2135 cm⁻¹ NB₁ band. In α -H195Q, the intensity pattern is reversed, and the asymmetry favors the NB₁ band, suggestive of two or more unresolved sub-species. Since a small change in side-chain at the 195 position affects the photolysis spectra, these results are at least consistent with locating the 'free CO' close to that position.

Molecular Dynamics

Inspired by the experimental evidence for a CO docking site in N₂ase, we began a molecular dynamics study of CO migration from that site–analogous to those used for myoglobin and related proteins.^{15, 46} In this work we took advantage of a force field for the FeMo-cofactor that had been previously validated by comparison with results from nuclear resonance vibrational spectroscopy (NRVS), as well as structural models for CO binding to the FeMo-cofactor from DFT calculations that have been tested against NRVS and EXAFS data.²³ We chose to initially place the CO near Fe2. Our previous work has shown the photolabile

"Hi-1" CO form has a terminal CO and a formyl-like species.⁴⁷ Likewise, Fe6 has been implicated as the most reactive site on the FeMo-cofactor,⁴⁸ therefore we presume Fe6 binds the more reduced formyl-like species and Fe2 binds the terminal CO species that photolyzes to free CO. The entire protein structure was then relaxed to minimize any forces resulting from CO insertion. This yielded an initial Fe2-CO distance of 4.2 Å. The CO was then allowed to migrate through the protein at various temperatures, including 10, 80, 150, 200, 250, and 300 K. The motion of the CO and the protein was followed out to 10 ns (Figure 3).

The molecular dynamics calculations find a candidate 'docking site' with the CO about 5 Å from Fe2 (Figure 1b and Figure 3). In this location, CO is within H-bonding distance to α -His-195 N ϵ (3.1 Å). Another residue within H-bonding distance is α -Y229 (O-O distance 4.1 Å). Finally, the critical α -V70 side-chain is in position to constrain the CO motion. This bears some resemblance to the docking site in myoglobin, where photolyzed CO is within H-bonding distance to H64 and is further trapped by aliphatic side-chains of L29 and I107 after photolysis.^{38, 39, 49} As illustrated by the time courses plotted for different temperatures (Figure 2), at 10K the CO remains in the docking site, but at higher temperatures it rapidly migrates to 3 distinct regions at progressively larger distances from the FeMo-cofactor.

Upon leaving the docking site, the overall motion was stochastic and bidirectional. However, the path that was most frequently taken was strikingly similar to the IS channel.¹⁴ Our result is compared with other proposed channels and pockets in Figure 2. The channel begins near key residues α -H195 and α -V70, and it ends near surface residues α -M78, α -V179, and α -V206 and α -I259 (Figure 2). Although this might only seem a confirmation of the Igarashi channel, from the dynamics calculation we see that the migration of CO is not uniform over time (Figure 2). Rather, just as in myoglobin and other proteins,⁵⁰ the CO was relatively stable in several distinct sites-the initial 'docking site' and 3 distinct 'pockets'. Passage from the docking site and between pockets is governed by occasional gate openings by key residues along the channel through thermodynamic fluctuations. We now discuss the sites where the CO spent most of the time.

The first pocket encountered (after leaving the docking site) has the CO center-of-mass (COM) 9.5 Å from the FeMo-cofactor central carbide, with a closest approach to Fe2 of 8.5 Å (Figure 4a). Key side-chains in this pocket include α -M279, α -V71, and α -S278. Unlike the later pockets, this first pocket also includes a tyrosine side-chain, α -Y229, a good candidate for hydrogen-bonding to CO. As the closest pocket to the docking site, we label this the proximal or P-pocket.

The second ('medial' or 'M') pocket seen in the MD simulations contains α -W253, α -S254, α -I282 (Figure 4b). It is gated from the previous pocket by α -N199, α -M279, α -V71 and α -A198. The four residues sterically prevent CO from returning to the previous pocket, but they can open during thermal fluctuations. When CO is trapped within the M-pocket, the Fe2 to CO distance is about 13.0 Å. We gain confidence in this predicted location for CO because it coincides with the Xe1 pocket seen by x-ray diffraction.²⁰ As shown in Figure 4b, our typical CO position overlaps nicely with the Xe position observed by Rees and coworkers.^{20,42}

Since N_2 is the natural substrate for N_2 ase, we also performed a simulation at 300K using N_2 as the diatomic in motion. As expected, N_2 behaved similarly to CO and bidirectionally traversed through the three pockets and into the solvent space (Figure S1). In this simulation, the N_2 spent most of its time in the M pocket–just as we observed with CO.

An Energy Surface

about 25 Å.

The motion of CO within N₂ase appears to involve capture within pockets interlaced with occasional gate openings. We decided to map the potential energy surface for this motion of CO within the protein by performing a potential of mean force (PMF) calculation using GROMACS.⁵¹ This involved sampling multiple Fe2-CO distances and evolving them individually in time. The results from these calculations are shown in Figure 4d. The first minimum is a shallow well at 5 Å that corresponds to the proposed docking site. At longer distances there are three distinct potential minima corresponding to each identified pocket in the channel. Beyond these wells we see the solvent interaction crater at 20 Å and solvent space at about 25 Å. An important result from these calculations is that the location identified as the medial pocket is the lowest energy location where CO can reside.

The overall scale of the PMF is similar to what is seen in previous work for a ligand in the different channel proposed recently,¹⁹ however the PMF defined in this work has potential wells with depths greater than what is seen for the that channel. The M pocket has the largest barriers in both direction: 1.9 kcal/mol towards the D-pocket and 7.4 kcal/mol for CO moving towards the P-pocket.

Discussion

Our calculations are the first use of molecular dynamics to include an experimentally constrained force field for the FeMo-cofactor. By combining these calculations with IR-monitored cryogenic photolysis, we have found the first evidence for a CO docking site in N₂ase. We favor this docking site as the most likely position for free CO following cryogenic photolysis. The docking site is adjacent to the FeMo-cofactor and has two possible hydrogen bonding partners for CO– α -H195 and α -Y229. In myoglobin, H-bonding to H64 is considered important for the 12 cm⁻¹ splitting of product bands B₁ and B₂ at 2131 and 2119 respectively. Substitution of histidine by leucine in the H64L variant in myoglobin causes the loss of that splitting and yields a single product band at 2126 cm⁻¹.³⁸ H-bonding to α -H195 or α -Y229 may play a similar role in splitting the 'free CO' bands of photolyzed N₂ase.

Using the MD calculations, by following the path of CO or N_2 starting near this docking site, we discovered a channel for these molecules from the FeMo-cofactor to the protein exterior, in a location consistent with a previous proposal from static CAVER calculations.¹⁴ A benefit from the dynamics simulations is that they reveal a novel gating mechanism for CO or N_2 migration through this hydrophobic channel–a feature not observable from static calculations. We find that CO spends most of its time in distinct pockets, and travel between pockets is enabled by gating motions of neighboring amino acids.

Although the exact mechanism of N₂ase catalysis is far from understood, all proposed reaction pathways require multiple electron and proton transfers to the FeMo-cofactor region before any ligand binding can occur. In particular, the popular Lowe-Thornley (LT) model for the N₂ase reaction pathway⁵² posits 3 or 4 electron/proton transfers to the FeMo-cofactor and/or its associated ligands before N₂ binding. In the absence of substrate binding, the LT model proposes that the FeMo-cofactor will oxidize through H₂ evolution.⁸

It has been argued that because N_2 ase is a slow enzyme, it has no need for a hydrophobic tunnel that would allow rapid gas access to the FeMo-cofactor.²⁸ However, the pockets and gates that we observe may play a role in optimizing N_2 ase catalytic efficiency. By trapping an N_2 molecule in the M-pocket, the enzyme would have substrate available for binding within a few nanoseconds of reaching the E_3 or E_4 levels. This might help minimize 'futile' H_2 production that might occur if N_2 were not immediately available. These gated "storage" pockets would allow for N_2 availability whenever the FeMo-cofactor reaches the appropriate level of reduction. In a similar vein, it has been proposed that large sections of tunnels serve as H_2 'gas reservoirs' in hydrogenases. ⁵³, ⁵⁴

Weyman and coworkers studied the α -V75I and α -V76I variants in a similar N₂ase from *Anabaena variabilis* (which would correspond to α -V70I and α -V71I in Av).⁵⁵ Although as expected, their α -V75I variant showed reduced N₂ fixation activity and α -V76I substitution had no effect. Since our proposal for gated access between pockets is dynamic, we can accommodate these findings by simply allowing for comparable gating by isoleucine or valine residues at the Av position 71.

Some of the other channels illustrated in Figure 2 presumably play a role in the egress of the mandatory H_2 co-product as well as NH_3 or NH_4 +. In this area we are agnostic, since Lautier has noted that hydrophobic molecules sometimes travel through hydrophilic channels and *vice versa*.⁵⁶ However, it seems unlikely that evolved H_2 would exit through our proposed substrate channel, since this might limit N_2 influx to the FeMo-cofactor. It seems logical that N_2 ase would have a mechanism or 'pressure relief' for the active site region to release H_2 from the active site without interfering with the catalytic process. Overall, N_2 ase has to handle the flow of electrons, protons, N_2 , H_2 , and NH_3 , and the nature of 'traffic control' in this remarkable enzyme remains to be completely understood.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Abbreviations

Av	Azotobacter vinelandii
N2	dinitrogen
СО	carbon monoxide
N2ase	nitrogenase
FeMo-co	the iron-molybdenum cofactor of the MoFe protein
MoFe protein	the larger molybdenum-iron containing protein of nitrogenase
Fe protein	the smaller iron containing protein of nitrogenase
IR	infrared
FT-IR	fourier transform infrared spectroscopy
MB	myoglobin
MBCO	myoglobin inhibited with CO
ns	nanoseconds
ps	picoseconds
PMF	potential of mean force
LT	Lowe-Thorneley model of nitrogen fixation

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Figure 1.

Left: 'free CO' IR bands in wild-type $N_2ase^{44} vs$. H195Q N_2ase . Curves represent photolysis difference (subtracting with a no photolysis background) spectra: baseline noise before photolysis (—), halfway through photolysis (—), and endpoint of photolysis (—). Right: a model for CO in the docking site in one of the two orientations that could give rise to the observed splitting.



Figure 2.

Some of the many pathways that have been proposed 1: Igarashi & Seefeldt,¹⁴ 2: Barney,¹⁷ 3: Smith,¹⁹ and Morrison,²⁰ Also, the pockets observed in this work P: Proximal, M: Medial, and D: Distal pockets. Residues labels are given to provide a frame of reference and outline the channel path. The CO in the dynamics calculations follows the channel predicted by Igarashi (labeled 1 in the figure).



Figure 3.

Typical time-dependent traces for CO migration at different temperatures (black 300K, red 250K, purple 150K, blue 10K).



Figure 4.

(a) Illustration of the proximal pocket, (b) the medial pocket with xenon overlayed from Morrison *et al.*,²⁰ (c) the distal pocket, and (d) a profile of the interaction energy between CO and the system *vs*. the Fe2-CO distance along the proposed channel from the FeMocofactor to the solvent space.