Electron Transfer and Substrate Reduction in Nitrogenase

Karamatullah Danyal
Utah State University

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ELECTRON TRANSFER AND SUBSTRATE REDUCTON IN NITROGENASE

by

Karamatullah Danyal

A dissertation submitted in partial fulfillment
of the requirements for the degree

of

DOCTOR OF PHILOSOPHY

in

Biochemistry

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UTAH STATE UNIVERSITY
Logan, Utah

2014
ABSTRACT

Electron Transfer and Substrate Reduction in Nitrogenase

by

Karamatullah Danyal, Doctor of Philosophy

Utah State University, 2014

Major Professor: Dr. Lance C. Seefeldt
Department: Chemistry and Biochemistry

Population growth over the past ~50 years accompanied by the changes in dietary habits due to economic growth have markedly increased the demand for fixed nitrogen. Aided by biological nitrogen fixation, the Haber-Bosch process has been able to fulfill these demands. However, due to its high temperature and pressure requirements, Haber-Bosch is an expensive process. Every year, approximately 2% of the total energy expenditure by man is used to manufacture fixed nitrogen. Biological systems, on the other hand, produce ammonia at ambient temperature and pressure with much higher efficiency than the Haber-Bosch process. Research in the field of biological nitrogen fixation could prove valuable in understanding the mechanism of the enzyme responsible, nitrogenase. This could eventually allow researchers to mimic the enzyme and fix nitrogen at standard temperature and pressure, which would lead to greater availability of fixed nitrogen and a better standard of living for mankind.
As part of this research, nitrogenase of *Azotobacter vinelandii* was studied to understand the order of events in reduction of substrates and the conformational changes in the enzyme responsible for its ability to reduce said substrates at room temperature and pressure. This knowledge was used to study variant forms of nitrogenase that could be activated using controlled external reductants. This freedom from the biological reductant of nitrogenase opens the door for further research into the understanding and development of enzyme mimics that can reduce substrates at room temperature and pressure.

(191 pages)
PUBLIC ABSTRACT

Electron Transfer and Substrate Reduction in Nitrogenase

by

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Utah State University, 2014

Major Professor: Dr. Lance C. Seefeldt
Department: Chemistry and Biochemistry

The reduction of nitrogen to ammonia by the industrial Haber-Bosch process is considered one of the major scientific breakthroughs of the last century. It is considered to be responsible for approximately one third of the world’s current population. This growth over the past 50 or so years accompanied by the changes in dietary habits due to economic growth have markedly increased the demand for fixed nitrogen in the form of fertilizer. The Haber-Bosch process and biological nitrogen fixation has been able to fulfil this demand. However, this industrial process is costly due to its high temperature and pressure requirements. Every year, approximately 2% of the total global energy is used to manufacture fixed nitrogen (ammonia). Microorganisms like the diazotrophs though, can fix nitrogen at a fraction of the cost. Research in the field of biological nitrogen fixation could prove valuable in understanding the mechanism of the enzyme responsible, nitrogenase which could eventually allow researchers to mimic the enzyme
and fix nitrogen at room temperature and pressure. In the current climate of energy awareness, this should be a high priority.

Nitrogenase, the enzyme responsible for nitrogen fixation in nature, has two component proteins that associate and dissociate multiple times to reduce nitrogen. This was studied to understand the order of events in association and dissociation such as; electron transfer, ATP hydrolysis, and phosphate release. When the two components associate, the induce movements within the enzyme complex. These movements, called conformational changes, are responsible for the ability of the enzyme to reduce nitrogen at room temperature and pressure. The conformational changes, and their effect on the reduction of substrates was also studied. This knowledge was used to find and study modified forms of one half of nitrogenase that could be activated using controlled external reductants. This freedom from the association and dissociation events opens the door for further research into the understanding and development of enzyme mimics that can reduce substrates at room temperature and pressure.
ACKNOWLEDGMENTS

I would like to thank first and foremost my loved ones. Mom and Dad (Ammi and Abbu) for being an inspiration in both the desire to learn and the work ethic that goes hand in hand with this desire. Also for the constant unavering support and love that you have provided me throughout my life. Listening to me whine about an enzyme that you didn’t know much about and reminding me that all I can do is my best and even if I don’t achieve that I still have your love. My wife, who seems to be the personification of love and support and patience towards my imbalanced life that leaves less time for her than nitrogenase. Hibba, your love and support have meant everything to me and I stand indebted to you for the rest of my life. I am thankful to my sister Bano and my brothers Anser, Ahsan and Jerry who have always been there to talk with me and brighten my day each and every time. May Allah bless you and yours for all eternity. Finally, I am thankful to the Almighty Allah for granting me my childhood wish of becoming a student of nature and science and blessing me with the best support system in the world in the form of my family.

There are people at work who have been amazing over the past seven years. First and foremost, my professor Dr. Seefeldt, you have been the coolest person to work with. I have loved coming to work, enjoyed every experiment I have done and reveled in the glory of scientific research because of the environment you nurture in your lab. Your support and guidance have always made me better than I was and I hope to maintain this relationship after I leave. My supervisory committee members, I know it can be hard when you have to be on a committee for seven years, but I thank you all for all your help
and guidance and above all your encouragement year round. Thank you all for your open door policy especially Dr. Johnson whom I have constantly bothered with structure related questions over the past seven years.

I would like to thank my friends James, Brad, Simon, Boyd, Kasia, Ashwini, Ameya, Sudipta, Nimesh, AJ and the numerous others that deserve to be thanked. Some of you I worked with, some of you were buddies at or after work. In this group I would like to specially thank James Danford who has been and continues to be my best friend. I would also like to thank Simon Duval for being a great friend to me over the past two years both in the lab and outside. I am lucky to have known and been friends with you all.

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Karamatullah Danyal
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<td>7-meG</td>
<td>7-methylguanine</td>
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<td>$^{32}$P-ATP</td>
<td>Radioactively labeled ATP on the $\alpha$ phosphate</td>
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<td>ADP</td>
<td>Adenosine diphosphate</td>
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<tr>
<td>AMPPCP</td>
<td>$\beta\gamma$-methylene Adenosine Triphosphate</td>
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<td>ATP</td>
<td>Adenosine Triphosphate</td>
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<td>Apo-MoFe protein</td>
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<td>DTPA</td>
<td>Diethylene triamine pentaacetic acid</td>
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<td>EDTA</td>
<td>Ethylenediaminetetraacetic acid</td>
</tr>
<tr>
<td>EGTA</td>
<td>Ethylene glycol tetraacetic acid</td>
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<tr>
<td>$E_m$</td>
<td>Midpoint potential</td>
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<tr>
<td>$E_n$</td>
<td>MoFe protein reduced by $n$ electrons ($n=1$-$8$)</td>
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<tr>
<td>ENDOR</td>
<td>Electron nuclear double resonance</td>
</tr>
<tr>
<td>EPR</td>
<td>Electron paramagnetic resonance</td>
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<tr>
<td>ET</td>
<td>Electron transfer</td>
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<td>F cluster</td>
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<td>Fe Protein</td>
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<td>FeMo-cofactor</td>
<td>Iron-molybdenum cofactor</td>
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<td>HEPES</td>
<td>4-[2-hydroxyethyl]-1-piperazineethanesulfonic acid</td>
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<tr>
<td>$K_m$</td>
<td>Michaelis constant</td>
</tr>
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<td>$k_{Pi}$</td>
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<td>$k_{ET}$</td>
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<tr>
<td>$k_{Diss}$</td>
<td>Rate of dissociation</td>
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<td>$k_{ATP}$</td>
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<td>Iron-molybdenum cofactor (FeMo-cofactor)</td>
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<td>MDCC</td>
<td>N-[2-(1-maleimidyl)ethyl]-7-(diethylamino)coumarin-3-carboxamide</td>
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<td>$M^N$</td>
<td>FeMo-cofactor in the dithionite reduced state</td>
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<td>MOPS</td>
<td>3-(N-morpholino)propanesulfonic acid</td>
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<td>$M^R$</td>
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<td>NHE</td>
<td>Normal hydrogen electrode</td>
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<td>P cluster</td>
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<td>PBP</td>
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<td>$P^{ox}$</td>
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<td>RFQ</td>
<td>Rapid freeze quench</td>
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<td>SDS-PAGE</td>
<td>Sodium dodecyl sulfate polyacrylamide gel electrophoresis</td>
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<tr>
<td>sKIE</td>
<td>Secondary kinetic isotope effect</td>
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<tr>
<td>Abbreviation</td>
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<tr>
<td>SF</td>
<td>Stopped-flow</td>
</tr>
<tr>
<td>UV-Vis</td>
<td>Ultra violet-visible</td>
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<tr>
<td>VFQ</td>
<td>Viscogen freeze quench</td>
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Due to its seemingly endless presence in many biomolecules such as proteins and nucleic acids, the importance of nitrogen ($N_2$) is undeniable. However, only the reduced form of nitrogen can be used for growth and development of organisms. Consequently, despite its abundance in our atmosphere (~80%), it represents the limiting nutrient for growth.\(^1\) There are many oxidation states of nitrogen (+5 to -3) and the conversion of one state to the other follows a cyclic path known as the nitrogen cycle (Figure 1-1).

Nitrogen fixation, the conversion of nitrogen ($N_2$) to ammonia ($NH_3$), is of particular interest in nitrogenase research. There are three known ways in which $N_2$ is converted to $NH_3$.

---

Approximately 1\% of the $3 \times 10^{14}$ gallons of total nitrogen fixed every year comes from lightning strikes\textsuperscript{2}. The industrial process developed by Fritz Haber in 1908 and commercialized by Carl Bosch is now used to produce approximately half of the total nitrogen fixed every year.\textsuperscript{3,4} In the Haber-Bosch process hydrogen and nitrogen react at high temperature and pressure (~450°C; 200 atmospheres) over iron based catalysts to produce ammonia for fertilizers.\textsuperscript{5,6} Biological nitrogen fixation by microbes called diazotrophs is responsible for the remainder of the nitrogen fixation on our planet. A common feature in all diazotrophs is the presence of a metalloenzyme responsible for the reduction of nitrogen to ammonia, called nitrogenase (Figure 1-2).

Nitrogenase enzymes are classified according to the metal composition of their active site cofactor, which is known to contain molybdenum and iron (MoFe), vanadium and iron (VFe), or iron only (FeFe).\textsuperscript{7,8} Out of these forms of the enzyme, the molybdenum containing enzyme (MoFe), isolated from \textit{Azotobacter vinelandii} is the enzyme of choice for most researchers. This is due to the established behavior of the enzyme and genetic tractability of \textit{Azotobacter vinelandii}. All known nitrogenases include a smaller component protein called dinitrogenase reductase, also known as the iron (Fe) protein.\textsuperscript{9}

Until recently the Fe protein was the only known redox active agent with the ability to transfer electrons to the larger component (MoFe protein) in a way that the latter can reduce substrates.\textsuperscript{10-13} It contains a [4Fe-4S] cluster and transfers the electrons to the MoFe protein in a MgATP dependent manner. The MoFe protein contains the P cluster and the M cluster. The P cluster is an [8Fe-7S] metal cluster and is known to be the immediate electron acceptor from the Fe protein. It also serves as an intermediate in
the overall electron transfer from the Fe protein to the MoFe protein \( \text{(Figure 1-2).} \)\(^2\) The M cluster also known as the FeMo-cofactor is the active site of nitrogenase where substrate binding and reduction occur \([7\text{Fe-9S-Mo-C-homocitrate}]\).\(^{14,15}\) The general \( \text{N}_2 \) reduction reaction catalyzed by the Fe-MoFe complex is presented in eq 1:\(^{16}\)

\[
\text{N}_2 + 8e^- + 16\text{MgATP} + 8\text{H}^+ \rightarrow 2\text{NH}_3 + \text{H}_2 + 16\text{MgADP} + 16\text{Pi}
\] (eq 1)

\textbf{Figure 1-2: Molybdenum based nitrogenase.} Representation of MoFe and Fe protein of nitrogenase. (Left) MoFe and Fe protein complex with the two catalytic halves (PDB ID 2AFK). (Right) Clusters of nitrogenase with the direction of electron flow.
The accepted mechanism of nitrogenase enzyme proposes that the Fe protein, in its reduced and MgATP bound state (reduced by ferrodoxin), binds to one side (αβ-unit) of the MoFe protein. The immediate consequence of this binding is the electron transfer and hydrolysis of the two MgATP molecules to two MgADP molecules.\textsuperscript{17,18} The order of the two events is unknown.\textsuperscript{19–25} Two possible pathways about the electron transfer events have been proposed. In one pathway the binding of the Fe protein to the MoFe protein initiates electron transfer from the Fe protein to the P cluster.\textsuperscript{26} In the other pathway, however, the binding initiates electron transfer from the P to the M cluster within the MoFe protein followed by the reduction of the P cluster from the Fe protein.\textsuperscript{27} A distance of approximately 30Å separates the Fe protein [4Fe-4S] cluster and FeMo-cofactor.\textsuperscript{28} The P cluster acts as a mediator in the electron transfer pathway and transfers electrons to the FeMo-cofactor.

It has now been nearly twenty years since the structure of the molybdenum containing nitrogenase was solved.\textsuperscript{28,29} This was followed by structures of many variants of nitrogenase.\textsuperscript{30–32} Important information such as the positions of the clusters inside the protein as well as their arrangement and interactions with the protein, was obtained due to these efforts. These structures have, however, re-emphasized several key questions about the enzyme, some of which have been the main target of my research. In the remainder of this chapter we will take a greater look at the two protein components in nitrogenase, the role of nucleotides in electron transfer, and substrate binding and reduction on the enzyme in the presence of the Fe protein.
Iron Protein:

The homodimeric Fe protein of nitrogenase is the most common redox-active agent that can transfer electrons to the MoFe protein for substrate reduction and is the only reductant for substrate reduction to the wild-type/unaltered MoFe protein. This protein is also important in the biosynthesis and insertion of FeMo-cofactor as well as the maturation of the P cluster into an [8Fe-7S] cluster from two [4Fe-4S] clusters.

Binding of one ATP to each monomer of the Fe protein prior to complex formation and electron transfer is required for the two events to occur. The two ATP molecules bind to the Fe protein sequentially with the second ATP binding with higher affinity due to cooperative binding. Binding of MgATP to the Fe protein causes the $E_m$ for the [4Fe-4S]$^{2+/1+}$ redox couple to shift by -120 mV from -300 mV to -420 mV. The same potential change occurs when the Fe protein binds MgADP though no electron transfer to the MoFe protein takes place.

Upon binding MgATP, the Fe protein undergoes conformational changes. The major consequence of these conformational changes is the outwardly movement of the [4Fe-4S] cluster by ~5 Å. The movement of this iron sulfur cluster ligated by four cysteine residues allows for a distance shortening between it and the P cluster of the MoFe protein.

The Fe protein [4Fe-4S] cluster is paramagnetic in the as-isolated state in the presence of dithionite and is in the 1+ oxidation state with $3\text{Fe}^{2+}$ and $1\text{Fe}^{3+}$. In this oxidation state, the [4Fe-4S] cluster exhibits a mixed spin state with an $S = 1/2$ and $S = 3/2$ (Figure 1-3). This results in an EPR spectrum with features centered in the $g = 2$ region and $g = 4.3$ region. The $g = 2$ ($S=1/2$) region signal is rhombic in lineshape, with
g values of 2.04, 1.94, and 1.89 while the S = 3/2 signal has inflections around $g \approx 5$. The proportions of the two spin states can be changed by the addition of either glycerol or urea to the sample prior to freezing. Adding glycerol shifts the spin equilibrium in favor of S = 1/2, while adding urea shifts the spin equilibrium in favor of S = 3/2.

The addition of nucleotides (MgATP or MgADP) change the lineshape of the \([4\text{Fe}-4\text{S}]^{1+}\) EPR spectrum. Adding MgADP results in subtle changes to the lineshape, whereas adding MgATP results in a shift in the lineshape from rhombic to largely axial (Figure 1-3).

**Figure 1-3:** Derivative EPR spectra of Fe protein. Trace 1 shows the EPR spectrum of Fe protein (70 µM) in the as-isolated state. Spectral conditions were 9.44 GHz microwave frequency, 1.26 modulation amplitude, and 15 mW microwave power at 12 K. Trace 2 shows the Fe protein in the presence of MgATP (7.5 mM) under the same spectra conditions as trace 1.

The oxidized \([4\text{Fe}-4\text{S}]^{2+}\) cluster is diamagnetic and therefore is silent in the EPR. The 1+ oxidation state can be reduced to an all ferrous state (0 oxidation state) by
treating the Fe protein with low potential electron mediators like Ti-citrate or reduced flavodoxin.\textsuperscript{46–48} The [4Fe-4S]\textsuperscript{0} state has 4Fe\textsuperscript{2+}, and a spin S = 4 state that can be observed in the parallel mode of an EPR spectrometer. The parallel mode spectrum shows a single inflection centered at g = 16.4.\textsuperscript{46–48}

Complex formation between Fe and MoFe protein leads to ATP hydrolysis, electron transfer, phosphate release and the eventual dissociation of the Fe protein from the MoFe protein.\textsuperscript{10,49} Crystallographic evidence supports the Fe protein rolling over the surface of the MoFe protein during these events (Figure 1-4).\textsuperscript{29} During these movements the [4Fe-4S] cluster of the Fe protein moves towards and eventually away from the MoFe protein. This implies that conformational changes are taking place while Fe and MoFe protein are in complex. This is supported by the change in EPR signal of the Fe protein upon binding with 2MgATP molecules from rhombic to axial.\textsuperscript{43} Further support for the rolling motion is presented in chapters 2,3,4 and 7 where these conformational changes have been studied for the different events in the iron protein cycle.

Figure 1-4: Nitrogenase complexes. Representation of the Fe protein in complex with the MoFe protein under differing conditions: (A) nucleotide free (PDB ID 2AFH), (B) $\beta\gamma$-methylene ATP bound (PDB ID 2AFK), and (C) ADP bound form(PDB ID 2AFI).\textsuperscript{29}
MoFe protein

The \( \alpha_2\beta_2 \) heterotetrameric MoFe protein with two catalytic halves contains a P cluster and an M cluster in each half. From the X-ray crystal structure of nitrogenase complexed with the Fe protein (PDB ID 1G21) we know that the P cluster (Figure 1-5) lies between the M cluster and the Fe protein (Figure 1-2). This represents the strongest evidence for the argument that the P cluster behaves as a capacitor that receives and stores the electrons from the Fe protein and donates them to the M cluster. A general lack of direct evidence demonstrating redox changes in the P cluster during catalysis has hindered the proper assignment of the redox role of the P cluster. Therefore, a conclusive statement about the order of transfer of electrons could not be made. Some redox properties of the P cluster, such as its different oxidation states, are known. It is also known that the P cluster undergoes conformational changes upon chemical oxidation to the \( P^{ox} \) state. As determined by Mössbauer spectroscopy, under dithionite reduced conditions, the P cluster is an all ferrous iron sulfur cluster. This is known as the \( P^N \) state of the P cluster (eq 2).

\[
P^N \quad -309 \text{ mV} \quad \rightarrow \quad P^{+1} \quad \rightarrow \quad P^{+2} \quad \leftarrow \quad P^{+3} \quad +90 \text{ mV}
\]

(eq 2)

A suggested \( P^R \) state (1e- reduction beyond the \( P^N \) state) has never been observed. This is consistent with all Fe atoms being in the ferrous state in \( P^N \). The \( P^N \) state, under the influence of the midpoint potentials of different mediators, can be oxidized sequentially to the \( P^{+1}, P^{+2}, \) and the \( P^{+3} \) oxidation state. Since it is not possible to reverse the \( P^{+3} \) state to \( P^{+2} \) in vitro, it is assumed that this state is biologically irrelevant. This has led researchers to propose an electron transfer model involving only the \( P^{+1/N} \) and the \( P^{+2/+1} \) redox couple. These two redox couples of the P cluster transfer either 1e- or
2e\textsuperscript{−} from the P cluster to the M cluster. The P\textsuperscript{N} state is EPR silent, while the P\textsuperscript{+1} and the P\textsuperscript{+2} states are EPR active. As described above the P cluster can be oxidized from the resting state to three additional states called P\textsuperscript{1+}, P\textsuperscript{2+}, and P\textsuperscript{3+}. The P\textsuperscript{+1} state is a mixed spin system of S = 1/2 and S = 5/2, with a rhombic signal with g = 2.06, 1.95, and 1.82 and minor inflections at g = 6.67 and 5.3. The P\textsuperscript{2+} oxidation state does not have a signal in perpendicular mode EPR. It has a parallel mode signal with g = 11.8. This is due to a non-Kramer system integer spin S ≥ 3. The P\textsuperscript{+3} oxidation state has a mixed spin S = 7/2 and S = 1/2 state with a signal with g = 2.0, and signals at g = 10.4 and 7.2.\textsuperscript{53,56} Even though the P\textsuperscript{+1} and P\textsuperscript{+2} states of the P cluster are paramagnetic, they have proven to be difficult to observe in nitrogenase turnover samples. This has led to a general lack of knowledge about the role of the P cluster during enzyme turnover.

Figure 1-5: P cluster of the MoFe protein. The reduced form of the P cluster (right) shows the structure of the P cluster [8Fe-7S] intact with ligands such as α-62\textsuperscript{Cys}, α-88\textsuperscript{Cys}, α-154\textsuperscript{Cys}, and β-153\textsuperscript{Cys}, β-70\textsuperscript{Cys} bound to the P cluster. β-188\textsuperscript{Ser} is also shown with the reduced P cluster. The oxidized P cluster (left) shows the conformational changes that the P cluster goes through upon oxidation. The Fe 6 of the P cluster moves away from the S1 by ~1 Å and binds to the β-188\textsuperscript{Ser}. The α-88\textsuperscript{Cys} backbone nitrogen ligates with the Fe 5 of the P cluster as well. These newly formed ligations stabilize the new oxidation state of the cluster.
Two models for the role of the P cluster in nitrogenase catalysis can be hypothesized. One in which the Fe protein donates an electron to the P cluster reducing it beyond its all ferrous state to a super reduced state and the P cluster then reduces the FeMo-cofactor. A second in which upon Fe protein binding, the P cluster, in its $P^N$ state, donates an electron to the M cluster, thereby creating a deficit of electrons in itself. This deficit can then be back-filled by the Fe protein restoring the $P^N$ oxidation state of the P cluster. The deficit spending model also explains the lack of any spectroscopic evidence due to the immediate back-filling by the Fe protein.$^{27,57}$

The M cluster (also known as FeMo-cofactor) [$7\text{Fe}-9\text{S}-\text{Mo-C-Homocitrate}$] is composed of two cubane substructures bridged by three sulfides.$^{15}$ One of the two sub-clusters is a $[4\text{Fe}-4\text{S}]$ cluster while the other is a $[3\text{Fe}-4\text{S}-\text{Mo}]$ cluster. The homocitrate is bound to the molybdenum atom via two oxo ligands. The cofactor is bound to the protein by $\alpha$-$442^{\text{His}}$ and $\alpha$-$275^{\text{Cys}}$ via the molybdenum and iron(Figure 1-6)$^{14}$.

The FeMo-cofactor is the active site in nitrogenase where substrate reduction takes place.$^{10}$ It can be reduced beyond the resting state when the Fe protein binds in the presence of ATP. The $M^N$ cluster is reduced to the $M^R$ state (Figure 1-7), which is an integer spin $S \geq 1$ EPR silent, diamagnetic state.$^{58}$ As the electron transfer from the Fe protein to the MoFe protein is not synchronized, it is likely that the $M^R$ state contains more than one oxidation state of the FeMo-cofactor. The P-cluster remains silent under turnover conditions, consistent with it remaining in the $P^N$, EPR silent state.$^{59}$
Figure 1-6: FeMo-cofactor. Structure of the FeMo-cofactor with \( R \)-homocitrate and \( \alpha-70^{Val}, \alpha-96^{Arg}, \alpha-195^{His}, \alpha-275^{Cys}, \alpha-442^{His} \) amino acids. The individual atoms of the cofactor are labeled.

**Trapping inhibitors and substrates on nitrogenase during turnover.**

The EPR spectrum of the M cluster in the resting state of the MoFe protein does not show any changes upon addition of substrates or inhibitors, indicating a lack of changes to the redox state or electronic properties of the M-cluster or P cluster which has been interpreted as a lack of binding of these compounds to the \( M^N \) oxidation state. Under turnover conditions, with the M cluster reduced to the \( M^R \) state, it is possible to trap a number of different substrates and inhibitors on FeMo-cofactor, yielding a substrate-FeMo-cofactor complex that is EPR active. Given that only paramagnetic states can be observed, these trapped states must arise from FeMo-cofactor in the \( M^N \) oxidation state or even number of electron reduced states. Turnover conditions require
both Fe protein and MoFe protein, ATP, MgCl₂, an ATP-regeneration system, and the substrate that is being trapped under anaerobic conditions. The ATP regeneration system commonly used includes phosphocreatine and creatine phosphokinase. Following mixing of all of the components, the samples are frozen in liquid nitrogen.

Figure 1-7: Derivative EPR spectra of MoFe protein in resting state and during turnover. Trace 1 shows the EPR spectrum of the MoFe protein (50µM) in its as-isolated state. Spectral conditions were 9.65 GHz microwave frequency, 1.26 modulation amplitude, and 1.0 mW microwave power at 4.8 K. Trace 2 shows the EPR spectrum of MoFe protein (50µM) and Fe protein (70µM) during turnover (30 seconds of turnover prior to freeze quenching the sample) under the same spectral conditions.

Proton. Protons can be trapped bound to FeMo-cofactor in the wild-type MoFe protein, but the EPR signals are of low intensity.⁶⁴ If the valine at position α-70 is
substituted by an isoleucine, a much stronger EPR signal is observed \( (\text{Figure 1-8, trace 1}) \). The new rhombic signal arises from an \( S = 1/2 \) spin state of FeMo-cofactor, with \( g \) values of \( (2.14, 2.00, 1.96) \). This state is trapped for the MoFe protein under steady-state turnover in argon atmosphere. Use of \(^1\)H and \(^2\)H ENDOR spectroscopy reveals that the bound intermediate is associated with one or more Fe atoms and likely represents two bound hydrides. More recent analysis using \(^{95}\)Mo has revealed that the intermediate is not bound to Mo, but rather to one or more Fe atoms.\(^{65,66}\)

**Figure 1-8: EPR spectra of nitrogenase trapped during reduction of protons and propargyl alcohol.** The top trace shows the EPR spectrum of the \( \alpha-70^{1\text{le}} \) MoFe protein trapped during proton reduction. Turnover was created by mixing 100 \( \mu \)M MoFe protein with 50 \( \mu \)M Fe protein with MgATP and a MgATP regeneration system. Spectral conditions were 9.65 GHz microwave frequency, 1.26 modulation amplitude, and 2 mW microwave power at 8 K. The lower trace shows the \( \alpha-70^{\text{Ala}} \) MoFe protein trapped during turnover with propargyl alcohol. Spectral conditions were 9.64 GHz microwave frequency, 2.0 mW microwave power, and 1.26 mT modulation amplitude at 5 K.
Alkynes. The non-physiological substrate propargyl alcohol (HC≡C-CH₂OH) is reduced to allyl alcohol (H₂C=CH-CH₂OH) in the α-70Val-Ala and α-70Val-Thr variant MoFe protein. When trapped during turnover with propargyl alcohol, a new S = 1/2 spin state spectrum is observed (Figure 1-8, trace 2)(Figure 1-9, trace 1). Other non biological and alkyne substrates have also been trapped on FeMo-cofactor (e.g., CS₂ and acetylene (Figure 1-9, trace 2).  

![EPR spectra of nitrogenase trapped during reduction of acetylene and propargyl alcohol.](image)

Figure 1-9: EPR spectra of nitrogenase trapped during reduction of acetylene and propargyl alcohol. The top trace shows the EPR spectrum of the α-70Thr MoFe protein trapped during propargyl alcohol reduction. Turnover was created by mixing 50 µM MoFe protein with 50 µM Fe protein with MgATP and a MgATP regeneration system. Spectral conditions were 9.64 GHz microwave frequency, 1.26 modulation amplitude, and 2 mW microwave power at 4.8 K. The lower trace shows the α-195Gln MoFe protein trapped during turnover with acetylen. Spectral conditions were 9.65 GHz microwave frequency, 2.0 mW microwave power, and 1.26 mT modulation amplitude at 8 K.
Dinitrogen. An intermediate derived from N$_2$ can be trapped on the wild-type MoFe protein.$^{70,71}$ This is achieved by mixing Fe and MoFe protein (with an ATP regeneration system) with a full atmosphere of N$_2$ and allowing a steady-state to be achieved before rapidly freezing in a liquid N$_2$ frozen slurry of hexane. The trapped state reveals a novel S = 1/2 EPR spectra with g values 2.08, 1.99, 1.97 (Figure 1-10, trace 1).$^{70,71}$ $^{14}$N$_2$ and $^{15}$N$_2$ have been used as substrates coupled with the appropriate ENDOR spectroscopy to show that the N derived from N$_2$ is bound to FeMo-cofactor in this trapped state. The presence of a second N atom has not been confirmed, but the current model is for this intermediate to represent an N$_2$ bound end-on to one or more Fe atoms of FeMo-cofactor.$^{70}$

Diazenes. The stepwise reduction of a N$_2$ bound to FeMo-cofactor is proposed to proceed through a partially reduced intermediate at the level of reduction of diazene (HN=NH).$^{72,73}$ Diazene can be reduced by nitrogenase and can be trapped bound to FeMo-cofactor.$^{74}$ Both the wild-type and α-195$^{Gln}$ MoFe protein can be trapped under turnover conditions with diazene or methylidiazene by rapidly freezing 30 s after initiating the reaction. In the wild-type protein, the signal is similar to the N$_2$ wild-type bound state with g = 2.06, and 1.98 with a minor inflection at g = 4.25 (Figure 1-10, trace 2). On the other hand, in the α-70$^{Ala}$/α-195$^{Gln}$ variant of the MoFe protein, the diazene trapped state shows an S =1/2 spin state with g = 2.07 and 2.01. (Figure 1-10, trace 3).$^{74,75}$
Figure 1-10: EPR spectra of nitrogenase trapped during reduction of N$_2$, diazene, and hydrazine. Trace 1 shows the wild type MoFe protein trapped during turnover with 1 atm of N$_2$. Spectral conditions were 9.65 GHz microwave frequency, 1.26 modulation amplitude, and 1 mW microwave power at 4.7 K. Trace 2 shows the wild-type MoFe protein trapped during turnover with diazene, while trace 3 shows the α-70$^{\text{Ala}}$/α-195$^{\text{Gln}}$ MoFe protein trapped during turnover with diazene. Trace 4 shows the α-70$^{\text{Ala}}$/α-195$^{\text{Gln}}$ MoFe trapped during turnover with hydrazine. In all cases, turnover conditions were achieved with 75 µM MoFe protein, 50 µM Fe protein, and MgATP including a regenerating system. Spectral conditions were 9.65 GHz microwave frequency, 1.26 modulation amplitude, and 20 mW microwave power at 8 K.

*Hydrazine.* Hydrazine (H$_2$N-NH$_2$) is a substrate for nitrogenase, being reduced to two ammonia molecules.$^{76}$ It is proposed that during the reduction of N$_2$, an intermediate
is bound to FeMo-cofactor that is at the level of reduction of hydrazine.\textsuperscript{77} It is possible to trap a hydrazine derived intermediate on FeMo-cofactor. This is best achieved in a MoFe protein variant with the \( \alpha-195^{\text{His}} \) substituted by a glutamine. Trapping this variant MoFe protein during steady-state turnover with 50 mM hydrazine by rapidly freezing results in a \( S = 1/2 \) spin state with EPR signals at \( g = 2.09, 2.01, \) and 1.93. The intensity of the signals can be increased by also substituting the \( \alpha-70^{\text{Val}} \) by an alanine in the \( \alpha-195^{\text{Gln}} \) background (\textbf{Figure 1-10}, trace 4).\textsuperscript{78}

\textbf{Figure 1-11: EPR spectra of nitrogenase trapped during turnover with high and low CO.} Trace 1 shows the Hi-CO EPR signal observed when MoFe protein (50 \( \mu \)M) and Fe protein (40 \( \mu \)M) are trapped during turnover under 0.5 atm CO. Trace 2 shows the Lo-CO EPR signal when nitrogenase is trapped under turnover with 0.08 atm CO. Analysis was conducted at 9.44 GHz microwave frequency, 1 mW microwave power, and 0.84 mT (8.4 G) modulation amplitude at 12.8 K for the Hi-CO while the Lo-CO sample was obtained at 9.44GHz microwave frequency, 20mW microwave power and 0.84mT (8.4 G) modulation amplitude at 13 K.
Carbon monoxide. The inhibitor CO can also be trapped bound to FeMo-cofactor.\textsuperscript{60} Two known signals for the binding of CO to the FeMo-cofactor have been observed.\textsuperscript{79,80} One signal (\textbf{Figure 1-11}, trace 1) is known as the hi-CO signal since it is achieved at 0.5 atm or higher partial pressure of CO, while the other signal is known as the lo-CO signal (\textbf{Figure 1-11}, trace 2) since it is trapped under lower CO concentrations (0.08 atm partial pressure of CO).\textsuperscript{80,81} The hi-CO signal has been assigned to two CO molecules bound to FeMo-cofactor, while the lo-CO signal has been assigned to one CO molecule bound.\textsuperscript{60} The hi-CO signal exhibits an axial EPR signal with g = 2.17, 2.06, while the lo-CO sample exhibits a rhombic EPR signal with g = 2.09, 1.97, 1.93.\textsuperscript{80}

During catalysis of N\textsubscript{2} to ammonia, the MoFe protein of nitrogenase goes through eight separate electron reductions by the Fe protein (eight iron protein cycles).\textsuperscript{49} Different substrates bind at different electronic states (E states) and get reduced by the enzyme.\textsuperscript{82} Nitrogen reduction begins at the E\textsubscript{4} state and results in the first ammonia molecule being produced at E\textsubscript{7} and the second one being produced at E\textsubscript{8}.\textsuperscript{82}

The turnover trapped states of nitrogenase have had a tremendous impact on our understanding of the reduction of substrates on the enzyme. By trapping the turnover trapped states of diazene, hydrazine and dinitrogen and evaluating the electronic states of the cofactors it can be stated that during its reduction of nitrogen, nitrogenase undergoes an alternating mechanism in which intermediates such as diazene and hydrazine are produced (\textbf{Figure 1-12}).\textsuperscript{60,65,70,71,74,78,81}
Figure 1-12: Alternating (A) vs. distal (D) pathway of $\text{N}_2$ reduction. Here M specifies FeMo-cofactor (No individual metals are specified). Straight arrows indicate the addition of $\text{H}^+/\text{e}^-$ to the substrates.\textsuperscript{82}
**MoFe-Fe Protein Complex**

The initiation of nitrogenase catalytic cycle (docking of Fe protein to the MoFe protein) is followed by a series of events of which only a few are known and understood. From studies conducted by examining the effects of salts on the formation of Fe-MoFe complex and from the use of cross-linking agents, important interface residues were identified. These residues play a role in the docking of the Fe protein to the MoFe protein. These studies also provided evidence for the interface residues on the Fe protein being near the [4Fe-4S] cluster. The nitrogenase enzyme complex forms transiently between the MoFe protein and the Fe protein. X-ray crystallography has revealed structures for the Fe-MoFe complex nucleotide free and with MgADP, β,γ-Methylene adenosine 5’-triphosphate, and MgADP+AlF₄ bound to the Fe protein. The 127Leu-Δ variant of the Fe protein binds to the MoFe protein in the absence of any nucleotides and creates a stable and tight complex. This Fe protein is capable of binding ATP though does not hydrolyze it. This has allowed the examination of many properties of the complex. When the 127Leu-Δ protein is in complex with the MoFe protein, the Eₘ of its [4Fe-4S] cluster changes by a value of -200mV resulting in a reduction potential of -620 mV. Nitrogenase complex structures of the 127Leu-Δ Fe protein both in the presence and absence of ATP have been obtained. The P cluster’s Eₘ is shifted by -80 mV in complex. Similar Eₘ shift has been observed in the [Fe2MgADP+AlF₄]:[MoFe] complex as well.

The crystal structure of the 127Leu-Δ variant of the Fe protein in complex with MoFe protein in the presence of ATP has been obtained. When the Fe protein from this complex is compared to the crystal structure of as-isolated Fe protein many details about
the nature of the changes in the Fe protein in complex become evident. The comparison of the two structures shows that significant conformational changes are observed in the switch I and II regions (segments of Fe protein homologous to other nucleotide binding protein e.g. Ras P-21 and myosin) of the Fe protein.\textsuperscript{17} This movement in the switch I and II regions of the Fe protein results in the forward movement of the [4Fe-4S] cluster towards the P cluster. Though these movements observed in complex cannot be specified to only the binding of ATP they inform us of possible changes occurring during the electron transfer event.\textsuperscript{28}

The nucleotide free, $\beta,\gamma$-Methylene adenosine 5'-triphosphate, and MgADP+AlF$_4$ bound complex crystal structures show the extent of the conformational changes and the motion of the Fe protein over the surface of the MoFe protein (Figure 1-13).\textsuperscript{29,50} The overall area of this rolling motion of Fe protein over the surface of the MoFe protein has an area of $\sim$2000 Å$^2$. During this motion the Fe protein hydrolyses the ATP, transfers the electron and dissociates from the MoFe protein after releasing the phosphate. The [4Fe-4S] cluster of the Fe protein moves towards and eventually away from the P cluster of the MoFe protein. (Figures 1-5 and 1-13).

The [4Fe-4S] cluster of the Fe protein is situated away from the P cluster when there are no nucleotides bound to it. It moves closer and occupies almost the same position in both the AMPPCP and the ADP-AlF$_4$ bound form. Finally the cluster moves away in the opposite direction from the nucleotide free form in the ADP bound form. This movement of the cluster in conjunction with the rolling motion has been studied for the electron transfer event and the phosphate release event (Chapters 2 and 7).
Figure 1-13: Movement of the [4Fe-4S] cluster. Movement of the nucleotide free, AMPPCP(βγ-methylene ATP), ADP-AlF₄, and ADP bound [4Fe-4S] cluster relative to the P cluster of the MoFe protein shows the cluster starting from a farther distance from the P cluster in the nucleotide free form and moving closer by ~5 Å in the AMPPCP and ADP-AlF₄ bound form followed by a movement away in the ADP bound form.²⁹

Iron Protein Cycle

As noted earlier the Fe protein reduces the MoFe protein while it is transiently associated in the form of a complex (2Fe:1MoFe). This event is coupled to an ATP hydrolysis event which is followed by phosphate release and dissociation of the Fe protein from the MoFe protein. The combination of association, conformational changes, ATP hydrolysis, electron transfer, phosphate release and dissociation is known as the Fe protein cycle. The order of all the events within this cycle was unknown. It was known that upon binding 2MgATP molecules the reduced iron protein undergoes conformational
changes. This is followed by the binding of the activated Fe protein to the MoFe protein. The order of events at this point became unclear. After the binding of the Fe protein to the MoFe protein 2ATP molecules are hydrolyzed. It was also known that an electron is transferred from the Fe protein to the MoFe protein during this time as well. A number of studies have attempted to determine the order of these two (ATP hydrolysis and electron transfer) events in the Fe protein cycle. This was attempted by monitoring phosphate release with quench flow or by observing change in fluorescence using MDCC labeled phosphate binding protein, proton production using pH indicators or by monitoring heat release utilizing stopped-flow calorimetry. All of these approaches are problematic. Using quench flow to study phosphate release does not establish a difference between the released phosphate vs phosphate still bound to the Fe protein. Heat and pH changes in the overall system are difficult to attribute to a specific step in the iron protein cycle. This is especially complicated by the conformational changes in the protein. The stopped-flow fluorometric approach utilizing MDCC-PBP clearly identifies the rate of phosphate release. This rate is faster than the rate of dissociation, however, it does not help in determining when the ATP hydrolysis event takes place.

ATP hydrolysis can occur independent of electron transfer, however, electron transfer seems to require ATP. There is evidence stating that the two events might not be coupled to each other at all. Experiments conducted with 2e-reduced Fe protein show that the stoichiometry of the electrons transferred to the MoFe protein changes from 1e/2ATP molecules to 2e/2ATP molecules. This suggests that ATP binding but not hydrolysis might be needed for electron transfer. Another study using 127Leu deletion
variant of the Fe protein shows electron transfer from the Fe protein to the MoFe protein in the absence of ATP. The crystal structure of this protein shows conformational changes within the Fe protein similar to the ATP bound form of the wild type Fe protein. Evidence presented in chapter 3 using $\beta_{18}^{\text{Ser-Cys}}$ variant of the MoFe protein also supports this argument.

However, contrary evidence also exists. ITC studies showing the ATP hydrolysis event taking place prior to electron transfer suggest that not only ATP binding but also its hydrolysis has to occur prior to electron transfer.

The order of ATP hydrolysis relative to electron transfer is not the only mystery in the Fe protein cycle. The P cluster’s positioning between the [4Fe-4S] cluster of the Fe protein and the M cluster of the MoFe protein suggests a role for the P cluster in the electron transfer from the Fe protein to the M cluster of the MoFe protein. This is supported by selective mutagenesis (around the P cluster) studies that show either reduction in or complete lack of enzymatic activity and serve as evidence for the role of P cluster. Variant proteins such as $\beta_{18}^{\text{Ser-Gly}}$ and $\beta_{153}^{\text{Cys-Ser}}$ show reduction in protein activity due to these mutations around the P cluster. All of this serves as evidence supporting the role of the P cluster during electron transfer. The order of electron transfer from the [4Fe-4S] cluster of the Fe protein to the P cluster and eventually to the M cluster is uncertain.

Prior to research presented in chapter 3 two hypothesis for the order of electron transfer existed. One states that the P cluster must be reduced first by the Fe protein to an as of yet unidentified so called P super reduced state. This super reduced P cluster can then transfer the electron to the M cluster (Figure 1-14-A). Another hypothesis states that
upon Fe protein binding the P cluster transfers the electron to the M cluster resulting in a deficit of an electron on the P cluster fulfilled, eventually, by the [4Fe-4S] cluster of the Fe protein (Figure 1-14-B).

Figure 1-14: Electron transfer in nitrogenase. (A) shows the P cluster super reduction model of electron transfer with the electron from Fe protein being received by the P cluster first (step 4) while (B) shows the electron transfer event from the P cluster to the M cluster occurring first (Step 4).

The order of ATP hydrolysis relative to electron transfer was also unknown. This resulted in a lack of knowledge about whether it was the binding or hydrolysis of ATP that provided the energy for the electron transfer event. If ATP hydrolysis was occurring prior to electron transfer then the energy from the event could be responsible for the electron transfer. However, if ATP hydrolysis followed the electron transfer event then the binding of ATP rather than the hydrolysis would result in the transfer of an electron from Fe protein to the MoFe protein.20,21,24,25
In **figure 1-15**, a scheme showing the true electron transfer event and ATP hydrolysis order is shown based on work presented in chapter 5. It shows that the ATP hydrolysis event (70 s\(^{-1}\)) is slower than and thus follows electron transfer which is a faster event at \(\sim 140 \text{ s}^{-1}\).

Fe protein bound to two MgATP molecules associates with the MoFe protein. This binding/association event between the two proteins creates conformational changes within the MoFe protein between the P and the M cluster (unpublished). This conformational change is accompanied by the transient association of the \(\beta\)-188\(^{\text{Ser}}\) residue and the P cluster which leads to the lowering of the P cluster reduction potential. This allows the P cluster to reduce the FeMo-cofactor. The electron transfer event between the P and the M clusters generates an electron deficit at the P cluster. This deficit is fulfilled by the Fe protein’s [4Fe-4S]\(^{2+}\) cluster. The rate constant of this series of events is \(\sim 140 \text{ s}^{-1}\) (Chapter 3).

The oxidation of the [4Fe-4S] cluster of the Fe protein is followed by the hydrolysis of ATP at the two sites on the Fe protein (70 s\(^{-1}\)) (Chapter 5). Upon hydrolysis the protein goes through another set of conformational changes (Chapter 7) which results in the release of inorganic phosphate from the Fe protein (30 s\(^{-1}\)). The dissociation of the Fe protein from the MoFe protein (6 s\(^{-1}\)) follows the release of phosphate. Upon dissociation the Fe protein releases the two MgADP molecules and gets re-reduced. This regenerates the Fe protein for another iron protein cycle.
Figure 1-15: The Fe protein cycle. Fe protein is shown as (blue ovals) with the two oxidation states of the [4Fe-4S] cluster (+1 (orange cube) or +2 (yellow cube)) binding to MoFe protein (α subunit red and β subunit green) with the P cluster (rectangle, grey $P^N$, red activated $P^{N*}$, and yellow $P^{+1}$ states) and the M cluster (diamond, grey $M^N$ and maroon $M^R$ state). Starting from the top left with the association of the Fe protein with the MoFe protein and ending at top right with the dissociation of the oxidized Fe protein from the reduced MoFe protein the scheme shows a complete Fe protein cycle. First order rate constant ($k$, s$^{-1}$) values for 25°C are shown. Conformational changes in the Fe protein and MoFe protein are denoted by changes in shape of the ovals or rectangles.
MoFe Protein Cycle

The MoFe protein cycle is an eight state cycle in which the MoFe protein is reduced for substrate reduction via successive single electron reductions (E states) each as a result of a Fe protein cycle and each coupled to proton transfer. Different substrates bind at different E states and are subsequently reduced. Acetylene undergoes two electron reductions by binding at E_2 once sufficient electrons have been accumulated in the form of hydrides bound to the cofactor. Hydrazine binds to E_1 and is subsequently reduced in the next two electron transfer steps. Nitrogen binds at the E_4 state and is reduced in multiple steps until its release in the form of ammonia (Figure 1-16).

There are four major questions in the field of nitrogenase. Other than understanding the biosynthesis of the clusters of nitrogenase, three major questions are regarding the mechanism. These include understanding the role of ATP binding and hydrolysis in the electron transfer event (Chapter 5), the role of the P cluster in the electron transfer event (Chapter 3) and the mechanism of substrate reduction at the active site (FeMo-cofactor) (Chapter 6). The work presented in this dissertation is aimed at contributing to the understanding of these open questions about the nitrogenase mechanism.
Scheme 1-16: The MoFe protein cycle. Shows eight consecutive steps of the MoFe protein reduction with substrates entering and departing at different steps (Blue). Fe protein is shown as a homodimer with two blue ovals and a cube [4Fe-4S] cluster. MoFe protein is shown as rectangles (Red and green) prior to conformational changes and rounded edge after conformational changes. Each of the eight steps includes an iron protein cycle. Alternating mechanism of reduction is used for the depiction of nitrogen reduction.
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CHAPTER 2

CONFORMATIONAL GATING OF ELECTRON TRANSFER FROM Fe PROTEIN TO MoFe PROTEIN*

ABSTRACT

The nitrogenase Fe protein contains a [4Fe-4S] cluster and delivers one electron at a time to the catalytic MoFe protein, during this electron delivery, the Fe protein in its [4Fe-4S]$^{1+}$ reduced state (Fe$^{\text{red}}$) binds two MgATP and forms a complex with the MoFe protein, with subsequent transfer of one electron to the MoFe protein in a reaction coupled to the hydrolysis of two ATP. Crystal structures with the nitrogenase complex in different nucleotide-bound states show major conformational changes which provide a structural underpinning to suggestions that inter-component electron transfer (ET) is ‘gated’ by conformational changes of the complex and/or of its component proteins. Although electron delivery is coupled to ATP hydrolysis, their connection is puzzling, for it appears that ET precedes both ATP hydrolysis and Pi release. We here test the gating hypothesis with studies of the intracomplex oxidation of Fe$^{\text{red}}$ by MoFe protein in the presence of a variety of solutes. Conformational control of this process (gating) is revealed by the finding that it responds to changes in osmotic pressure (but not viscosity), with no fewer than 80 waters being bound during the reaction. The absence of a solvent kinetic isotope effect further implies that ATP hydrolysis does not occur during the ET step.

The Mo-dependent nitrogenase\textsuperscript{1,2} is comprised of two component proteins, denoted the Fe and MoFe proteins. The Fe protein contains a [4Fe-4S] cluster and delivers one electron at a time to the MoFe protein, which contains the multimetallic catalytic cluster [7Fe, Mo, 9S; X], designated FeMo-co, as well as the auxiliary [8Fe, 7S] P cluster, which may possibly mediate electron transfer from Fe protein to FeMo-co. During this electron delivery, the Fe protein in its [4Fe-4S]\textsuperscript{1+} reduced state (Fe\textsuperscript{red}) binds two MgATP and forms a complex with the MoFe protein, with subsequent transfer of one electron to the MoFe protein in a reaction coupled to the hydrolysis of two ATP.

**Figure 2-1:** Representation of complexes between Fe protein (top) and MoFe protein (bottom) in ATP-like (dashed) and ADP-like bound forms; (adapted from Tezcan and coworkers\textsuperscript{3}).
Crystal structures with the nitrogenase complex in different nucleotide-bound states show that major conformational changes occur upon ATP hydrolysis, Figure 2-1.\textsuperscript{3,4} This provides a structural underpinning to suggestions\textsuperscript{1,5,6} that inter-component electron transfer (ET) is “gated”\textsuperscript{7-9} by conformational changes of the complex and/or of its component proteins. Although electron delivery is coupled to ATP hydrolysis, their connection is puzzling, for it appears that ET precedes both ATP hydrolysis\textsuperscript{10} and Pi release.\textsuperscript{11} Furthermore, none of the X-ray structures of the Fe protein-MoFe protein complex, [Fe: MoFe], reveals any perturbation of either of the two clusters within the MoFe protein that might produce conformational activation of electron transfer.\textsuperscript{4}

To determine whether conformational changes control ET within the [Fe: MoFe] protein complex, we have measured oxidation of Fe\textsuperscript{red} by MoFe protein in the presence of solutes that increase the viscosity (0) and osmotic pressure of the solution. For a dynamical conformational transition in condensed media, an increase in viscosity alters the rate constant according to the formula, $k(0) \sim k^{1/0}$.\textsuperscript{12} Changes in osmotic pressure instead modulate the energetics of reactions that change the number of bound waters.\textsuperscript{13} In such a case, the observed rate constant for ET, $k_2$, varies exponentially with the molality ($m$) of added solute according to the equation, $k_2 \% \exp[-(\Delta n/55.6)m]$ where $\Delta n$ is the number of waters absorbed in the conformational transformation.

Figure 2-2 shows typical traces that monitor the oxidation of Fe\textsuperscript{red} by resting-state MoFe protein in solutions with varying concentrations of sucrose as viscogen/osmolyte. The ET reaction was initiated by the addition of MgATP to a solution of [Fe\textsuperscript{red}] and [MoFe] in the stopped flow. MgATP was added in sufficiently high concentrations to ensure that Fe\textsuperscript{red} binds two ATP and then binds to MoFe within the dead-time of the
As a result, the absorbance changes in these pre-steady-state experiments are wholly associated with Fe^{\text{red}} oxidation within the [Fe^{\text{red}}(\text{MgATP})_2: \text{MoFe}] complex.\textsuperscript{15,16}

**Figure 2-2:** Stopped-flow oxidation of Fe^{\text{red}} within the [Fe^{\text{red}}(\text{MgATP})_2: \text{MoFe}] complex. Sucrose concentrations vary from 0 to 2 \text{m}. Traces for \(m = 0\) in H\textsubscript{2}O (shown) and D\textsubscript{2}O (not shown) overlay and give the same kinetic constants within error.

The increases in absorbance of the stopped-flow traces in **Figure 2-2** caused by the oxidation of Fe^{\text{red}} are well-described by an exponential rise; the rate constant for aqueous buffer, \(k_2 = 158 \text{ s}^{-1}\), is in excellent agreement with previously reported measurements.\textsuperscript{1,17} Progressive additions of either sucrose (**Figure 2-2**), glucose and raffinose, PEGs 300, 600, or glycerol all cause progressively large decreases in the ET rate constant. This solute control of ET reveals that intra-complex oxidation of Fe^{\text{red}} by
resting-state MoFe protein is indeed ‘gated’ by a conformational transition that activates ET.

**Figure 2-3** presents plots of log $k_2$ vs the molality for each solute employed. Each plot is linear, revealing that osmotic pressure effects generate the changes in $k_2$. This is confirmed by noting that, in all cases, the changes are larger than could be generated by viscosity effects alone. For example, with sucrose as solute, for $m = 2$ the increased viscosity could at most cause $k$ to decrease by $1/0 \approx 1/5$, whereas the measured decrease is more than $1/10$; more dramatically, for glycerol as solute, with $m = 3.2$ the increased viscosity could at most cause a reduction of $\sim 1/2$, whereas $k_2$ decreases by $\sim 1/15$. In fact, it appears that viscosity plays *no* role in the solute-induced changes in ET. The plots of $k_2$ vs $m$ for PEG300 and 600 completely overlay, although the viscosities differ by roughly a factor of two at any given molality, and thus the slopes of the plots should differ likewise if the ET rate constant varied inversely with viscosity.

When $k_2$ is controlled by osmotic pressure effects, the slope of the plot of log $k$ vs $m$ is proportional to $[-\Delta n]$ for the conformational conversion (see above). The negative slopes in **Fig 2-3** thus correspond to an *uptake* of waters, $\Delta n > 0$. Multiple osmolytes were employed because the value of $\Delta n$ measured in osmotic pressure experiments in general differs among solutes because they can interact with the protein surface in different ways, displacing different numbers of bound waters and yielding different apparent values for $\Delta n$. The osmolyte whose slope is greatest interacts least and displaces the fewest waters, thereby giving the best value for $\Delta n$. In this study that osmolyte is sucrose, and its slope in **Figure 2-3** gives, $\Delta n \sim + 80$ as the best available
value lower bound for the number of waters that bind to newly exposed surfaces during
the conformational transition.

![Image](image.png)

**Figure 2-3:** Log of the rate constants for oxidation of Fe$^{\text{red}}$ within the [Fe$^{\text{red}}$(MgATP)$_2$:MoFe] complex as a function of osmolyte concentration.

Taking roughly one water to be bound per $\Delta A^0 \sim 10 \Delta^2$ of exposed surface, the
binding of $\Delta n \sim 80$ waters would correspond to a conformational transition in which $\Delta A$
$\sim 800 \Delta^2$ of surface becomes exposed. To calibrate this value, the ‘ADP’ structure of the
[Fe: MoFe] complex (Figure 2-1) exposes $\sim 2000 \Delta^2$ more protein surface than does the
“ATP” structure.\(^3\)\(^4\) Thus, the conformational change revealed by the present
measurements can be plausibly attributed to the kind of large-scale motion of the Fe
protein relative to MoFe protein shown in Figure 2-1, but likely one that is of somewhat
lesser extent and thus exposes correspondingly less surface. However, one cannot rule out
other types of structural changes.
Is this gated oxidation of Fe\textsuperscript{red} coupled to ATP hydrolysis? To test this we measured the solvent kinetic isotope effect, \( sKIE = \frac{k_2(H_2O)}{k_2(D_2O)} \), for Fe\textsuperscript{red} oxidation. ATP hydrolysis is accompanied by the release of 0.5 protons/ATP\textsuperscript{10} so if it were involved in the intra-complex oxidation of Fe\textsuperscript{red}, one would expect \( sKIE \gg 1 \). However, within error, we find that the oxidation of Fe\textsuperscript{red} has no kinetic isotope effect: \( sKIE = 1 \) (Figure 2-1). This indicates that ATP is not hydrolyzed during Fe\textsuperscript{red} oxidation.

How does conformational activation facilitate ET? One possibility is that it generates a transition state for Fe\textsuperscript{red} oxidation whose structure is optimized for direct ET from the [4Fe-4S]\textsuperscript{1+} cluster of Fe\textsuperscript{red} to FeMo-co. If the transition state occurs along an (imagined) reaction coordinate whose beginning and end points are the ATP-like and ADP-bound structures, Figure 2-1, it is unlikely that the conformational changes would enhance ET by decreasing the donor-acceptor distance: this transition increases the distances from the [4Fe-4S] cluster of the Fe protein to both the P cluster and FeMo-co\textsuperscript{3}. Among other possibilities, one may, imagine that the \( \Delta n \) waters being bound include ordered water in the interface of the active complex, and that those enhance ET\textsuperscript{19}.

Alternatively, one would expect an absence of viscosity effects if the solute effects were indeed wholly energetic, and did not operate on a dynamical process. This could occur if the rate-limiting step were preceded by a rapid pre-equilibrium between the energetically favored structure of the ATP-bound form of the complex and a higher-energy structure activated for ET, eq 1:

\[
[\text{Fe}^{\text{red}}: \text{MoFe}] \xrightarrow{K^*} [\text{Fe}^{\text{red}}: \text{MoFe}]^* \xrightarrow{k} [\text{Fe}^{\text{ox}}: \text{MoFe}^{\text{red}}]
\] (1)
The gating limit embodied in eq 1 has been termed “conformational coupling.” If the activated structure binds Δn additional waters, then $K^*$, and thus the observed ET rate constant, $k_2 = kK^*$, would be independent of viscosity but vary exponentially with the osmolyte molality ($m$), in agreement with experiment (Figure 2-3).

Experiments are under way to test the assignment of the gating motions to rearrangements such as those in Figure 2-1, the alternate mechanisms noted above, and the role of the P cluster. It seems likely to us that the nitrogenase complex employs a “compound” ET gate, with the gating motions revealed here being followed by as yet unknown conformational changes, at least some within the MoFe protein, which trigger events such as intra-MoFe ET, ATP hydrolysis, Pi release, and dissociation of the complex.

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(18) Indeed, it has been noted that sucrose behaves as an ideal solute for osmotic pressure studies, as it forms H-bonds with water with similar energetics to water-

CHAPTER 3

ELECTRON TRANSFER WITHIN NITROGENASE: EVIDENCE FOR A DEFICIT SPENDING MECHANISM

ABSTRACT

The reduction of substrates catalyzed by nitrogenase utilizes an electron transfer (ET) chain comprised of three metalloclusters distributed between the two component proteins, designated as the Fe protein and the MoFe protein. The flow of electrons through these three metalloclusters involves ET from the [4Fe-4S] cluster located within the Fe protein to an [8Fe-7S] cluster, called the P cluster, located within the MoFe protein and ET from the P cluster to the active site [7Fe-9S-X-Mo-homocitrate] cluster called FeMo-cofactor, also located within the MoFe protein. The order of these two electron transfer events, the relevant oxidation states of the P-cluster, and the role(s) of ATP, which is obligatory for ET, remain unknown. In the present work, the electron transfer process was examined by stopped-flow spectrophotometry using the wild-type MoFe protein and two variant MoFe proteins, one having the α-188Ser residue substituted by cysteine and the other having the β-153Cys residue deleted. The data support a "deficit-spending" model of electron transfer where the first event (rate constant 168 s⁻¹) is ET from the P cluster to FeMo-cofactor and the second, “backfill”, event is fast ET (rate constant >1700 s⁻¹) from the Fe protein [4Fe-4S] cluster to the oxidized P cluster. Changes in osmotic pressure reveal that the first electron transfer is conformationally gated, whereas the second is not. The data for the β-153Cys deletion MoFe protein variant

provide an argument against an alternative two-step “hopping” ET model that reverses the two ET steps, with the Fe protein first transferring an electron to the P cluster, which in turn transfers an electron to FeMo-cofactor. The roles for ATP binding and hydrolysis in controlling the ET reactions were examined using \( \beta\gamma \)-methylene-ATP as a pre-hydrolysis ATP analog and ADP + AlF\(_4^–\) as a post-hydrolysis analog (a mimic of ADP + P\(_i\)).

**INTRODUCTION**

The reduction of N\(_2\) to two ammonia molecules by nitrogenase requires eight electrons and eight protons, with two of the electrons and protons resulting in one H\(_2\) formed for each N\(_2\) reduced (eqn 1) (1).

\[
N_2 + 8 e^- + 8 H^+ + 16 MgATP \rightarrow 2 NH_3 + H_2 + 16 MgADP + 16 Pi \quad \text{(eqn 1)}
\]

For the Mo-dependent nitrogenase, electrons are delivered to substrates via an electron transfer (ET) chain comprised of three metal clusters that starts at the reduced [4Fe-4S]\(^{1+}\) cluster in the Fe protein (designated as F\(^{1+}\) cluster) (2), then goes to the [8Fe-7S] (P-cluster) in the MoFe protein, (3, 4) and ends on the [7Fe-9S-X-Mo-homocitrate] cluster or FeMo-cofactor (M cluster) in the MoFe protein (5), where substrates bind and are reduced (Figure 3-1). Transfer of a single electron to the FeMo-cofactor is initiated when the Fe protein transiently associates with the MoFe protein, with each ET event coupled to the hydrolysis of two MgATP molecules (1). This ET event has recently been shown to be conformationally gated (6–8). The order of MgATP hydrolysis and ET and the nature of their coupling remain unknown, although there is evidence to suggest that MgATP hydrolysis follows ET (1, 6, 9, 10). After ET and hydrolysis of MgATP, the oxidized Fe protein dissociates from the MoFe protein, enabling the MoFe protein to bind
another reduced, MgATP-containing, Fe protein (11). This cycle must be repeated a sufficient number of times to support the reduction of the bound substrate.

At least three different models can be considered for the ATP-coupled delivery of an electron from the Fe protein to the catalytic site FeMo-cofactor. In "direct" ET, the Fe protein would reduce FeMo-cofactor directly, skipping the P cluster. The known distances and driving forces make this option theoretically possible (12, 13), but there are several lines of evidence against such a model, including evidence for changes in oxidation state of the P cluster during turnover (3, 4), and the location of the P cluster ‘in-line’ between the F and M clusters in X-ray structures of Fe protein-MoFe protein complexes (14–16). In a "sequential" or “hopping” model for ET, the rate-limiting step is the initial transfer of an electron from the reduced [4Fe-4S]1+ cluster of the Fe protein (F1+) to the resting state of the P cluster (PN), generating a ‘super-reduced’ P-cluster (PS). This initial inter-component ET step would then be followed by rapid intra-molecular ET from PS to FeMo-cofactor (MN), resulting in the resting PN state and reduced FeMo-cofactor (MR). This model is not attractive because it would require the reduction of the all ferrous P cluster (PN) to a state (PS) that has never been observed for the P cluster (17, 18). In fact, the reduction of an FeS cluster beyond the all ferrous state has never been observed for any protein (19–22).

A third model can be denoted "deficit-spending." In this model, the first ET event involves the transfer of an electron from the PN-cluster to FeMo-cofactor (PN → MN), resulting in the formation of a P-cluster state that has been oxidized by one electron relative to the resting PN state (P1+) and reduction of the M cluster by one
Figure 3-1: Nitrogenase proteins and electron transfer between metal clusters.

**Panel A** The Fe protein (left) is shown docking to an $\alpha\beta$-unit of the MoFe protein (right). **Panel B** The two electron transfer steps: from the Fe protein [4Fe-4S] cluster (F) to the MoFe protein P cluster (P) ($k_{FP}$) and from the P cluster (P) to the FeMo-cofactor (M) ($k_{PM}$). Two ATP are hydrolyzed to two ADP and two Pi during each docking event. **Panel C** The relevant oxidation states of the three metal cluster are shown, with the resting state (in the presence of dithionite) shown in the middle, and more reduced states shown going up and more oxidized states going down. Colors are Fe in rust, S in yellow, C in gray, Mo in magenta, and O in red. The unknown atom X at the center of FeMo-cofactor is shown in blue. The MoFe protein structure was prepared from PDB file 1M1N.pdb and the Fe protein was prepared from the PDB file 1FP6.pdb using the program PyMol.
electron (M^R). The electron “deficit” in the P-cluster is then erased by "back-fill" ET from the Fe protein to the oxidized P cluster (F^{1+} → P^{1+}). This model is the most attractive of the three, being consistent with the available information, although this model has yet to be experimentally tested.

In the present work, we have conducted stopped-flow kinetic measurements of ET from the reduced Fe protein to MoFe protein of nitrogenase that test both the deficit-spending and hopping models of ET and examine the roles of ATP binding/hydrolysis and protein conformational activation in ET. Key to these studies are two MoFe protein variants with different amino acid substitutions. In one, the β-153^{Cys} ligand to the P cluster was deleted, a change which tests aspects of the hopping model for ET. In the other, the β-188^{Ser} located in close proximity to the P cluster (Figure 3-2) was substituted by cysteine, giving a MoFe protein that contains a P cluster that exists extensively in an oxidized state (P^{1+}). This feature was explained by a negative shift in E_m for the P^{1+/N} couple (estimated to be -90 mV) that leaves P^{1+} as the majority state (~ 65%) even in the presence of the reductant dithionite, with the remaining 35% in the P^N state. It was also suggested that E_m of the P cluster shifts because β-188^{Cys} forms a P-cluster ligand, thereby stabilizing the P^{1+} state. The availability of a catalytically competent MoFe protein whose P-cluster exhibits both P^N and P^{1+} states in the resting enzyme presented an opportunity to test the deficit-spending model by kinetic analysis of inter-component ET from Fe protein to the substituted MoFe protein through experiments both in aqueous buffer and in solutions of varying osmotic pressure. The roles for ATP binding and hydrolysis in controlling the electron transfer reactions were examined using 3γ-methylene-ATP as a pre-hydrolysis ATP analog and ADP + AlF_4^- as a post-hydrolysis
analog (a mimic of ADP + P_i).

**Figure 3-2: P-cluster in the oxidized and reduced states.** Shown is the P-cluster in the reduced (P_N) state (left) and in the oxidized (P^1+) state (right) with the amino acids β-153^{Cys}, β-188^{Ser}, and α-88^{Cys} shown. Structures were generated from the PDB files 1M1N and 2MIN.

**Materials and Methods**

**Materials and protein purification.** All reagents were obtained from Sigma-Aldrich Chemicals (St. Louis, MO). *Azotobacter vinelandii* strains DJ995 (wild-type MoFe protein), DJ1190 (β-188^{Cys} MoFe protein), DJ1193 (β-188^{Cys} MoFe protein expressed in a ΔnifB background, resulting in a protein lacking FeMo-cofactor, called the apo- β-188^{Cys} MoFe protein), DJ1158 (β-153^{deletion} MoFe protein), and DJ1003 (wild-type MoFe protein expressed in a ΔnifB background, resulting in a protein lacking FeMo-
cofactor, called the apo-wild-type MoFe protein) were grown and nitrogenase MoFe proteins were expressed as described previously (23). All MoFe proteins contained a seven-histidine tag on the α-subunit. This allowed purification of each protein using the previously described metal affinity chromatography protocol (23). Based on SDS-PAGE analysis using Coomassie blue staining, all proteins were found to be greater than 95% homogeneous. The rates for proton reduction were determined using previously established protocols (24). The activity for wild-type MoFe was found to be ~2100 nmols H₂/min/mg MoFe protein while the activity of ß-188Cys was found to be ~1200 nmols H₂/min/mg MoFe protein, similar to previously reported values (4). Septum sealed vials, degassed and under an argon atmosphere, were used for all manipulation of proteins. Gastight syringes were used to transfer all gasses and liquids.

**Stopped-flow spectrophotometry.** Electron transfer (ET) from Fe protein to MoFe protein was monitored by the increase in absorbance at 430 nm. This increase in absorbance results from the oxidation of the Fe protein [4Fe-4S] from its reduced (1+) state to its oxidized (2+) state. The change in absorbance was monitored as a function of time after mixing in a Hi-Tech SF61 stopped-flow UV-visible spectrophotometer equipped with a data acquisition and curve fitting system (Salisbury, Wilts, U.K.). The sample handling unit was kept inside an N₂-filled glovebox while the temperature of the reaction solutions was controlled using a circulating water bath kept outside the glovebox (25). All reactions were carried out at a temperature of 25°C, in a 100 mM HEPES buffer, pH 7.4, with 10 mM dithionite. The Fe protein and MoFe protein, when present, were contained in one drive syringe of the stopped-flow. The other drive syringe contained buffer with or without a nucleotide. The final concentrations of proteins and nucleotides
were chosen to provide two Fe proteins for each MoFe protein along with excess ATP as specified in the legends to the figures. The instrument dead-time following mixing was approximately 2 ms. Earlier work showed that the absorbance increase associated with the $F^{1+} \rightarrow F^{2+}$ oxidation can be followed by slower absorbance decreases related to re-reduction of $F^{2+}$ (26). As we are concerned only with the $F^{1+}$ oxidation step, the overall time-course of a stopped-flow trace was fit to the equation for a first-order sequential reaction corresponding to $F^{1+} \rightarrow F^{2+} \rightarrow F^{1+}$ reaction; we report only the observed rate constant, $k_{\text{obs}}$, for the $F^{1+} \rightarrow F^{2+}$ step, as the follow-up reaction involves multiple processes, including dissociation of the Fe-MoFe protein complex. The fits to this kinetic scheme were carried out in Sigmaplot (Systat Software Inc, San Jose, CA). Osmotic pressure effects on the ET rate were examined as described (6).

**Results**

**ET from Fe protein to MoFe protein.** Pre-steady state ET from the Fe protein to the resting-state MoFe protein is monitored through measurement of the increase in the absorbance at 430 nm that accompanies the oxidation of $F^{1+}$ to $F^{2+}$ (27, 28). Stopped-flow mixing of reduced Fe protein with wild-type resting-state MoFe protein in the presence of saturating MgATP yields stoichiometric oxidation of $F^{1+}$ (Figure 3-3) in a process that is exponential, with a rate constant, $k_{\text{obs}} = 168 \text{ s}^{-1}$, that is consistent with earlier studies (26).
Figure 3-3: Electron transfer from the Fe protein to MoFe proteins monitored by
stopped-flow spectroscopy. Shown is the absorbance at 430 nm plotted against the time
after mixing Fe protein (75 μM) and MoFe proteins (20 μM) in one syringe against
different solutions (without or with nucleotides at 18 mM). Shown is wild-type MoFe
protein and Fe protein mixed with MgATP (WT+ATP, green); β-188Cys MoFe protein
and Fe protein mixed with MgATP (β-188Cys + ATP, blue); β-188Cys MoFe protein and
Fe protein mixed with MgADP (β-188Cys + ADP, orange); and β-188Cys MoFe protein
and Fe protein mixed with buffer without nucleotides (β-188Cys - nucleotides, light
green). The top two sets of data were fit to a single exponential (solid lines) for the wild-
type MoFe protein data and to a single exponential with a decline for the β-188Cys MoFe
protein data, with determined rate constants of 168 s⁻¹ (WT+ATP) and 187 s⁻¹ (β-188Cys +
ATP). The mixing dead-time is ~ 2 ms. Other parameters and equations for fits are
described in Materials and Methods.
When the β-188Cys MoFe protein is mixed with reduced Fe protein plus MgATP, the absorbance change indicates that ET associated with oxidation of F^{1+} also is stoichiometric (Figure 3-3). However, unlike reaction with the wild-type MoFe protein, the Fe protein oxidation by β-188Cys MoFe protein shows an ET "burst", with ~65% of the F^{1+} oxidation occurring during the instrument dead-time (2 ms). The remaining 35% oxidation occurs with a resolved time-course that can be fit to an exponential with a rate constant $k_{obs} = 187$ s^{-1}, essentially the same as observed for the wild-type MoFe protein (25). The β-188Cys MoFe protein in dithionite solution was earlier shown to exist as an equilibrium mixture of 3 states of the P cluster (P^{1+}, P^{2+}, and P^N), with the total contribution from the two oxidized forms, being ~65% of the enzyme present with ~35% existing as the resting oxidation state, P^N (4). The kinetically resolved phase of the stopped-flow trace, with rate constant equivalent to that for wild-type MoFe protein, is thus assigned to oxidation of F^{1+} by β-188Cys MoFe protein in the P^N/M^N state, with the "burst" ET phase being assigned to rapid oxidation of F^{1+} by P^{1+} of the β-188Cys MoFe protein in the P^{1+}/M^N state. This assignment is confirmed below.

In a separate study, when the β-153Cys of the MoFe protein was deleted, this MoFe protein variant showed stoichiometric oxidation of F^{1+} with a rate constant essentially unchanged from that observed to the wild-type MoFe protein.

**The roles of nucleotides in ET.** We next consider the contributions of nucleotides to the ET events to the β-188Cys MoFe protein. Both the binding and hydrolysis of ATP within the Fe protein is intimately linked to ET from the Fe protein to the wild-type MoFe protein during substrate reduction. MgADP and MgATP analogs do not support the ET reaction (29). As shown in Figure 3-3, MgADP also does not support
oxidation of the Fe protein by the β-188Cys MoFe protein: neither the burst nor the resolved kinetic phases are observed. Thus, it can be concluded that MgATP binding, at a minimum, is required for both the burst-phase reduction of P^{1+} of the β-188Cys MoFe protein in the P^{1+}/M^N state as well as the resolved ET to the P^N/M^N state.

To further differentiate the respective roles of ATP binding and hydrolysis in ET from the Fe protein to the β-188Cys MoFe protein, ET reactions were monitored in the presence of ATP analogs. The complex between Fe protein and MoFe protein that contains the non-hydrolyzable analog βγ-methylene-ATP was earlier shown to induce conformational changes in the Fe protein similar to those induced by MgATP, yet the analog is not hydrolyzed by nitrogenase (30–33). As previously reported (30), βγ-methylene-ATP does not support ET from the Fe protein to the wild-type MoFe (Figure 3-4). Likewise, this analog does not support the resolved phase of Fe protein oxidation by the β-188Cys MoFe protein. However, this ATP analog does support the burst ET from the Fe protein to P^{1+}. Thus, the conformational changes in the Fe protein and/or in the Fe protein-MoFe protein complex induced by binding of βγ-methylene-ATP are sufficient to enable the rapid $F^{1+} \rightarrow P^{1+}$ ET, but not the ET from $F^{1+}$ to the MoFe protein in the $P^N/M^N$ state.

It has been proposed for several ATP-hydrolyzing enzymes (34, 35), including nitrogenase, that the state following ATP hydrolysis, but before Pi release, [MoFe-Fe(2MgADP + 2 Pi)], is simulated when MgADP + AlF_4^- is bound (15). In the wild-type nitrogenase, this analog induces a tight complex between the Fe protein and the MoFe protein (15, 36). This complex does not exhibit the normal Fe protein oxidation, only very slow ET from Fe protein to MoFe protein. In agreement with this, we observe no
ET over the normal observation times (0.5 s) for a complex that includes wild-type MoFe protein, reduced Fe protein, and this nucleotide analog (Figure 3-5, panel B). Likewise, the resolved phase of Fe protein oxidation is suppressed in the β-188Cys MoFe protein-Fe[MgADP + AlF₄⁻] protein complex; however, the complex shows that the burst oxidation of the Fe protein during the mixing time is preserved (Figure 3-5, panel A). Thus, both the non-hydrolyzable ATP analog, βγ-methylene-ATP, and the analog for ADP + Pi, MgADP + AlF₄⁻, promote the rapid $F^{1+} \rightarrow P^{1+}$ ET reaction, whereas neither analog supports the resolved ET from the Fe protein to either wild-type or β-188Cys MoFe protein that contains $P^N$.

![Figure 3-4](image.jpg)

**Figure 3-4: Electron transfer from Fe protein to MoFe proteins with the nucleotide analog βγ-methylene ATP.** Shown is the absorbance at 430 nm plotted against the time after mixing Fe protein (75 μM) and MoFe proteins (20 μM) in one syringe with the nucleotide analog βγ-methylene-ATP (18 mM). Shown are β-188Cys MoFe protein and Fe protein mixed against βγ-methylene-ATP (β-188Cys + βγ-methylene-ATP); and wild-type MoFe protein and Fe protein mixed against βγ-methylene-ATP (WT + βγ-methylene-ATP).
Figure 3-5: Electron transfer from Fe protein to MoFe proteins with the nucleotide analog MgADP-AlF_{4}⁻. Shown is the absorbance at 430 nm plotted against the time after mixing MoFe protein (20 µM) and Fe protein (75 µM) with buffer (without or with nucleotide, 10 mM) in the stopped-flow. Shown in panel A is β-188Cys MoFe protein and Fe protein mixed with MgADP-AlF_{4}⁻ (β-188Cys + MgADP-AlF_{4}, purple); with MgADP (β-188Cys + MgADP, green); without nucleotide (β-188Cys - nucleotide, light green). Also shown is the difference spectrum ((β-188Cys+ADP-AlF_{4}) - (β-188Cys + ADP), grey). Shown in panel B is wild-type MoFe protein and Fe protein mixed with MgADP (WT + ADP, green); with ADP-AlF_{4}⁻ (WT + ADP-AlF_{4}, purple); and without nucleotide (WT - nucleotide, light green).

The role of FeMo-cofactor in the ET. Electrons transferred into the MoFe protein ultimately end up on FeMo-cofactor. If the ET burst observed when β-188Cys MoFe protein and Fe protein are mixed with MgATP indeed reflects ET from the Fe protein [4Fe-4S] cluster to P^{1+}, then the FeMo-cofactor should not be involved in this ET event and its absence should have no influence on the observed ET burst. To test this, we purified variants of both wild-type and β-188Cys MoFe proteins that do not contain FeMo-cofactor (called apo-wild-type and apo-β-188Cys MoFe proteins) (4, 23). When apo-wild-type MoFe protein is used, no ET is observed (Figure 3-6). When the apo-β-188Cys MoFe protein and Fe protein were mixed against MgATP, the stopped-flow trace exhibited an
ET burst that was indistinguishable from that seen for the FeMo-cofactor containing \( \beta \)-188\text{Cys} MoFe protein (Figure 3-6), while the slower, resolved phase was absent. These observations confirm that the kinetically resolved ET represents \( P^N \rightarrow M^N \) ET, whereas the burst observed for the \( \beta \)-188\text{Cys} is \( F^{1+} \rightarrow P^{1+} \) ET.

**Figure 3-6: Electron transfer from Fe protein to apo-MoFe proteins.** Shown is the absorbance at 430 nm plotted against the time after mixing Fe protein (75 \( \mu \)M) and MoFe protein (42 \( \mu \)M) with buffer (without or with nucleotides, 10 \( \mu \)M) in the stopped-flow. Shown are apo-\( \beta \)-188\text{Cys} MoFe and Fe protein mixed with MgATP (Apo-\( \beta \)-188\text{Cys} + ATP); apo-wild-type MoFe protein and Fe protein mixed with MgATP (Apo-WT + ATP); and apo-\( \beta \)-188\text{Cys} MoFe and Fe protein mixed with buffer lacking nucleotides (apo-\( \beta \)-188\text{Cys} - nucleotides). See the legend to Figure 3-6 for other details.

**Conformationally gated ET.** Recently, it was shown that the rate of ET from the Fe protein to the wild-type MoFe protein is sensitive to osmotic pressure, indicating that the rate-limiting step in the ET reaction is gated by large protein conformational changes (6). Changes in osmotic pressure modulate the energetics of reactions that involve conformational changes in which the number of bound waters is changed. The rate constant \( (k_m) \) for such a process varies exponentially with the molality \( (m) \) of added
solute according to the equation, $k_m \propto \exp[-(\Delta n/55.6)m]$, where $\Delta n$ is the number of waters absorbed in the transformation (6).

![Figure 3-7: Osmotic pressure effects on electron transfer reactions from Fe protein to MoFe protein.](image)

Repeating this experiment with $\beta$-$188^{\text{Cys}}$ MoFe protein shows that the relative proportions of the burst and resolved ET phases are independent of the presence of sucrose as osmolyte up to a sucrose concentration of $\sim 2 \text{ m}$ (Figure 3-7). This indicates that the presence of the osmolyte does not alter the reduction potential of P cluster, which would alter the relative proportions of $P^{1+}$ and $P^N$ forms and thus the relative contributions of the two kinetic phases. Unlike the burst phase, the resolved phase of the ET is dependent on osmotic pressure. The inset to Figure 3-7 shows that rate constants ($k_{obs}(m)$) for ET from Fe protein to either the wild-type MoFe protein or the resolved
phase ET to the $\beta$-188$^{\text{Cys}}$ MoFe protein have essentially identical responses to sucrose addition, with the rate constants depending exponentially on the molality ($m$) of added sucrose with a slope corresponding to $\Delta n \sim +80$ waters. Taking roughly one water to be bound per $\sim 10 \text{ Å}^2$ of exposed surface, $\Delta n \sim 80$ waters would correspond to a conformational transition in which $\sim 800 \text{ Å}^2$ of surface becomes exposed (6).

The present results thus indicate that ET to the MoFe state that contains $\text{P}^\text{N}$ and $\text{M}^\text{N}$ is conformationally gated in both wild-type MoFe and $\beta$-188$^{\text{Cys}}$ MoFe protein, and that the amino acid substitution of $\beta$-188$^{\text{Ser}}$ by a cysteine does not alter the conformational gate. In contrast, the presence of sucrose osmolyte up to 2 $m$ does not affect the burst ET to the point where it becomes visible after the dead-time of the instrument. These results suggest that whereas the resolved oxidation of $\text{F}^{1+}$ by the MoFe protein in the $\text{P}^\text{N}/\text{M}^\text{N}$ state is conformationally gated, the rapid $\text{F}^{+1} \rightarrow \text{P}^{1+}$ ET event is not.

**Discussion**

While a deficit-spending scheme for nitrogenase ET events is the most reasonable of the three proposed, this model has proven difficult to test experimentally. For example, until now it has not been possible to measure the contributions of the individual inter- and intra-molecular ET events postulated by this model ($k_{FP}$ and $k_{PM}$, respectively, [Figure 3-1, panel B]) to the observed ET rate constant ($k_{obs}$). We discuss how the experiments described in this report support a deficit-spending model of ET in nitrogenase and further argue against a hopping ET scheme, and provide insights into the roles of nucleotides in ET. We conclude by presenting a model for the sequence of events for the entire nitrogenase ET cycle that integrates the present results with earlier findings.
ET by a deficit-spending mechanism. In the present work, the deficit spending model was addressed by using the β-188Cys-substituted MoFe protein, which has P^{1+} in the resting state (4). This situation offered the opportunity to directly monitor the F^{1+} → P^{1+} ET event (backfill) without the need for a prior P^N → M^N ET step. For this substituted protein, the rapid intermolecular ET event from the Fe protein to the MoFe protein could be assigned to the F^{1+} → P^{1+} ET event, which was found to be faster than the dead-time of the stopped-flow instrument with a rate constant that can be estimated $k_{fp} > 1700 \text{ s}^{-1}$. This rate constant is far greater than $k_{obs} = 168 \text{ s}^{-1}$ observed for ET from the Fe protein to MoFe protein in the P^N/M^N state, indicating that intramolecular P^N → M^N ET is the rate-limiting step.

In the ET traces for β-188Cys-substituted MoFe protein, the initial ET burst assigned to F^{1+} → P^{1+} is followed by a slower resolved ET phase. This slower phase is assigned to ET from the Fe protein to the β-188Cys MoFe protein in the P^N/M^N state, and its rate constant is the same as that observed for ET from the Fe protein to the wild-type MoFe protein in the P^N/M^N state. Thus, measurement of electron transfer using the β-188Cys-substituted MoFe protein reveals both the F^{1+} → P^{1+} ET event observed as a burst and the P^N → M^N ET event observed as a slower, resolved phase.

These observations are in full accord with expectations based on the deficit-spending model. In this model, the P^N → M^N ET event is the rate-limiting step in ET from the Fe protein to the MoFe protein, and the observed ET rate constant for the oxidation of F^{1+} by the MoFe protein in the P^N/M^N state is assigned to this event: $k_{obs} = k_{pm}$. The subsequent ‘backfill’ ET ($k_{fp}$) reaction, F^{1+} → P^{1+}, does not contribute to the observed ET because it is at least 10-times faster. This sequence of ET events, with the
corresponding large differences in rate constants, provides an explanation for the fact that only very low levels of a P\textsuperscript{i+} or P\textsuperscript{2+} state have been observed by electron paramagnetic resonance (EPR) spectroscopy when the nitrogenase complex is freeze-trapped during steady-state turnover \((I, 37, 38)\). Although the P\textsuperscript{i+} state is EPR active, it cannot build up to any appreciable concentration in the deficit-spending mechanism because the rate of reduction of P\textsuperscript{i+} by the Fe protein (F\textsuperscript{i+} → P\textsuperscript{i+}) is much faster than the rate of oxidation of P\textsuperscript{N} by M cluster (P\textsuperscript{N} → M\textsuperscript{N}), resulting in mostly the P\textsuperscript{N} state (which is EPR silent) under steady-state turnover.

**Gated ET.** The rate constant for ET from the Fe protein to the MoFe protein \((k_{\text{obs}})\) is sensitive to osmotic pressure, which has been assigned to protein conformational changes that gate the intermolecular ET event (6). This conformational change was calculated to involve exposure of a substantial (>800 \(\text{A}^2\)) protein surface area (6). The present study shows no modulation of the rapid backfill ET (F\textsuperscript{i+} → P\textsuperscript{i+}) by changes in osmotic pressure, suggesting that this ET step is not gated by protein conformational changes; in any case, this step is so rapid that it cannot be involved in determining the observed rate of ET. Thus, it can be concluded that the protein conformational gate for ET applies specifically to the P\textsuperscript{N} → M\textsuperscript{N} intramolecular ET step.

What conformational processes could be coupled to the P\textsuperscript{N} → M\textsuperscript{N} ET reaction? Examination of X-ray structures of various complexes of the Fe protein with the MoFe protein (with or without nucleotides bound) reveal large changes in the Fe protein-MoFe protein interface, involving changes in exposed protein surface area, but do not reveal any significant changes within the MoFe protein that could explain how the Fe protein with two bound 2ATP might activate the P\textsuperscript{N} → M\textsuperscript{N} ET (14). An attractive “compound
gate" model can be developed, however, from examination of the X-ray structure of an oxidized MoFe protein (P\textsuperscript{1+}, M\textsuperscript{N}) (39). From this structure, the P\textsuperscript{2+} cluster is stabilized by the coordination of the side-chain of β-188\textsuperscript{Ser} to an Fe atom of the P cluster (39). We suggest that conformational activation of resting-state MoFe protein by an as-yet unidentified change in the Fe protein-MoFe protein interface could cause the β-188\textsuperscript{Ser} side-chain to transiently coordinate to an Fe atom of P\textsuperscript{N}, thereby creating an activated state of the P cluster (designated P\textsuperscript{N*}) with a lowering of the potential to the point that P\textsuperscript{N*} →M\textsuperscript{N} ET becomes favorable. For the β-188\textsuperscript{Cys} MoFe protein, the assumption is that a cysteine at this position shifts the equilibrium in favor of the state with the cysteinate bound to the Fe of the P cluster, and that the redox equilibrium between P\textsuperscript{N} and P\textsuperscript{1+} in the presence of dithionite also involves a ligation equilibrium, with ∼65% of the P cluster in the form of P\textsuperscript{1+} with bound cysteine, and ∼35% in the form of P\textsuperscript{N} with cysteine not bound as in the wild-type MoFe protein. This picture would account for the observation that the resolved ET phase for the β-188\textsuperscript{Cys} MoFe protein behaves the same as that for the wild-type MoFe protein. Lowe and Thorneley (26) further treat ET as occurring only as part of the transfer of an ‘H-atom’, namely an electron and a proton.

Two recent reports describing ET to the MoFe protein in the absence of the Fe protein also support an activation step that could facilitate the intramolecular P\textsuperscript{N*} →M\textsuperscript{N} ET step. It has been found that attaching a Ru-ligand complex onto the surface of the MoFe protein near the P cluster can be used to photoinduce ET from the Ru-ligand complex into the MoFe protein, allowing acetylene reduction (40). In the absence of the Fe protein, the rates of acetylene reduction were very low (0.2% of the rate observed with Fe protein and ATP), consistent with the need for some activation of the P → M ET
process to promote full substrate reduction rates. In a separate study, it was found that a low potential Eu(II)-ligand complex could be used to drive electrons into the MoFe protein in the absence of the Fe protein in support of reduction of hydrazine (H$_2$N-NH$_2$) to two ammonia (41). In this system, substrate reduction was observed only when an amino acid residue located between the P cluster and M cluster was substituted by another amino acid, suggesting that the conformational switch normally activated by Fe protein binding involves changes in residues located between the P cluster and FeMo-cofactor and that the key conformational changes can be partially mimicked by an amino acid substitution (41).

Further evidence against a hopping model. The strongest argument against a hopping or sequential ET model for nitrogenase is the improbability that the P cluster can be reduced beyond the all ferrous P$^N$ state, as would occur following an initial ET event: F$^{1+} \rightarrow$ P$^N$. Super-reduction of all-ferrous P$^N$ would require the formation of an Fe(I) with tetrahedral coordination by sulfur, an unknown Fe state. However, our results show that this step would have to be gated, and one might imagine that the activation process might involve a transient loss of a cysteinate ligand from one of the Fe atoms. This would create an activated P$^{N+}$ cluster state that has a three-coordinate Fe(II) and a more positive reduction potential. Such a state might accept an electron from F$^{1+}$, and then re-binding of the cysteinate could create a strongly reducing P$^S$ state that undergoes prompt follow-up P$^S \rightarrow$M$^N$ ET.

To test this possibility, we examined ET from F$^{1+}$ to a MoFe protein where one of the cysteine ligands to the P cluster was removed (β-153CysΔ). It was found that the rate
of oxidation of $F^{1+}$ to this MoFe protein variant is unchanged from the rate to the wild-type MoFe protein, providing further evidence against the hopping ET model.

**Role for nucleotides.** MgATP binding and hydrolysis are both essential to intermolecular ET from the Fe protein to the MoFe protein (42). In the absence of ATP hydrolysis, the Fe protein does not transfer an electron to the resting-state MoFe protein ($P^N, M^N$). The role of ATP hydrolysis in this ET process cannot be filled by the non-hydrolyzable ATP analog $\beta\gamma$-CH$_2$-ATP or the ADP+Pi analog, ADP-AlF$_4$. However, results reported here reveal that ATP hydrolysis is not necessary for the backfill ET ($F^{1+} \rightarrow P^{1+}$). Rather, the burst ET to the $\beta$-188$^{Cys}$ MoFe protein is supported by both $\beta\gamma$-CH$_2$-ATP and ADP-AlF$_4$. Thus, the backfill electron transfer event ($F^{1+} \rightarrow P^{1+}$) is not conformationally gated and requires nucleotide binding but not hydrolysis.

In contrast to this finding, neither of the ATP analogs supported the slower, resolved phase ET assigned to the $P^N \rightarrow M^N$ ET reaction, pointing to this ET step as requiring an ATP-bound state that has not yet been achieved. Both MgATP and MgADP binding to the Fe protein have been shown to equally lower the midpoint reduction potential of the $[4Fe-4S]^{2+/1+}$ couple (33), yet MgADP does not support either the $P^N \rightarrow M^N$ or the $F^{1+} \rightarrow P^{1+}$ ET events, revealing that it is not simply an issue of driving force for the ET. Thus, the studies presented here pinpoint (i) the conformational gate and (ii) the specific need for ATP binding/hydrolysis to the $P^N \rightarrow M^N$ ET step.

**Summary sequence of events.** The findings presented here, taken with earlier findings, can be used to propose an overall model for the events associated with ET from the Fe protein to the MoFe protein (Figure 3-8). In this model, the ET process is initiated by the association of the Fe protein (2MgATP) with the resting state of the
MoFe protein (Figure 3-8, step 1). Although the binding of 2 MgATP to the Fe protein causes a major structural change and lowers its reduction potential by -120 mV (43), the osmotic pressure dependence of the oxidation of F\textsuperscript{1+} indicates that the complex initially formed between Fe protein-2MgATP and MoFe protein is not competent for ET (6). Instead, ET is initiated by a conformational activation (Figure 3-8, step 2), presumably involving motion at the Fe protein-MoFe protein interface, with exposure of as many as ~80 water molecules (6). In this model, the interface motion is coupled to a conformational change (second gate) (Figure 3-8, step 3) within the MoFe protein that shifts the P cluster to an activated state (P\textsuperscript{N*}) that transiently favors intramolecular ET from the P\textsuperscript{N*} state to the M\textsuperscript{N} cluster (Figure 3-8, step 4) (6). The final step in this sequence involves the rapid ‘backfill’ ET to the oxidized P cluster (F\textsuperscript{1+} \rightarrow P\textsuperscript{1+}), resulting in an oxidized Fe protein and a reduced MoFe protein that has its metal clusters in the P\textsuperscript{N} and M\textsuperscript{R} states (Figure 3-8, step 5).

The specific contributions for MgATP binding and hydrolysis to the individual steps remains to be resolved, but our studies show that the backfill ET process requires nucleotide binding but not hydrolysis, while the activated P\textsuperscript{N*} \rightarrow M\textsuperscript{N} ET initial step is not facilitated by any of the ATP analogs examined, requiring ATP itself. Studies of a wide range of ATP utilizing enzymes, in particular transport proteins, have led to the general
Figure 3-8: Models for nitrogenase mechanism. Shown is one catalytic cycle for a functional half of nitrogenase composed of an αβ-unit of the MoFe protein (rectangles) and a homodimeric Fe protein (ovals) with the associated metal clusters: \( F = [4\text{Fe-4S}] \) in the 1+ or 2+ oxidation states, \( P = \text{P cluster} \) in the N or ox oxidation states, and \( M = \text{FeMo-cofactor} \) in the N or R oxidation states. The cycle starts at the top with reduced Fe protein +2 MgATP associating with the αβ-unit of the MoFe protein (1). The remaining steps are numbered (clockwise): (2) the conformational gate, (3) the second gate, (4) the first ET \( (k_{PM}) \), (5) the backfill ET \( (k_{FP}) \), (6) ATP hydrolysis, (7) release of 2 Pi, (1) and replacement of oxidized Fe protein + 2MgADP with reduced Fe protein + 2MgATP.

View that energy transduction in these proteins occurs during the binding of MgATP, which induces a conformational state activated for the particular function of the ATPase (‘taut’ state). ATP hydrolysis occurs after the functional/catalytic step, initiating
relaxation from the taut state to a relaxed state with ADP bound, which in turn leads to a resetting of the system for another round of catalysis (34, 35).

While the order of ET and MgATP hydrolysis in nitrogenase has not been well established by previous work, the consensus model is that MgATP hydrolysis follows the ET events (1, 44, 45), in agreement with the current views of ATPase function in other proteins. MgATP binding to the Fe protein is known to induce protein conformational changes that increase the affinity of the Fe protein for binding to the MoFe protein, but given that further activation is needed to induce ET, it would appear that conversion to some as yet uncharacterized ATP binding state (perhaps a true transition state) is required to activate the P^N→M^N ET. In this model, the MgATP hydrolysis reaction is placed late in the reaction, after all ET events, although as just noted, this order has not been clearly resolved (Figure 3-8, step 6). The present results reveal that the backfill ET from F^{+1}→P^{1+} requires an ATP-bound state, but does not require ATP hydrolysis. It thus seems reasonable to suggest that nitrogenase functions similarly to other ATPases, and that ATP hydrolysis during nitrogenase ET is a late event that relaxes the Fe protein for dissociation from the MoFe protein. The rate limiting step for the overall reduction of substrates has been shown to be dissociation of the oxidized Fe protein with two bound 2MgADP from the MoFe protein (Figure 3-8, step 7), with a rate constant of ~ 6 s^{-1} (26).

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CHAPTER 4

TEMPERATURE INVARIANCE OF THE NITROGENASE ELECTRON TRANSFER MECHANISM

ABSTRACT

Earlier studies on electron transfer (ET) from the nitrogenase Fe protein to the MoFe protein concluded that the mechanism for ET changed during cooling from 25°C to 5°C, based on the observation that the rate constant for Fe protein to MoFe protein ET decreases strongly, with a non-linear Arrhenius plot. They further indicated that the ET was reversible, with complete ET at ambient but with an equilibrium constant near unity at 5°C. These studies were carried out with buffers having a strong temperature coefficient. We have examined the temperature variation in the kinetics of oxidation of the Fe protein by the MoFe protein at constant pH = 7.4 fixed by the buffer MOPS, which has a very small temperature coefficient. Using MOPS, we also observe temperature dependent ET rate constants, with non-linear Arrhenius plots. But, we find that ET is gated across the temperature range by a conformational change that involves the binding of numerous water molecules, consistent with an unchanging ET mechanism. Furthermore, there is no sKIE throughout the temperature range studied, again consistent with an unchanging mechanism In addition, the non-linear Arrhenius plots are explained by the change in heat capacity caused by the binding of waters in an invariant gating ET mechanism.

mechanism. Together, these observations contradict the idea of a change in ET mechanism with cooling. Finally, the extent of ET at constant pH does not change significantly with temperature, in contrast to the previously proposed change in ET equilibrium.

**INTRODUCTION**

Nitrogenase catalyzes the multi-electron reduction of dinitrogen (N₂) to two ammonia (NH₃), a reaction that is the major contributor to N₂ fixation in the biosphere.⁴ The enzyme comprises two component proteins, called the Fe protein and the MoFe protein, which associate and dissociate each time an electron is delivered from the Fe protein to the MoFe protein, which contains the catalytically active FeMo-cofactor, [7Fe, Mo, 9S; C; homocitrate].⁴ This process of electron transfer (ET) from Fe to MoFe protein can be summarized by the deceptively simple kinetic scheme of Eq 1: the reduced Fe protein with two bound ATP [Fe₆(MgATP)₂] rapidly forms an activated complex with the MoFe protein, where the rate constant for binding is k₁ and that for dissociation is k₋₁.

$$\text{Fe}_6(\text{MgATP})_2 + \text{MoFe} \xleftrightarrow{k_1 \quad k_-1} \text{Fe}_6(\text{MgATP})_2;\text{MoFe} \xleftrightarrow{k_2 \quad k_-2} \text{Fe}_6(\text{MgAXP})_2;\text{MoFe}^{\text{red}}$$

(1)

ET from the Fe protein to the MoFe protein then creates a one-electron reduced MoFe protein denoted MoFe^{red}. This forward ET is characterized by a single ET rate constant k₂; the possibility of reverse ET is incorporated by inclusion of rate constant k₋₂.
A key challenge in defining the mechanism of nitrogenase is to understand the steps in the kinetic scheme of eq 1 at a detailed molecular level. The process begins when the ATP-bound Fe protein with a reduced [4Fe-4S] cluster docks to the MoFe protein, **Figure 4-1.** The subsequent intracomplex ET from the Fe protein to the MoFe protein was shown to be gated in a study that employed variations in viscosities/osmotic pressures.\(^5\) The variation of the ET rate constant with osmotic pressure showed that ~80 waters bind to the protein surfaces during the reaction. This binding induces conformational changes to the MoFe protein that initiate ‘gated’ electron transfer (ET)\(^6,\)\(^7\) within the MoFe protein, from the P cluster [8Fe-7S] to FeMo-cofactor.\(^5\) This process in turn creates an electron ‘deficit’ on the P cluster, which is back-filled by rapid electron transfer from the Fe protein to the oxidized P cluster.

X-ray crystal structures of MoFe protein:Fe protein complexes have been solved in various states of nucleotide binding.\(^8,\)\(^9\) From these structures, it is clear that the Fe protein exhibits major conformational differences in different nucleotide-bound states, and that it binds at different sites on the surface of the MoFe protein in the different states, suggesting that motion between sites is involved in the gating and inspiring the scheme implied in **Figure 4-1.** However, these structures do not show any differences within the MoFe protein, thus leaving open the nature of the motions within the MoFe protein that contribute to the gated ET.\(^10,\)\(^11,\)\(^12\)

This ET process, which has been termed “deficit spending”\(^13\) is linked to the hydrolysis of two ATP molecules bound to the Fe protein. Although ATP hydrolysis may well follow the ET reaction, in analogy to other NTPases,\(^14\) the order of electron transfer
Fig 4-1. Events proposed as occurring at the interface of the MoFe protein and the Fe protein as the latter delivers an electron to the former. Here MoFe protein is represented by the semi-circle. The Fe protein (coming in from the left) binds two ATPs. The binding of ATP to the Fe protein causes conformational changes in the Fe protein which bring the [4Fe-4S] cluster closer to the surface of the protein. Conformational changes subsequent to binding ‘gate’ the conversion to an ET-active state. As the order of ET and ADP hydrolysis is uncertain, in this state the nucleotide is written‘AXP’. The final step is the departure of the Fe protein (right).
and ATP hydrolysis has not been clearly established,\(^{(2, 15, 16)}\) and so in Fig 4-1 the state of the nucleotide in the ET-active conformation is not specified (X = T or D). The absence of a solvent kinetic isotope effect (sKIE)\(^{(5)}\) at ambient temperature implied that ATP hydrolysis, which should be slowed by the presence of D versus H, is not involved in the rate-limiting step of ET.

Equation 1 is written as though the ET step is reversible. Although ET is quantitative at ambient temperature, previous studies found that the absorbance change associated with the oxidation of Fe\(^{\text{red}}\) by MoFe protein (ΔA) decreases as the temperature is lowered.\(^{(2, 16)}\) This was interpreted as indicating that ET is an equilibrium process, with an equilibrium constant at ambient temperature that is large, \(K_2 = k_2/k_{-2} \gg 1\), corresponding to quantitative ET, but that decreases to \(K_2 \sim 1\) at 5°C. If ET is indeed reversible, then either it precedes ATP hydrolysis, or the hydrolysis also must be reversible.

A number of studies investigated the temperature dependence of ET as a means of exploring mechanism.\(^{(2, 16, 17)}\) They suggested that the actual mechanism of ET from the Fe protein to the MoFe protein changes upon cooling. Thus, a non-linear Arrhenius plot of the observed ET rate constant versus inverse temperature was interpreted as having a ‘break’ at ~14-16°C, and used as the basis for a suggestion that one mechanism operates at higher temperatures, another at lower, each with a characteristic activation enthalpy. The proposal of a change in mechanism was supported by stopped-flow (SF) isothermal calorimetry (ITC) measurements, which were interpreted as showing that in the low-temperature regime the hydrolysis of ATP precedes the electron transfer event, whereas at ambient temperature the ET event occurs prior to ATP hydrolysis.\(^{(16)}\)
Here, we have re-examined the temperature variations at constant pH in the extent and rate constants for oxidation of the Fe protein by the MoFe protein within the pre-formed complex, and have tested for the persistence of gating at low temperature and for the appearance of an sKIE on ET. Our findings show that the ET mechanism does not change during cooling at constant pH.

**MATERIALS AND METHODS**

All reagents for these experiments were purchased from Sigma-Aldrich Chemicals (St. Louis, MO). Wild type MoFe protein was expressed in *Azotobacter vinelandii* strain DJ995, while wild type Fe protein was obtained from strain DJ884 grown as described previously.\(^{(18)}\) MoFe protein purification was performed using previously described metal affinity chromatography method due to the seven-histidine tag on the \(\alpha\)-subunit of the protein while Fe protein was purified, with some modifications, using ion exchange and size exclusion as previously described.\(^{(18,19)}\) Both proteins utilized in these experiments were found to be of greater than 95% purity based on SDS-PAGE analysis using Coomassie blue staining. Septum sealed degassed vials, under an argon atmosphere were used to determine the activity of the proteins. The wild type Fe protein activity was found to be \(~1750\) nmols while the wild type MoFe protein activity was found to be \(~2100\) nmols. Both activities were similar to previously reported values for the activity of these proteins.\(^{(20)}\) All liquids were transferred using gastight syringes.

The increase in absorbance at 430 nm was monitored to follow the ET event from the Fe protein to the MoFe protein. This increase in absorbance occurs due to the Fe protein’s oxidation by one electron from its reduced \((1^+\) state to the \((2^+\) state; there is no
significant accompanying change in absorbance of the MoFe protein. The change of absorbance as a function of time after the mixing of the two proteins was observed using a Hi-Tech SF61 stopped flow (SF) UV-visible spectrophotometer equipped with data acquisition software (Salisbury, Witts, U.K.). The SF unit of the spectrophotometer was housed inside a nitrogen filled glovebox. The temperature during the experiments was controlled by a circulating water bath housed outside the glove box.\(^{(21)}\) All reactions, except for the change in temperature experiments, were carried out at 25°C. Differing buffer systems, HEPES and MOPS, pH 7.4 were used with 10mM dithionite. Due to the change in pH with temperature in the HEPES buffer, MOPS was utilized for the temperature dependency experiments to keep the pH constant.\(^{(22)}\) During the SF experiments Fe and MoFe protein were contained in one drive syringe of the unit while the other syringe contained MgCl\(_2\) and ATP. Both syringes contained buffer (HEPES/MOPS 100mM pH 7.4) and 10mM dithionite. The dead time of the Hi-Tech SF61 SF system is 2 ms.

In the present experiments, the formation of the \([\text{Fe}^{\text{red}}(\text{MgATP}*)_2; \text{MoFe}]\) complex is essentially quantitative and complete within the deadtime of the experiment. Thus, the SF trace monitors the intra-complex oxidation, \(\text{Fe}^{\text{red}} \rightarrow \text{Fe}^{\text{ox}}\), by MoFe protein, the second step in Eq 1. In the SF experiment, the absorbance of the mixed solutions prior to \(\text{Fe}^{\text{red}} \rightarrow \text{Fe}^{\text{ox}}\) oxidation is obtained with a non-reactive ‘push’ in which ATP is omitted, then the instrument output is ‘nulled’ by subtraction of this absorbance. As a result, the SF traces represent the time-variation of the absorbance-difference caused by reaction, relative to that of the initial mixed, unreacted solution.
Previous studies typically obtained the rate constant for the oxidation of Fe\textsuperscript{red}, denoted here $k_{\text{obs}}$, by fitting the initial portion of the SF traces with an exponential rise-to-maximum.\textsuperscript{(25, 16, 17)} However, as can be seen in the data presented below (Figure 4-2) and in SI, the SF traces for the oxidation of Fe\textsuperscript{red} by MoFe protein in general have a more complex shape, which indicates that the oxidation of Fe\textsuperscript{red} is followed by a slower reaction to a state with a smaller absorbance-difference. Haaker and coworkers attribute this follow-up reaction to hydrolysis of ATP. Alternatively, we suggest this may represent a partial re-reduction of the resulting Fe\textsuperscript{ox} back to Fe\textsuperscript{red} that occurs as part of a complex set of reactions that involves dissociation of Fe\textsuperscript{ox}, re-reduction to Fe\textsuperscript{red}, and re-binding to MoFe.\textsuperscript{(23)}

In this report we are interested only in the rate constant and absorbance-difference associated with the initial intra-complex oxidation, Fe\textsuperscript{red} $\rightarrow$ Fe\textsuperscript{ox}, by MoFe protein. As shown in SI, neither of these parameters is accurately derived with a rise-to-maximum fit if the followup phase of the SF trace is appreciable, so we have employed the simplest possible kinetic model to treat the observed traces. We find that the overall traces can be well-modeled by a kinetic scheme that includes two sequential first-order reactions, Eq 2,

$$
[\text{Fe}^{\text{red}}, \text{MoFe}^N] \rightarrow [\text{Fe}^{\text{ox}}, \text{MoFe}^{\text{red}}] \rightarrow \text{C}
$$

(2)

the first representing the initial oxidation of Fe\textsuperscript{red} by MoFe protein, the second a followup reaction to an unspecified state, denoted C. This model leads to an equation for the resolved time-course of the absorbance difference, $\Delta A(\tau)$, given by Eq 3:
\[ \Delta A(\tau) = \Delta A_0 + \Delta A \left[ \left( \frac{k_{\text{obs}}}{k_{\text{obs}} - k_{\text{Dr}}} \right) e^{-k_{\text{obs}} \tau} - e^{-k_{\text{obs}} \tau} \right] + \Delta A_{\text{Dr}} \left[ 1 - \left( \frac{-k_{\text{Dr}}}{k_{\text{obs}} - k_{\text{Dr}}} \right) e^{-k_{\text{obs}} \tau} + \left( \frac{k_{\text{obs}}}{k_{\text{obs}} - k_{\text{Dr}}} \right) e^{-k_{\text{obs}} \tau} \right] \] (3)

where \( \tau = t - t_{\text{DT}} \), with \( t_{\text{DT}} \) being the time at which mixing artifacts have subsided (dead-time). In this equation, \( k_{\text{obs}} \) is the rate constant for the oxidation of Fe\textsuperscript{red} by MoFe protein, and \( k_{\text{Dr}} \) is the rate constant for the follow-up process. This equation takes into account possible reaction during the dead-time leading to an absorbance-difference, \( \Delta A_0 \). As a result of such a phenomenon, the total absorbance-difference associated with the first-step intracomplex oxidation of Fe\textsuperscript{red} (Eq 2), denoted here \( \Delta A \), equals the absorbance change associated with the initial oxidation step in the resolved phase of Eq 3, \( \Delta A_r \), plus the absorbance change from the portion of the oxidation reaction that occurs during the dead-time, \( \Delta A_0: \Delta A = \Delta A_0 + \Delta A_r \). As incorporated into Eq 3, State C is characterized by a net absorbance-difference, \( \Delta A_{\text{Dr}} \), relative to the initial state. In the Results section we discuss only the parameters associated with the initial oxidation of Fe\textsuperscript{red} by MoFe protein: \( k_{\text{obs}} \) and \( \Delta A \); for completeness only, values for the parameters associated with the unspecified followup process, \( k_{\text{Dr}} \) and \( \Delta A_{\text{Dr}} \), are given in SI.

As discussed in detail in the SI, the fits to Eq 3 give as much as a two-fold smaller rate constant, \( k_{\text{obs}} \), than a rise-to-maximum fit. However, we note that such trends as changes with temperature or osmotic pressure, which are the focus of this work, are comparably represented by the two approaches (Figure 4-S1).
RESULTS AND ANALYSIS

Influence of temperature and solvent isotope on ET

*Figure 4-2* shows the SF traces at different temperatures for the oxidation of Fe$^{\text{red}}$ by the MoFe protein in H$_2$O (*top*) and D$_2$O (*bottom*) solutions with MOPS as the buffer. As this buffer has a negligible temperature coefficient of pKa vs T,$^{(22)}$ these measurements thus are carried out with essentially invariant pH = pD = 7.4 over the temperature range, 25 - 5°C. As discussed in *Materials and Methods*, at temperatures above 5°C the absorbance changes associated with the initial oxidation of Fe$^{\text{red}}$ by MoFe do not show a simple rise-to-maximum; the initial absorbance increase is followed by a decrease in absorbance associated with re-reduction of Fe$^{\text{red}}$ or the change in absorbance of Fe$^{\text{ox}}$ due to hydrolysis of ATP. To obtain the rate constants for the initial Fe$^{\text{red}}$ oxidation, the traces were fit to a sequential model in which the oxidation of Fe$^{\text{red}}$ is followed by a slower re-reduction process, as described in *Materials & Methods*. In discussing these results, we consider only the rate constants and absorbance changes associated with the initial Fe$^{\text{red}}$ oxidation.

To test for the congruence of the current results to those reported earlier, the traces of *Figure 4-2* were truncated and fit to a rise-to-maximum. As mentioned in *Materials and Methods* and shown in *SI*, such fits typically give larger values for k$_{\text{obs}}$, in keeping with those reported earlier,$^{(2,16,17)}$ but the trends with temperature are completely equivalent,

*Figure 4-S1*. Thus, trends reported in this study can be appropriately compared to those in earlier reports. However, we believe that the magnitudes of k$_{\text{obs}}$ based on fits to Eq 3 which are notably lower than earlier values, are more appropriate.
The temperature responses of the rate constants and absorbance changes associated with the initial Fe\textsuperscript{red} oxidation for D\textsubscript{2}O and H\textsubscript{2}O solutions (MOPS pH = pD = 7.4), denoted $k_{\text{obs}}$ and $\Delta A$, are plotted, respectively, in Figure 4-3 and Figure 4-4. The observed ET rate constants decrease strongly with temperature (Figure 4-3), and the $k_{\text{obs}}$ values for H\textsubscript{2}O and D\textsubscript{2}O solutions with MOPS are the same within experimental error. The finding that $k_{\text{obs}}(\text{H}_2\text{O}) = k_{\text{obs}}(\text{D}_2\text{O})$ at ambient temperature confirms our earlier report that there is no sKIE at 25°C; the equivalence of the rate constants at all temperature shows that an sKIE does not emerge upon cooling.
Figure 4-2. Temperature-response stopped-flow traces for oxidation of Fe^{red} within [Fe^{red}(MgATP)_{2};MoFe] complex in H_{2}O and D_{2}O buffers. Top: H_{2}O, Bottom: D_{2}O. Plots are labeled with the temperature (°C) from 25°C to 5°C in the respective color of the traces. White lines through each trace are fits as described in *Materials and Methods*.

Figure 4-3. Semi-logarithmic plot of the observed rate constants, k_{obs} for D_{2}O (blue symbols) and H_{2}O (red symbols). Individual experiments are visualized by open, closed, or crossed points. Purple solid line is a fit to all data sets, assuming that there is no difference in k_{obs} for H_{2}O and D_{2}O buffers.
Temperature dependent ET equilibrium?

Thorneley et al. reported that the absorbance change (ΔA) associated with Fe\textsuperscript{red} → MoFe ET in H\textsubscript{2}O buffer diminishes with decreasing temperature,\textsuperscript{(2)} a finding corroborated by Mensink and Haaker.\textsuperscript{(16)} The absorbance decrease was interpreted in terms of a temperature dependence of an ET equilibrium between the two proteins, as represented by the second step of Eq 1. It was proposed that ET was essentially quantitative at 25°C, \( k_2 \gg k_{-2} \) (Eq 1), but by 5°C only ~50% of the complexes showed ET (\( k_2 \approx k_{-2} \)).

There are multiple reasons for concern about these conclusions. First, the experiments were carried out with solutions that used TES/NaOH buffer, and as a result were not performed at constant pH as the temperature was lowered. TES buffers have a large temperature coefficient,\textsuperscript{(22)} causing an increase in pH from 7.4 to ~ 7.7 as the temperature is lowered from 25°C to 5°C. Second, the kinetic traces were fit to a simple rise-to-maximum function, which does not necessarily give a good representation of the maximum absorbance change when the reaction involves more than one stage (Eq 1). In particular, as has been reported, the kinetic traces often show a rise-to-maximum, slight fall, then slow rise.\textsuperscript{(2)}

We thus analyzed the temperature dependence of the absorbance changes associated with the ET kinetics measurements discussed above. Figure 4-4 shows that these absorbance changes fluctuate appreciably from one experiment to the next, even though the kinetic constants do not (Figure 4-3). However, within the error of our measurements the absorbance changes with MOPS as buffer are the same for H\textsubscript{2}O and D\textsubscript{2}O solutions, and are invariant with temperature; the same is true for the absorbance changes obtained from rise-to-maximum fits (Fig. S2). We tested whether the implied
constancy in the extent of ET is modified by a change with cooling of the difference between the extinction coefficients of oxidized and reduced Fe protein, \( \Delta \varepsilon = \varepsilon(\text{Fe}^{\text{ox}}) - \varepsilon(\text{Fe}^{\text{red}}) \). As shown in Figure 4-S3 within error, this difference is temperature invariant down to 5°C. We thus interpret the results of Figure 4-4 as showing that the quantitative ET in MOPS buffer at pH 7.4 is essentially unchanged during cooling at constant pH from 25°C to 5°C. In the description of nitrogenase ET given by Eq 1, the observed ET rate constant is related to the microscopic rate constants by, \( k_{\text{obs}} = k_2 + k_{-2} \). However, when ET is quantitative, the ET rate constants obey the inequality, \( k_2 \gg k_{-2} \), and thus we may assign \( k_{\text{obs}} = k_2 \) throughout the temperature range examined.
**Figure 4-4**: Temperature dependence of absorbance changes $\Delta A$, (Eq 3) obtained from intracomplex oxidation of $\text{Fe}^{\text{red}}(\text{ATP})_2$ by MoFe as a function of temperature. Purple dashed line is the average of all values.

To test whether the previously reported temperature response of $\Delta A$ associated with $\text{Fe}^{\text{red}} \rightarrow \text{MoFe}$ ET in H$_2$O buffer could be attributed to a change in pH upon cooling, the temperature dependence of ET was reexamined using samples prepared with HEPES/H$_2$O buffer (Figure 4-S4). HEPES has a temperature coefficient similar to that of TES, used by Haaker and coworkers; with HEPES as buffer the pH increases from 7.4 to ~ 7.8 as the temperature is lowered from 25°C to 5°C. As shown in Figure 4-S5, the absorbance change associated with $\text{Fe}^{\text{red}} \rightarrow \text{MoFe}$ protein ET for H$_2$O solutions of HEPES buffer indeed drops by roughly 50% from 25 to 5°C, whereas the absorbance change does not decrease for the D$_2$O solutions of HEPES buffer (Figure 4-S4), and if anything increases slightly with decreasing temperature. The differing behavior for the H$_2$O and D$_2$O solutions with HEPES presumably reflect an equilibrium isotope effect on the pH dependence of the extent of ET. This inference is supported by measurements over a range of pH values in HEPES buffer, all carried out at 5°C (Figure 4-S6 and 4-S7). These show that $k_{\text{obs}}$ does not change significantly with pH for either buffer; there is a substantial variation in $\Delta A$ with pH for the H$_2$O buffers, but negligible change for the D$_2$O buffers.

**Gating at Low Temperature?**

To reveal if conformational gating of ET from the Fe protein to the MoFe protein persists at temperatures below the apparent break temperature ($T_b \sim 14\,^\circ\text{C}-16\,^\circ\text{C}$) as concluded in earlier studies, we examined the influence of osmolytes on $k_{\text{obs}}$ for ET at
6°C. Sucrose, glycerol, and glucose were studied as osmolytes because they generated the strongest, intermediate, and weakest effects seen at ambient temperatures. As illustrated for sucrose as solute in Figure 4-5, inset, progressive additions of each osmolyte cause progressive decreases in $k_{\text{obs}}$ at 6°C, thus indicating that ET is under conformational control at this low temperature as well as at ambient temperatures. Given the observed control of ET by osmotic pressure effects at 6°C as well as at 25°C, we conclude that the gating mechanism that controls ET persists throughout the temperature range.

Figure 4-5. Plot of the solute-dependent osmotic effect on nitrogenase. Black = glycerol, red = glucose, blue = sucrose. Points and solid lines represent results at 6°C; dashed lines represent the previously published results at 25°C; glycerol alone could be examined to higher $m$ and the additional points are shown in Figure 4-S1. Inset:
Stopped-flow oxidation of Fe$^{\text{red}}$ within [Fe$^{\text{red}}$(MgATP)$_2$]:MoFe] complex with increasing amounts of sucrose.

Changes in osmotic pressure modulate the energetics of reactions that change the number of bound waters. The rate constant for such a process varies exponentially with the molality of added solute ($m$) according to the equation,\(^{(24)}\) $k(m) \propto \exp[-(\Delta n/55.6)m]$, where $\Delta n$ is the number of waters absorbed in the transformation. Logarithmic plots of $k_{\text{obs}}$ at 6°C vs molality for each solute employed are linear, as they are at ambient, Figure 4-5, revealing that the osmotic pressure effects that control ET at ambient persist at reduced temperatures. This linearity persists to higher osmolalities, as shown in Figure 4-S8. The slopes of these lines, which differ for the different osmolytes, reflect the numbers of waters absorbed during ET: $\text{slope} = -(\Delta n/55.6)$. At 25°C, $\Delta n = 50$-$80$ for the three osmolytes; at 5°C, $\Delta n = 30$-$50$. The decreased values of $\Delta n$ at 6°C suggest that the osmolytes interact more strongly with the surface as the temperature is decreased.

The absence of an H$_2$O vs. D$_2$O solvent kinetic isotope effect ($s$KIE) for ET between the Fe protein and the MoFe protein at 6°C indicates that ATP hydrolysis is not rate limiting at low temperatures, just as it is not rate limiting at 25°C. In combination with the unchanging osmotic pressure effects over the temperature range, the results indicate that the order of the ATP hydrolysis event is unaffected by the change in temperature, and are consistent with there being no change in the mechanism with changing temperature. This conclusion is contrary to the earlier ITC findings that ATP hydrolysis precedes electron transfer at lower temperatures.
Does the temperature variation of $k_{\text{obs}}$ imply a change of mechanism?

The Arrhenius plots of $\ln(k_{\text{obs}})$ vs $1/T$ presented in Figure 4-3 for solutions carried out with both $\text{H}_2\text{O}$ and $\text{D}_2\text{O}$ MOPS solutions are non-linear, consistent with earlier findings for experiments carried out with $\text{H}_2\text{O}$ buffer solutions.\(^{2,17}\) The earlier work interpreted this nonlinearity to indicate a change in ET mechanism as the temperature is lowered from ambient. Those temperature variations were analyzed in terms of an apparent “break” at $T_b \sim 14^\circ\text{C}-16^\circ\text{C}$ that separates two linear segments in the plots of $\ln(k_{\text{obs}})$ vs $1/T$. Each segment was fit to its own slope, $(-\Delta H^\dagger/R)$; each temperature range was assigned a different mechanism with a characteristic enthalpy of activation, $\Delta H^\dagger$, that is constant over the range. However, a real ‘break’ in slope is not possible for a finite system.\(^{25}\) Instead, our finding that waters become bound to the protein-protein complex during the ET process requires that the Arrhenius plot be curved, even though the mechanism does not change. The binding of $\Delta n$ waters during the ET reaction causes a change in the heat capacity, $\Delta C_p$, which one can approximate as,

$$\Delta C_p \sim \Delta n[C_{\text{pm}}(\text{ice}) - C_{\text{pm}}(\text{liquid})] \sim \Delta n[-37 J/K*\text{mol}].$$

This heat capacity difference in turn makes the enthalpy change, associated with the ET event temperature dependent, which induces curvature into the Arrhenius plot.

Taking the ET reaction as being governed by a conformational gating process that involves water binding,\(^{5}\) we now show that the temperature variation of the ET rate constant $k_{\text{obs}} = k_2$ can be described with a single set of thermodynamic parameters over the entire range of temperatures examined. Taking as a reference temperature that of the apparent “break”, $T_b$, one can write the enthalpy change for the ET reaction at any
temperature within this range as the sum of a temperature-independent enthalpy change, \( \Delta H(T_b) \), plus the temperature-dependent contribution associated with water binding, \(^{(26)}\)

\[
\Delta H(T) = \Delta H(T_b) + \Delta C_p (T - T_b)
\]

A Gibbs-Helmholtz equation for \( k_{\text{obs}} \) then leads to a temperature dependence, \( \text{Eq } 5 \). \(^{(26)}\)

\[
\ln \left( \frac{k_{\text{obs}}(T)}{k_{\text{obs}}(T_b)} \right) = -\frac{\Delta H(T_b)}{R} \left( \frac{1}{T} - \frac{1}{T_b} \right) + \frac{\Delta C_p}{R} \left[ \ln \left( \frac{T}{T_b} \right) + \frac{T_b}{T} - 1 \right]
\]

As shown in \textbf{Figure 4-3}, the combined temperature responses of \( k_{\text{obs}} \) in H2O and D2O buffers are indeed well described by the continuous function, \( \text{Eq } 5 \), with a single set of enthalpic parameters, \( \Delta H(T_b) \) and \( \Delta C_p \), and thus by a single mechanism.

The resulting constant contribution to the enthalpy change, \( \Delta H(T_b) = 130 \text{ kJ} \cdot \text{mol}^{-1} \), of \( \text{Eq } 5 \), is roughly the average of the values that would be obtained from a partition of the experimental data into two linear segments, with separate fits for each segment. The value, \( \Delta C_p = -5.5 \text{ kJ} \cdot \text{mol}^{-1} \cdot \text{K}^{-1} \), corresponds to the uptake of \( \Delta n \sim +150 \) waters. Although \( \Delta n \) is considerably larger than \( \Delta n \) as determined in the osmotic pressure measurements, given that the latter suggest that \( \Delta n \) may change with \( T \) and the simplified form of \( \text{Eq } 4 \), we take the osmotic pressure measurements and temperature dependence of the ET rate constants as being satisfactorily self-consistent.

**DISCUSSION**

In this report, we have examined the temperature variation in the kinetics of oxidation of the Fe protein by the MoFe protein and tested for osmotic pressure effects and sKIE. The reported findings can be summarized as follows. (i) As the temperature is
lowered from 25 to 5°C at a constant pH fixed by MOPS buffer, the observed rate constant for Fe$_{\text{red}}$ → MoFe protein ET decreases strongly, but with a non-linear Arrhenius plot, consistent with previous reports that employed buffers with high temperature coefficients. (ii) ET remains gated at the lowest temperature by a conformational change that involves the binding of numerous water molecules, and cooling does not introduce a solvent kinetic isotope effect ($sKIE$). (iii) These observations contradict the earlier proposals of a change in ET mechanism with cooling. (iv) In support of this mechanistic invariance, it is shown that the nonlinear Arrhenius plot for the ET rate constant arises from the change in heat capacity associated with the binding of waters, and does not indicate a change in mechanism. (v) Finally, it is shown that the Fe protein to MoFe protein extent of ET at constant pH does not change substantially with temperature, in contrast to the changing ET equilibrium inferred previously from experiments where cooling is accompanied by an increase in pH. Overall, these findings address multiple issues associated with inter-component oxidation of Fe$_{\text{red}}$ by MoFe and overturn some long-held views about this process.

Our earlier study of intracomplex ET at ambient temperature employed the addition of viscogen/osmolyte solutes to reveal that the process is gated and responds to osmotic pressure effects, not changes in viscosity. The changes in $k_{\text{obs}}$ with osmolality were interpreted as reflecting the binding of $\Delta n = \sim 50$-$80$ waters during ET, and it was inferred that this is associated with motions at the Fe-MoFe protein interface. We here show that osmotically controlled gating persists to 5°C. The osmolality dependence of $k_{\text{obs}}$ at 5°C indicates that somewhat fewer waters bind upon cooling, $\Delta n = +30$-$50$, a result
that may be attributed to an increased interaction of osmolytes with protein surfaces at the lower temperatures.

The finding that ET remains gated down to the lowest temperature examined, along with the finding that ET is independent of solvent isotopic composition over the entire temperature range, suggests that the ET mechanism does not change upon cooling. Moreover, the observed osmotic pressure dependence provides the foundation for the explanation of the non-linear Arrhenius plots of $k_{\text{obs}}$ in terms of a single, *unchanging* mechanism. The proposed analysis in terms of two linear segments that intersect at a ‘break’ temperature of 15°C that separates two temperature regions in which different mechanisms operated,\(^{16, 17}\) is thermodynamically untenable, while the finding that waters become bound during ET requires that the Arrhenius plot be nonlinear even *without* a change in mechanism. When waters bind during the reaction, the enthalpy of activation, $\Delta H^\dagger$, becomes temperature dependent because the heat capacity is different for free and bound waters (Eq 4). Application of the Gibbs-Helmholtz equation to the ET rate constant then necessarily yields a non-linear Arrhenius plot, Eq 4. As shown in Figure 4-3, the temperature response of $k_{\text{obs}}$ can be well described by a fit to Eq 5, with a plausible value for the number of waters bound during ET. This eliminates the basis of the earlier suggestions for a change in mechanism.

This report further gives an alternative explanation to earlier reports that the extent of ET decreases from ~ 100% at ambient temperature to ~ 50% as the temperature decreases to ~ 5°C. The present experiments carried out with MOPS, a buffer whose pKa changes negligibly with temperature, indicate that at constant pH the extent of ET does *not* change within error upon cooling from 25 to 5 °C, and that this invariance is
independent of solvent isotope (H/D). Thus, our measurements indicate that the extent of ET at constant pH is not temperature dependent.

We do corroborate the finding that the extent of ET does change upon cooling of solutions prepared with buffers with a large temperature coefficient (HEPES buffer in our experiments, TES buffer in previous work). The pH changes by roughly 0.5 units upon cooling these buffer solutions to 5°C, and thus the changes in behavior upon cooling can be assigned not to the temperature variation of ET but to a pH dependence of the extent of ET at 5°C, which has been observed explicitly (SI). A determination of the origin of this dependence is beyond the scope of this report.

Conclusions: The “deficit spending” mechanism for $\text{Fe}^{\text{red}} \rightarrow \text{MoFe}$ protein ET mechanism that operates at ambient temperature does not change upon cooling to 5°C at constant pH. This mechanism likely is associated in some part with motions at the interface between the two proteins of the type schematized in Figure 4-1. The change in heat capacity caused by binding of waters during the ET process leads to a non-linear Arrhenius plot of the rate constant for ET versus inverse temperature, invalidating earlier interpretations of this non-linearity in terms of a change in ET mechanism. The extent of ET observed at ambient temperature does not change during cooling at constant pH, but likely varies with pH at constant, low temperature, where it also depends on solvent isotope. The origin of this last effect, suggestive of an equilibrium solvent isotope effect, remains to be determined.
REFERENCES


(10) Although binding of water at an interface can enhance ET (refs 11 and 12), and thus may contribute to gating, we surmise it is the unknown motions within the MoFe protein that are most important here.


(20) Seefeldt, L. C., Morgan, T. V., Dean, D. R., and Mortenson, L. E. (1992) Mapping the site(s) of MgATP and MgADP interaction with the nitrogenase of
*Azotobacter vinelandii*. Lysine 15 of the iron protein plays a major role in MgATP interaction. *J. Biol. Chem.* 267, 6680-6688.


(25) An abrupt "break" in a plot such as that of Fig 4 would imply the existence of a 2nd order phase change, and could not actually occur in a molecular system.

(26) If the ET rate constant corresponds to that for the conformational gate, then its temperature response is characterized by the activation enthalpy. If ET involves a conformational preequilibrium, then the temperature variation would be described by the sum of the enthalpy change for the gating equilibrium and the ET activation enthalpy. In either case, in a precise application of transition-state theory the ratio, \( [k_{\text{obs}} (T)/k_{\text{obs}} (T_b)] \) would be replaced by \( [T_b(k_{\text{obs}}(T))/T(k_{\text{obs}}(T))] \).
but over the narrow range of temperatures examined, the parameters derived from such a fit differ insignificantly from that to the more intuitive Eq 5.
APPENDIX

Supporting Information

TEMPERATURE INVARIANCE OF THE NITROGENASE ELECTRON TRANSFER MECHANISM
Chart S1 and Fits to Eq 2

Prompted by comments from a reviewer, we incorporate Chart 4-S1 from a Mathcad spreadsheet that demonstrates why fitting many of the traces to Eq 2 is essential, by explicitly comparing a plot of an exponential rise-to-maximum to plots of Eq 2. This graph demonstrates that it is not appropriate to fit the multiphasic traces of Figure 4-2 to a rise-to-maximum.

![Chart 4-S1](image)

The red trace in the Chart corresponds to a simple exponential rise-to-maximum (set $\Delta A_r = 1$) for Fe$^{\text{red}}$ oxidation, rate constant $k_{\text{obs}}$ (consider $k_{\text{obs}} = 1$, or view the x-axis as being $x = k_{\text{obs}}*t$).

The blue trace is a plot of Eq 2 for the same $k_{\text{obs}} = 1$ and $\Delta A_r = 1$, but with a slow followup reaction having $k_{\text{Dr}} = k_{\text{obs}}/5$ to a product with $\Delta A_{\text{Dr}} = 1/10$. The shape of this trace is qualitatively like many in our paper.

The green trace normalizes the blue trace for comparison to the simple exponential rise.

Firstly, one sees that the absorbance maximum of the blue trace is suppressed; a fit of this trace to a rise-to-maximum would underestimate $\Delta A_r$. 
Secondly, comparison of the \textit{red} (exponential rise) and \textit{green} (normalized Eq 2) traces shows that the apparent rise rate in the \textit{green} trace for Eq 2 is greater than the $k_{\text{obs}}$ rise. Thus fitting an experimental trace to a rise-to-maximum would give a rate constant larger than $k_{\text{obs}}$, as well as an absorbance change too small.

Thus, to get the proper absorbance change and rate constant for oxidation of Fe$^{\text{red}}$ from multiphasic SF traces like those of Figure 4-2 one should use Eq 2.

However, as we explicitly state in Materials and Methods, the trends in the resulting parameters obtained by earlier workers as a consequence of changing temperature/osmotic pressure (including ourselves; see below), \textit{are} correctly represented.

We further comment on the absorbance change during the deadtime, incorporated in eq 2 as $\Delta A_o$. Given the relative timescales for the first and second step of the two-step reaction, this absorbance change can indeed be assigned to intracomplex oxidation of Fe$^{\text{red}}$. However, the state formed will convert slowly to C, and thus this term, in principle, has a time dependence (rate constant, $k_{\text{Dk}}$) as well. However, we have explored the consequences of including this phenomenon in the fits, and find that this added complexity causes negligible change in the parameters of importance, $k_{\text{obs}}$ and $\Delta A$. Therefore we have chosen not to incorporate what amounts to a pseudo-sophistication.
Figure 4-S2: Semi-logarithmic plot of the observed rate constants, $k_{obs}$, vs inverse temperature for D$_2$O and H$_2$O when traces are truncated to 0.5s and fit to a rise-to-maximum. Purple solid line is a fit to all data sets, assuming that $k_{obs}$ is the same for H$_2$O and D$_2$O buffers.
Figure 4-S3. Temperature profile of the optical change in the Fe protein in the presence of the MoFe protein. Colors represent different temperatures equally spaced within the range, 25°C-5°C, and overlay within error.
Figure 4-S4. Temperature-response stopped-flow traces for oxidation of Fe\textsuperscript{red} within [Fe\textsuperscript{red}(MgATP)\textsubscript{2};MoFe] complex in HEPES buffer for H\textsubscript{2}O and D\textsubscript{2}O. Arrow in H\textsubscript{2}O plot points in the direction of decreasing temperature, from 25°C to 5°C. D\textsubscript{2}O plot has corresponding color scheme with temperature. Inset: Logarithmic plot of the observed rate constants, $^{l}k_{obs}$ for $L = D$ (D\textsubscript{2}O) and $L = H$ (H\textsubscript{2}O). Black solid line is a fit to both data sets, assuming that there is no difference in $^{l}k_{obs}$ for $L = H$, $D$, and $\Delta n = +136$. 

 ![](image.png)
Figure 4-S5: Temperature dependence of absorbance changes $\Delta A$, (Eq 2) obtained from intracomplex oxidation of Fe$^{\text{red}}$(ATP)$_2$ by MoFe as a function of temperature in HEPES buffer. Blue circles: D$_2$O, red circles: H$_2$O. Blue dashed line is a linear fit to D$_2$O data. Black arrow indicates the direction of increasing pH/pD with temperature in HEPES buffer.
Figure 4-S6: Temperature dependence of absorbance changes $\Delta A$, (Eq 2) obtained from intracomplex oxidation of $\text{Fe}^{\text{red}}\text{(ATP)}_2$ by MoFe as a function of temperature for traces that have been truncated to 0.5s and fit to a rise-to-maximum. Purple dashed line is the average of all values.
Fig 4-S7: pH dependence of ET kinetic traces for reaction carried out in HEPES at 5°C.

pH 6.6 (red) pH 7.0 (blue) and pH 7.4 (Dark red) are shown.
**Fig 4-S8**: pH/pD dependence of absorbance changes $\Delta A$, in HEPES buffer at 5°C. Black arrow indicates the direction of increasing pH/pD.
Figure 4-S9. Plot of the solute-dependent osmotic effect on Nitrogenase. **Black** = 
glycerol, **red** = glucose, **blue** = sucrose. Points and solid lines represent results at 6°C;
dashed lines represent the previously published results at 25°C.
Figure 4-S10: Semi-logarithmic plot of the rate constant vs inverse temperature for the re-reduction of $\text{Fe}^{\text{ox}}$ to $\text{Fe}^{\text{red}}$, $k_{Dr}$ Eq 2, for $\text{D}_2\text{O}$ and $\text{H}_2\text{O}$ buffers. Blue circles and triangles represent $\text{D}_2\text{O}$, while red circles represent $\text{H}_2\text{O}$. Open and closed circles and triangles represent separate experiments.
Figure 4-S11: Temperature dependence of absorbance changes $\Delta A_{Dr}$, (Eq 2) associated with the re-reduction of $Fe^{ox}$ to $Fe^{red}$ as a function of temperature. Purple dashed line is the average of all values.
CHAPTER 5
ELECTRON TRANSFER PRECEDES ATP HYDROLYSIS DURING NITROGENASE CATALYSIS*

ABSTRACT

The biological reduction of N₂ to NH₃ catalyzed by Mo-dependent nitrogenase requires at least eight rounds of a complex cycle of events associated with ATP-driven electron transfer (ET) from the Fe protein to the catalytic MoFe protein, with each ET coupled to the hydrolysis of two ATP molecules. While steps within this cycle have been studied for decades, the nature of the coupling between ATP hydrolysis and ET, in particular the order of ET and ATP hydrolysis, has been elusive. Here, we have measured first-order rate constants for each key step in the reaction sequence, including direct measurement of the ATP hydrolysis rate constant: \( k_{\text{ATP}} = 70 \text{ s}^{-1}, 25^\circ\text{C} \). Comparison of the rate constants establishes the reaction sequence involves four sequential steps: (i) conformationally gated ET \( (k_{\text{ET}} = 140 \text{ s}^{-1}, 25^\circ\text{C}) \), (ii) ATP hydrolysis \( (k_{\text{ATP}} = 70 \text{ s}^{-1}, 25^\circ\text{C}) \), (iii) Pi release \( (k_{\text{Pi}} = 16 \text{ s}^{-1}, 25^\circ\text{C}) \), and (iv) Fe protein dissociation from the MoFe protein \( (k_{\text{diss}} = 6 \text{ s}^{-1}, 25^\circ\text{C}) \). These findings allow the completion of a comprehensive thermodynamic model for the cycle undergone by the Fe protein, revealing that it is not the energy of ATP hydrolysis, but rather the energy of ATP binding and protein-protein formation that drives ET.

INTRODUCTION

Biological nitrogen fixation catalyzed by the Mo-dependent nitrogenase has a limiting reaction stoichiometry shown in eq 1 (1, 2):

\[
N_2 + 8e^- + 16\text{MgATP} + 8\text{H}^+ \rightarrow 2\text{NH}_3 + \text{H}_2 + 16\text{MgADP} + 16\text{Pi} \quad (\text{eq 1})
\]

The ATP-driven reduction of one \(N_2\) with evolution of one \(H_2\) requires a minimum of \(8 e^-\) and the hydrolysis of 16 ATP molecules in a complex cascade of events in which electron transfer (ET) from the nitrogenase Fe protein to the catalytic MoFe protein is coupled to the hydrolysis of two ATP molecules (1, 3, 4). The Fe protein is a homodimer with a single [4Fe-4S] cluster and two nucleotide binding sites, one in each subunit (5). The MoFe protein is an \(\alpha_2\beta_2\) tetramer, with each \(\alpha\beta\) pair functioning as a catalytic unit that binds an Fe protein (6). Each \(\alpha\beta\) unit contains an [8Fe-7S] cluster (abbreviated P-cluster) and a [7Fe-9S-Mo-C-R-homocitrate] cluster (abbreviated FeMo-cofactor or M-cluster) (6–10). In each ET event, the Fe protein, in the reduced (1+) state with two bound ATP, first associates with the MoFe protein (Figure 5-1). In a recent model, termed “deficit-spending,” it is proposed that this association triggers a two-step ET event (11, 12). The first ET step occurs inside the MoFe protein, involving ET from the P cluster resting state (\(P^N\)) to the resting FeMo-cofactor (\(M^N\)) cluster, resulting in an oxidized P cluster (\(P^{1+}\)) and a reduced FeMo-cofactor (\(M^R\)) (12). This ET event is conformationally gated (11) with an apparent first order rate constant (\(k_{\text{ET}}\)) between 100-140 s\(^{-1}\) (11, 12). In the second ET step, an electron is transferred from the Fe protein [4Fe-4S] cluster to the oxidized \(P^{1+}\) cluster, resulting in the return of the P cluster to the resting oxidation state (\(P^N\)) and an oxidized [4Fe-4S]\(^{2+}\) cluster in the Fe protein (12). This second step is fast, having a rate constant greater than 1700 s\(^{-1}\) (12).
Figure 5-1: Order of events in nitrogenase complex. (A) The Fe protein subunits are shown as two blue ovals (left) with an ATP bound in each subunit and the [4Fe-4S] cluster (green cubane). The MoFe protein α subunit (orange) and beta subunit (green) are shown on the right, with the P^N cluster shown as a gray box and the FeMo-cofactor (M^N) shown as a gray diamond. (B) From left to right, order of events in the nitrogenase electron transfer is shown with rate constants (s^{-1}) displayed where known.

Transfer of one electron from the Fe protein to an αβ-unit of MoFe protein is known to be coupled to the hydrolysis of the two ATP molecules bound to the Fe protein, yielding two ADP and two Pi (2). Following the hydrolysis reaction, the two Pi are released from the protein complex with a first order rate constant (k_{Pi}) of 22 s^{-1} at 23^\circ C (13). The last event in the cycle is the release of the oxidized Fe protein with two ADP bound, [Fe^{ox}(ADP)_2] from the MoFe protein with a rate constant (k_{diss}) of ~ 6 s^{-1}, the rate-limiting step in catalysis at high electron flux (14). After dissociation from the MoFe
protein, the \([\text{Fe}^{\text{ox}}(\text{ADP})_2]\) protein is prepared for a second round of electron delivery by one-electron reduction to \([\text{Fe}^{\text{red}}(\text{ADP})_2]\) and replacement of the two MgADP by MgATP. This cycle is repeated until enough electrons are transferred to the MoFe protein to achieve substrate reduction (15).

Although the energetic coupling between ET and ATP hydrolysis is firmly established (1, 3, 4, 16), the nature of this coupling has remained unresolved: does ATP hydrolysis itself provide the principal energy input for the conformational change(s) that drive ET from Fe protein to the MoFe protein, or, does the bound ATP induce the formation of a reactive, “activated” conformation of the complex, with ET being driven by the free energy of ATP-activated protein-protein binding. These alternatives are characterized by different orders of ET and ATP hydrolysis, but the order has never been established. Some studies have indicated that ATP hydrolysis occurs after ET (13, 17, 18), while other studies have suggested just the opposite; namely that ATP hydrolysis occurs before ET (15, 16, 19, 20). One of the reasons for this lack of clarity in the order of these key events is the absence of direct measurement of ATP hydrolysis rates by nitrogenase within a single catalytic cycle. The rate constant for Pi release during one cycle has been measured, thereby establishing a lower limit on the rate constant for ATP hydrolysis (13). However, the rate constant for ATP hydrolysis could be much faster than Pi release, and could be faster than the rate constant for ET.

Here, we have directly measured the rate constant for ATP hydrolysis for a single nitrogenase turnover cycle, as well as measuring the rate constants for each of the other key steps under the same conditions, thereby allowing an unequivocal assignment of the
order of events in a single electron-transfer cycle. Establishing the order of events allows a full thermodynamic Fe protein cycle to be constructed.

RESULTS

ATP hydrolysis and Pi release. The hydrolysis of the 2 ATP bound in the Fe protein to 2 ADP + 2 Pi is initiated during the transient association of the Fe protein with the MoFe protein; ATP hydrolysis is not catalyzed by the Fe protein alone (1). Pre-steady state rates of ATP hydrolysis during a single Fe-MoFe protein association event have never been reported, although steady-state rates have been reported in several earlier studies, with typical rates ranging from 3600-4500 nmols.min⁻¹.(mg MoFe protein)⁻¹ (21). These prior steady-state measurements relied on quantification of released ADP or Pi over multiple turnovers, and thus provide little insight into the rate of ATP hydrolysis during the first turnover (21).

Establishing a pre-steady state rate for ATP hydrolysis is challenging, requiring quantification of ATP consumed and ADP formed on a millisecond time scale. To make this measurement, we employed a rapid chemical-quench approach to stop ATP hydrolysis at selected times after initiating the reaction, with an instrument dead-time of 4 ms and time resolution of 1 ms. The ATP and ADP in each quenched solution were quantified by utilizing ³²P-ATP (labeled on the α-phosphate) as a tracer. For each sample, ATP was separated from ADP by thin-layer chromatography, with quantification of each nucleotide accomplished by counting of the ³²P. Using this approach, it was possible to establish the precise concentration of ADP formed as a function of time after
Figure 5-2: Kinetics of key nitrogenase catalytic steps. (A) Electron transfer is monitored as the change in absorbance at 430 nm as a function of time (open circles). The data were fit to the sequential kinetic model (eqn 2) (line), with a calculated $k_{ET}$ of 140 s$^{-1}$. (B) ATP hydrolysis is monitored as the formation of $\alpha$-$^{32}$P-ADP as a function of time (closed circles). The data were fit to eqn 2 (line), with a calculated $k_{ATP}$ of 70 s$^{-1}$. (C) Phosphate (Pi) release is monitored by the increase in fluorescence from Pi binding to MDCC-labeled phosphate binding protein as a function of time (open square). The data were fit to eqn 2 (line), with a calculated $k_{Pi}$ of 16 s$^{-1}$. (D) Dissociation of Fe from the MoFe protein is monitored by the decrease in absorbance at 430 nm as a function of time (open triangle). The data were fit to a single exponential equation (solid line) with a first order rate constant ($k_{diss}$) of 6 s$^{-1}$. All four experiments were carried out under the same experimental conditions at 25 °C.
initiating the reaction by rapid-mixing (Figure 5-2B). Over the time range examined (up to 200 ms), the quantity of ADP formed was observed to rise rapidly to a plateau. These data were initially fit to an exponential rise to maximum to get an estimated rate constant, that was found to be noticeably less than \( k_{ET} \), and then the ATP hydrolysis data were fit to the kinetic model described in Materials and Methods (eq 2), yielding a rate constant for ATP hydrolysis, \( k_{ATP} = 70 \pm 7 \text{ s}^{-1} \) at 25°C.

The rate constant for ATP hydrolysis determined here can be compared to the rate constant for Pi release. In earlier studies, pre-steady state Pi release was found to have a rate constant \( (k_{pi}) \) of \( \sim 22 \text{ s}^{-1} \), roughly three-fold less than \( k_{ATP} \) (13). The conditions used in the earlier studies were different from the conditions used in the ATP hydrolysis studies reported here, so the Pi release experiment was conducted under the same conditions used for the measurement of ATP hydrolysis. Pi release was detected by binding of the free Pi to a phosphate-binding protein labeled with a fluorescent reporter (MDCC-PBP) that shows a 7-fold increase in fluorescence upon Pi binding, thereby allowing real-time monitoring of Pi release using a stopped-flow fluorometer (Figure 5-2C) (22). The concentration of Pi released as a function of time was fit to the kinetic model described in Materials and Methods (eq 2), yielding a rate constant, \( k_{pi} = 16 \pm 1 \text{ s}^{-1} \), which is consistent with earlier reports. Figure 5-2C shows that the kinetic model fits the Pi release data up to almost 150 ms quite satisfactorily, but it was also recognized that at later times the Pi release was more complex due to the achievement of a steady state. As indicated by the result, \( k_{pi} < k_{ATP} \), and as visualized when the progress curves for ATP hydrolysis and Pi release are plotted together on a logarithmic time plot (Figure 5-3), Pi
is not released promptly upon ATP hydrolysis, but rather occurs as a subsequent kinetic step with a detectable delay after ATP hydrolysis.

**Figure 5-3: Time course for key steps in the nitrogenase cycle.** The data for ET (○), ATP hydrolysis (●), and Pi release (□) are plotted as a function of the logarithm of time. Each data set was fit to the model in eqn 2 (solid lines), with the calculated rate constants noted. The protein-protein dissociation is represented by a simulation (dashed green line) generated using the kinetic model (eqn 2), with rate constants fixed at $k_{ET} = 140 \text{ s}^{-1}$, $k_{ATP} = 70 \text{ s}^{-1}$, and $k_{Pi} = 16 \text{ s}^{-1}$.

**ATP hydrolysis compared to electron transfer.** The rate constant for ET from the Fe protein to the MoFe protein ($k_{ET}$) has been reported to be between 100-150 s$^{-1}$ (23), depending on reaction conditions and the proteins used. This rate constant was re-examined in the present study under identical conditions to the Pi release and ATP hydrolysis measurements. ET is monitored by the absorbance increase that accompanies
oxidation of the Fe protein [4Fe-4S] cluster after initiation of the reaction (Figure 5-2A). Fit of these data to the kinetic model (eq 2) yields an apparent first order rate constant of \( k_{\text{ET}} = 140 \pm 10 \text{ s}^{-1} \). This rate constant reflects the oxidation of the Fe protein as an electron is delivered to the MoFe protein, a process recently shown to be gated by protein conformational changes (11). In the deficit-spending model of ET, the observed ET is the last ET event, following conformational gating and ET from the P cluster to FeMo-cofactor (12). Thus, the observed first order rate constant of 140 s\(^{-1}\) reflects both protein conformational changes and all ET events.

The rate constant for ATP hydrolysis of \( k_{\text{ATP}} = 70 \text{ s}^{-1} \) is half that of the measured ET rate constant, clearly placing ATP hydrolysis as a distinct kinetic step that follows ET. The order of events is visualized by comparing ET, ATP hydrolysis, and Pi release data on a logarithmic time plot (Figure 5-3): ET occurs first, followed by ATP hydrolysis, followed by Pi release.

*Protein-protein dissociation.* Earlier studies have shown that Fe protein dissociation from the MoFe protein is the last event in the cycle, with a first order rate constant \( (k_{\text{diss}}) \) ranging from 5 to 10 s\(^{-1}\) (14). We directly determined this dissociation rate constant, as described in Materials and Methods, under the same conditions as the other measurements (Figure 5-2D), finding a first order rate constant of \( k_{\text{diss}} = 6 \pm 2 \text{ s}^{-1} \) from a fit to an exponential of the absorbance change associated with Fe(ADP)\(_2\) reduction subsequent to mixing of [Fe\(^{ox}\)(ADP)\(_2\);MoFe] upon mixing. This value is consistent with protein-protein dissociation being a discrete kinetic step and the last event during the catalytic cycle. The dissociation event is simulated with the kinetic model (eq 2), by fixing the ET, ATP hydrolysis, Pi release, and dissociation rate constants. The simulation (presented as a rise to maximum) clearly places the dissociation as the last event in the series, in a kinetic step distinct from Pi release (Figure 5-3).
DISCUSSION

Model for Events in the Nitrogenase Cycle. The measurement of the pre-steady state rate constant for ATP hydrolysis reported here defines the order of the key events involved in electron transfer within the Fe protein-MoFe protein complex. As is evident in the kinetic progress curves shown in Figure 5-3, the order of events is established as: (i) ET, (ii) ATP hydrolysis, (iii) Pi release, followed by (iv) dissociation of the Fe protein from the MoFe protein.

The findings of the current study can be combined with earlier findings to construct a model of the key events that occur during one round of Fe protein-MoFe protein association (Figure 5-4). The process begins with the formation of a complex between Fe\textsuperscript{red}(ATP)\textsubscript{2} and an αβ catalytic unit of the MoFe protein (Figure 5-4, upper left). This association is known to be fast, with a second order rate constant ($k_{assoc}$) of $5 \times 10^7$ M$^{-1}$ s$^{-1}$ (24). At the concentration of Fe protein used in the present study, the association event is much faster than subsequent events, so all the measurements described here correspond to first-order, intra-complex steps.

Recent studies on the effects of osmotic pressure on the rate of ET have established that following protein-protein association, large scale (~ 800 Å$^2$ changes in buried surface) conformational changes within the complex gate subsequent ET events (11). It has been suggested that these conformational changes result in activation of the P cluster to a state (P*) that transfers an electron to FeMo-cofactor, forming the oxidized P cluster (P$^{1+}$). In this model, subsequent ET from Fe\textsuperscript{red}(ATP)$\textsubscript{2}$ to the oxidized P cluster is fast and not gated by protein conformational changes, and generates the final product of intra-complex ET, Fe\textsuperscript{ox}(ATP)$\textsubscript{2}$ bound to MoFe protein containing a resting P cluster and
Figure 5-4: Model of nitrogenase order of events. Shown are the Fe protein (blue ovals) with the [4Fe-4S] cluster in the +1 (green cube) or +2 (yellow cube) oxidation state. The MoFe protein (α subunit orange and β subunit green) with the P cluster (rectangle, grey $P^N$, red activated $P^{N*}$ and yellow $P^{+1}$ states) and the M cluster (diamond, grey $M^N$ and maroon $M^R$ state). The reaction sequence starts at the top left with the association of the Fe protein with the MoFe protein and ends at top right with the dissociation of the oxidized Fe protein from the reduced MoFe protein. First order rate constant (k, s$^{-1}$) values for 25°C are shown. Conformational changes in the Fe protein and MoFe protein are denoted by changes in shape of the ovals or rectangles.
one-electron reduced FeMo-cofactor \( (M^R) \). The overall rate constant for the multiple steps in the ET process is \( 140 \text{ s}^{-1} \) (12).

As established here, hydrolysis of the two ATP molecules to ADP and Pi occurs as a distinct kinetic step that follows this ET process. Data shown in Figure 5-3 further establishes that Pi is not released from the protein-protein complex immediately upon ATP hydrolysis, but Pi is released in a distinct, subsequent step. It seems reasonable to conclude that upon ATP hydrolysis, the complex undergoes conformational changes, although the extent of such changes has not been established. Likewise, it seems reasonable that the release of Pi is accompanied by additional protein conformational changes, but again such changes have not been measured directly.

The final step in the ET process is the release of the \( \text{Fe}^{\text{ox}}(\text{ADP})_2 \) from the one-electron reduced MoFe protein (Figure 5-4) in a process that is not synchronous with Pi release, but follows as a discrete step with a rate constant of \( 6 \text{ s}^{-1} \), as has been established in earlier studies (1, 13, 14). The MoFe protein that contains a one electron reduced FeMo-cofactor is then free to be reduced additional times by repeating the process presented in Figure 5-4, beginning with the binding of a second \( \text{Fe}^{\text{red}}(\text{ATP})_2 \) (15). The released \( \text{Fe}^{\text{ox}}(\text{ADP})_2 \) is ready to be returned to the \( \text{Fe}^{\text{red}}(\text{ADP})_2 \) state by reduction of its \( [4\text{Fe-4S}]^{2+} \) cluster reduced back to the +1 oxidation state, and replacement of ADP by ATP.

*The Fe-protein Cycle.* By revealing that intra-complex \( \text{Fe}^{\text{red}}(\text{ATP})_2 \rightarrow \text{MoFe} \) protein ET is followed by ATP hydrolysis (Figure 5-4), this work also completes the overall ‘Fe-protein cycle’ first formulated by Lowe and Thorneley (LT) to summarize the steps undergone by the Fe protein as it delivers an electron to the MoFe protein (1, 17).
Figure 5-5 presents the key Fe-protein reactions formulated as a thermodynamic cycle. The cycle begins with free Fe$^{ox}$ (ADP)$_2$ (upper right), the final form of Fe protein at the end of a previous ET cycle, and ends with a return to this state upon completion of the cycle. Figure 5-5 is further annotated to emphasize the function of the Fe protein as a ‘nucleotide switch’ (25, 26) that uses ATP binding/hydrolysis to ‘transport’ an electron from an electron source/reductant to the MoFe protein. Viewed in this way, Figure 5-5 introduces a correspondence of the thermodynamic cycle for ATP-dependent ‘electron transport’ by the Fe protein with the thermodynamic cycle for ATP-dependent substrate export by ABC transporters (27–30).

**Figure 5-5: Thermodynamic Fe-protein cycle.** Drawn with a focus on Fe protein electron transport to MoFe protein, it can be viewed as starting at upper right. MoFe$^{red}$ represents reduced MoFe protein. Dashed arrows and notation that surround the cycle refer to analogy with ABC exporter thermodynamic cycle (see text).
The Fe-protein cycle of Figure 5-5 (top, right to left) begins with the reduction of Fe$^{\text{ox}}$(ADP)$_2$ to Fe$^{\text{red}}$(ADP)$_2$, which triggers rapid replacement of the two ADP by two ATP (1, 17). The structure of the Fe protein undergoes major changes upon ATP binding, leading to numerous changes in its properties, but historically interest has focused on an $\sim -120$ mV change in the reduction potential of the [4Fe-4S] cluster of nucleotide-free Fe protein upon ATP binding, and the possibility that this change in potential is central to initiating the transfer of an electron to MoFe protein. Although the more negative midpoint potential would favor ET (1), for present purposes it is important to note (Figure 5-5) that the second step in the thermodynamic Fe protein-cycle is not ATP binding to free Fe protein, but replacement of bound ADP by ATP. Namely, the free Fe$^{\text{red}}$ does not appear in the thermodynamic cycle regardless of the mechanistic details of the nucleotide interchanged. In fact, the reduction potentials of Fe$^{\text{ox}}$(ADP)$_2$ to Fe$^{\text{red}}$(ADP)$_2$ are the same (31). As a result, ATP/ADP exchange is iso-potential, and so the potential lowering upon nucleotide binding cannot contribute to the thermodynamic coupling between ET and ATP binding and hydrolysis.

The MoFe protein catalytic unit next binds to Fe$^{\text{red}}$(ATP)$_2$ to form the ET-active [Fe$^{\text{red}}$(ATP)$_2$, MoFe] complex (Figure 5-5, left, down). Structural differences between Fe$^{\text{red}}$(ADP)$_2$ and Fe$^{\text{red}}$(ATP)$_2$ must have thermodynamic consequences for the Fe cycle as the free energies for binding between the MoFe protein and Fe protein alone or Fe protein with bound ADP or ATP are different (1). The excess free energy for binding MoFe protein to Fe$^{\text{red}}$(ATP)$_2$ relative to Fe$^{\text{red}}$(ADP)$_2$ contributes to the thermodynamic coupling between ET and ATP binding/hydrolysis.
The \([\text{Fe}^{\text{red}}(\text{ATP})_2, \text{MoFe}]\) complex undergoes inter-protein “transport” of the electron, ATP hydrolysis, Pi release, and dissociation. In the absence of a known ordering of these steps, two alternative proposals have been considered to explain the coupling of ET to ATP binding and hydrolysis. In one, ATP hydrolysis itself provides the principal energy input for the conformational change(s) that drive \(\text{Fe}^{\text{red}} \rightarrow \text{MoFe}\) protein ET; in the other, the bound ATP induces the formation of a reactive, “activated” conformation of the complex, and it is the free energy of ATP-activated protein-protein binding that drives ET. The present measurements of the rate constants for all of the key steps discriminates between these two alternatives, and in doing so sheds further light on the energy transduction through ATP hydrolysis: (i) That the ATP hydrolysis is temporally decoupled from and follows ET (Figure 5-3) shows that ET is not driven by the free energy of ATP hydrolysis, and that it must be the ATP-dependent free energy of protein-protein binding that drives ET. (ii) That ATP hydrolysis and Pi release occur as temporally separated steps, both of which precede the dissociation of the product \([\text{Fe}^{\text{red}}\text{ATP}_2, \text{MoFe}^{\text{red}}]\) complex (Figure 5-4) (13), demonstrates that both hydrolysis and Pi release are required to completely relax the conformationally activated complex, thereby inducing dissociation of the Fe protein from the reduced MoFe\(^{\text{red}}\) and completing the cycle.

Viewed in this way, the Fe protein thermodynamic cycle is analogous to the four-step thermodynamic cycle of the ABC (ATP-binding cassette) trans-membrane substrate transporters, in particular the exporter sub-class (27–30). These transporters catalyze substrate transport by a process dependent on conformational changes driven by ATP binding, with the system being reset for another cycle by ATP hydrolysis. The ABC
trans(ex)porter cycle begins with binding of the substrate to the protein, which triggers ATP binding (Step I); in nitrogenase the analogous step is ‘electron binding’ by Fe$^{ox}$$(ADP)_2$ protein, namely reduction to Fe$^{red}$(ADP)$_2$, followed by exchange of ADP by ATP. In the transporters, ATP binding free energy drives a sequence of conformational changes that results in an activated state that undergoes trans-membrane substrate transport (Step II); in nitrogenase, ATP binding free energy drives the formation of an activated [Fe$^{red}$(ATP)$_2$, MoFe] complex that undergoes deficit-spending Fe$^{red}$ $\rightarrow$ MoFe electron transport. In the transporters, ATP hydrolysis to ADP and subsequent Pi release (Step III) elicit conformational relaxation (Step IV), thereby ‘resetting’ the transporter for another catalytic cycle; in nitrogenase, ATP hydrolysis and subsequent Pi release relax the complex, resulting in dissociation of the reduced MoFe protein, leaving the Fe$^{ox}$(ADP)$_2$ protein ready for another Fe protein cycle. Thus, the concept of Fe protein as a “nucleotide switch” must be expanded to include the idea that the Fe protein-MoFe protein complex toggles between two conformational states – “activated” and “relaxed”-- in response to changes in nucleotide state, thereby driving nucleotide-dependent “electron transport”.

*Nucleotide-Dependent Structural Changes.* How do nucleotide-dependent changes in the conformation of the [Fe; MoFe] complex drive ET? The X-ray structures of Fe-MoFe protein complexes with a suite of nucleotides (32), including nucleotide-free, ADP-bound, the ATP analogue β-γ-methylene ATP (AMPPCP)-bound, and the ATP-hydrolysis transition state represented by ADP-AlF$_4$-bound complex, have suggested progressive changes in the structure of Fe protein accompanied by rearrangement at the protein-protein interface as a function of nucleotide state. We interpreted the osmotic-
pressure dependence of the observed $\text{Fe}^{\text{red}} \rightarrow \text{MoFe}$ ET process as reflecting motions of the Fe protein relative to MoFe (11), consistent with the crystallographic demonstration that changes in nucleotide state drive large changes in the buried protein-protein interface area (32). However, it must be that activation for ET includes conformational changes within the MoFe protein that have yet to be observed in crystal structures.

Neither the structure of the complex that binds the ATP analogue, AMPPCP, nor that of the complex that binds the analogue for the transition state of ATP hydrolysis, ADP-$\text{AlF}_4^-$, shows any significant perturbations within the MoFe protein relative to isolated MoFe protein. However, the ABC-transporter-like mechanism encapsulated in the cycle of Figure 5-5 offers a possible resolution to this puzzle. As ATP hydrolysis/Pi release occurs only after ET, and thus ‘only’ contribute to relaxation of the ‘activated’ conformational state associated with the ATP complex, there is no requirement that the transition state for ATP hydrolysis be associated with a high-energy state within the MoFe protein.

Two further observations can be considered regarding the absence of changes within the MoFe protein in the structure of the AMPPCP-bound complex: (i) AMPPCP in fact is not a truly faithful ATP analogue in the nitrogenase system, as shown by the inability to promote proper ET and (ii) as the deficit-spending ET process is conformationally gated, it may well be that the anticipated state in which FeMo-cofactor and its environs are activated for substrate reduction are only formed by a conformational fluctuation and never exists in high occupancy for trapping and direct characterization.

Summary. By establishing the pre-steady state rate constant for ATP hydrolysis in nitrogenase catalysis, the order of sequential events in the nitrogenase cycle is finally
established to be: (i) conformationally gated ET, (ii) ATP hydrolysis, (iii) Pi release, and finally (iv) Fe protein dissociation from the MoFe protein. Establishing this order of events provide insights into the functions of ATP in the nitrogenase mechanism, revealing that it is the free energy of ATP binding and protein-protein association that control the ET events, rather than the free energy of ATP hydrolysis.

MATERIALS AND METHODS

Materials, Protein Purification and Activity Assays. All reagents, unless stated otherwise, were purchased from Sigma Aldrich Chemicals (St. Louis, MO). Nitrogenase proteins were expressed in Azotobacter vinelandii strain DJ995 (wild-type MoFe with His tag), and DJ884 (wild-type Fe protein) as described previously (33). The MoFe protein contained a seven-histidine tag near the carboxyl terminus of the α-subunit allowing purification using the previously described metal affinity chromatography method (33). Fe protein was purified using ion exchange and size-exclusion liquid chromatography (33, 34). Both proteins were greater than 95% pure based on sodium dodecyl sulfate-polyacrylamide gel electrophoresis separation followed by Coomassie blue staining. Manipulation of proteins was done in septum-sealed serum vials under argon atmosphere. All transfers of gases and liquids were done using gastight syringes.

Stopped-flow Spectrophotometry and the Oxidation of Fe Protein. Stopped-flow spectrophotometry was performed at 25°C with a Hi-Tech SF61 stopped-flow (SF) UV-visible spectrophotometer equipped with data acquisition software (Salisbury, Witts UK). The SF unit of the spectrophotometer was housed inside a nitrogen-filled glove box (< 5 ppm O₂). The temperature during the experiments was controlled by a circulating water bath housed outside the glove box (18). In these experiments, 75 μM Fe and 20 μM
MoFe protein were contained in one drive syringe while the other syringe contained 18 mM MgCl₂ and 10 mM ATP. Both syringes contained buffer [100 mM HEPES (pH 7.4) and 10 mM sodium dithionite]. Primary electron transfer from the Fe protein to the MoFe protein was monitored by an increase in absorbance at 430 nm that occurs as the [4Fe-4S] cluster of the Fe protein becomes oxidized during turnover within the [Fe₅⁺(ATP)²;MoFe] complex.

The data were fit using KinTech Explorer (Kintek Corp., Austin, TX) to a sequential A → B → C → D → E kinetic model (eqn 2).

\[
\begin{align*}
k_{ET} & \quad k_{ATP} & \quad k_{Pi} \\
[Fe^{\text{red}}(ATP)_2;MoFe] & \rightarrow [Fe^{\text{ox}}(ATP)_2;MoFe^{\text{red}}] & \rightarrow [Fe^{\text{ox}}(ADP,Pi)_2;MoFe^{\text{red}}] & \rightarrow \\
[Fe^{\text{ox}}(ADP)_2;MoFe^{\text{red}}] + Pi & \rightarrow [Fe^{\text{ox}}(ADP)_2] + [MoFe^{\text{red}}]
\end{align*}
\] (eqn 2)

In this model, \(k_{ET}, k_{ATP}, k_{Pi}\) and \(k_{diss}\) are the rates constants for ET, ATP hydrolysis, phosphate release, and [Fe₅⁺(ADP)²;MoFe] complex dissociation, respectively. All steps of the ET process can be taken as irreversible except for ATP hydrolysis, and here the rate constant for the reverse reaction is so low that the reverse process can be ignored in the present experiments (1, 17). As a result the sequential model of eqn 2 is appropriate (in the fitting process all reverse rate constants are fixed to a value of zero). To fit the ET data, the other rate constants were fixed at \(k_{ATP} = 70 \text{s}^{-1}, k_{Pi} = 16 \text{s}^{-1}, k_{diss} = 6 \text{s}^{-1}\).

**Quench Flow Studies for ATP Hydrolysis.** Pre-steady ATP hydrolysis assays were performed at 25°C on a rapid chemical quench-flow instrument (KinTek Corp, Austin, TX) housed in a nitrogen-filled glove box (< 5 ppm O₂). A 18 µL volume of 10 µM MoFe and 20 µM Fe (syringe A) was mixed with a 18 µL volume of 1 mM ATP with [α-
$^{32}$P]ATP (1.5 µCi) from syringe B, with varying times of reaction. Reactions were rapidly quenched with 45 µL of 0.5 M EDTA added from syringe C. Aliquots (0.9 µL) of the quenched reaction were spotted onto a thin layer chromatography (TLC) plate and developed in 0.6 M potassium phosphate buffer, pH 3.4 for 45 minutes. The [$\alpha^{32}$P]ATP and the [$\alpha^{32}$P]ADP were detected with a Storm PhosphorImager (Molecular dynamics, Sunnyvale, CA) and quantified using the ImageQuant software (Molecular Dynamics, Sunnyvale, CA). The data were fit to the kinetic model (eqn 2), with fixed values of $k_{ET} = 140 \text{ s}^{-1}$, $k_{Pi} = 16 \text{ s}^{-1}$, $k_{diss} = 6 \text{ s}^{-1}$.

**Kinetics of Inorganic Phosphate Release.** The time-course of phosphate release was determined in a stopped-flow (Auto SF-120, Kintek Corp, Austin, TX) using the coumarin (N-[2-(1-maleimidyl)ethyl]-7-(diethylamino) coumarin-3-carboxamide) labeled phosphate binding protein assay (MDCC-PBP) (22). Briefly, Pi binding to MDCC-PBP results in an increase in fluorescence (13, 22) ($\lambda_{\text{exitation}} = 430 \text{ nm}$, $\lambda_{\text{emission}} > 450 \text{ nm}$). The experiments were carried out at 25°C in SF-buffer (0.5 mM sodium dithionite and 25 mM HEPES; pH 7.4). Prior to each experiment, the stopped-flow syringes and flow lines were treated with a Pi-MOP (SF-buffer with 300 µM 7-methylguanine (7-meG), and 0.2 units/mL purine nucleoside phosphorylase (PNPase) for 15 min to remove contaminating Pi (22) and then rinsed with buffer. Two µM MoFe and 6 µM Fe was rapidly mixed with a solution of 10 µM PBP-MDCC, 20 mM MgCl$_2$ and 2mM ATP and the change in fluorescence was monitored over time. A control time course, conducted without nitrogenase, was used to correct for the presence of contaminating phosphate. PBP-MDCC fluorescence enhancement was converted to [Pi] after calibration in the stopped-
flow using [NaH₂PO₄] standards as described (22). The Pi release data were fit to the sequential kinetic model (eqn 2) with $k_{ET} = 140 \text{ s}^{-1}$, $k_{ATP} = 70 \text{ s}^{-1}$, and $k_{diss} = 6 \text{ s}^{-1}$ fixed.

The Fe Protein-MoFe Protein Dissociation Rate Constant. The dissociation of the Fe protein from the MoFe protein was determined by following the release of the Fe protein from the MoFe protein in the stopped flow spectrophotometer. Fe$^{ox}$(ADP)$_2$ is not reduced by dithionite within the [Fe$^{ox}$(ADP)$_2$;MoFe] complex. Thus, the rate constant for the dissociation of the [Fe$^{ox}$(ADP)$_2$;MoFe] complex was determined by measuring the decrease in absorbance at 430 nm during the reduction of Fe$^{ox}$ to Fe$^{red}$ by dithionite after Fe$^{ox}$(ADP)$_2$ dissociates from the MoFe protein (14). Rate constants for dissociation were determined by fitting the data to a single-exponential equation (eqn 3)

$$A = e^{-k_{diss}t}$$

where $A$ is the amplitude of the absorbance change and $k_{diss}$ is the rate constant for dissociation of the complex. Sodium dithionite was removed from the as-isolated MoFe and Fe proteins and exchanged into a 50 mM HEPES buffer with 200 mM NaCl, pH 7.4, by passage over a Sephadex G-25 column. Oxidized Fe protein was generated by adding increments of a 25 mM IDS solution to the Fe protein until a blue color remained. The excess IDS was removed by passing the sample over a Dowex ion-exchange resin and Sephadex G-25 column equilibrated with 50 mM HEPES buffer, pH 7.4. For each experiment, syringe A contained 40 µM Fe$^{ox}$ protein, 40 µM MoFe protein, 5 mM MgADP in a 50 mM HEPES buffer, pH 7.4, and syringe B contained 200 µM Fe$^{red}$ protein, and 20 mM sodium dithionite in 50 mM HEPES buffer, pH 7.4.
REFERENCES


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CHAPTER 6
UNCOUPLING NITROGENASE: CATALYTIC REDUCTION OF HYDRAZINE TO AMMONIA BY A MoFe PROTEIN IN THE ABSENCE OF Fe PROTEIN-ATP

ABSTRACT

The catalytic reduction of hydrazine (N$_2$H$_4$) to ammonia by a β-98$^{\text{Tyr\rightarrow His}}$ MoFe protein in the absence of the Fe protein or ATP is reported. The reduction of N$_2$ or other substrates (e.g., hydrazine, protons, acetylene) by nitrogenase normally requires the transient association of the two nitrogenase component proteins, the Fe protein and the MoFe protein. The Fe protein, with two bound MgATP molecules, transfers one electron to the MoFe protein during each association, coupled to the hydrolysis of two MgATP. All substrate reduction reactions catalyzed by nitrogenase require delivery of electrons by the Fe protein coupled to the hydrolysis of MgATP. We report that when a single amino acid within the MoFe protein (β-98$^{\text{Tyr}}$) is substituted by His, the resulting MoFe protein supports catalytic reduction of the nitrogenous substrate hydrazine (N$_2$H$_4$) to two ammonia molecules when provided with a low potential reductant, polyaminocarboxylate ligated Eu$^{\text{III}}$ ($E_m$ -1.2 V vs. NHE). The wild-type and a number of other MoFe proteins with amino acid substitutions do not show significant rates of hydrazine reduction under these conditions, whereas the β-98$^{\text{His}}$ MoFe protein catalyzes hydrazine reduction at rates up to 170 nmol NH$_3$/min/mg MoFe protein. This rate of hydrazine reduction is 94% of the rate catalyzed by the β-98$^{\text{His}}$ or wild-type MoFe protein when combined with the Fe protein, ATP, and reductant under comparable conditions. The β-98$^{\text{His}}$ MoFe protein
reduction of hydrazine in the absence of the Fe protein showed saturation kinetics for the concentration of reductant and substrate. The implications of these results in understanding the nitrogenase mechanism are discussed.

Reduction of $N_2$ to ammonia by the Mo-dependent nitrogenase requires the action of two component proteins called the Fe protein and the MoFe protein.\(^1\) The MoFe protein contains the multimetallic FeMo-cofactor catalytic cluster $[7\text{Fe}-9\text{S}-\text{Mo}-X-R-$homocitrate] at which substrates bind, and the auxiliary P-cluster $[8\text{Fe}-7\text{S}]$ that is proposed to mediate electron transfer from the Fe protein (Figure 1).\(^2\) The Fe protein contains a $[4\text{Fe}-4\text{S}]$ cluster and two MgATP binding sites.\(^3\) During catalysis by nitrogenase, the Fe protein transiently associates with the MoFe protein and a single electron is transferred from the Fe protein to the MoFe protein in conjunction with the hydrolysis of two MgATP molecules.\(^4\) The Fe protein-2MgADP then dissociates from the

Figure 6-1: MoFe protein metal clusters and key amino acid residues are shown. The Fe protein association site is to the left outside of the view. Colors are Mo in magenta, Fe in rust, S in yellow, C in gray, O in red, and N in blue. Based on the protein data base file 1M1N.pdb.\(^2\)
MoFe protein, allowing another reduced Fe protein-2MgATP to bind to the MoFe protein. This cycle is repeated to accumulate electrons in the MoFe protein to support substrate reduction. The Fe protein with bound ATP is the only known reductant of the MoFe protein that supports reduction of N\textsubscript{2} or any of a number of other small substrates.\textsuperscript{1} Although several small electron transfer proteins (e.g. flavodoxin or ferredoxin) or electron mediators can donate electrons to oxidized metal clusters in the MoFe protein, none of these molecules can support substrate reduction.

These observations show that the Fe protein is more than a simple reductant of the MoFe protein, suggesting instead a specific task such as inducing conformational changes within the MoFe protein that allow substrate binding and/or electron transfer to FeMo-cofactor. This is supported by a recent report that electron transfer between the Fe protein and MoFe protein is conformationally gated, involving protein conformational changes.\textsuperscript{5} A number of X-ray structures of Fe protein-MoFe protein complexes have been solved in various nucleotide-bound states,\textsuperscript{6} yet no structural changes have been observed within the MoFe protein that might explain the role of the Fe protein/ATP.

In recent years, we have probed MoFe protein function by substituting specific amino acids for different amino acids and then characterizing the properties of the resulting MoFe protein variants.\textsuperscript{4} We reasoned that it might be possible to explore the putative Fe protein conformational change within the MoFe protein by means of amino acid substitution of residues near the P cluster, FeMo-cofactor, or in between these two clusters (Figure 6-1). Here, we report that when the $\beta$-98\textsuperscript{Tyr} residue, located within the MoFe protein between the P-cluster and FeMo-cofactor, is substituted by histidine, the
resulting MoFe protein (β-98\textsuperscript{His}) can reduce the nitrogenous substrate hydrazine (N\textsubscript{2}H\textsubscript{4}) to ammonia in the absence of the Fe protein and ATP, with electrons being supplied instead by a Eu\textsuperscript{II}-polyaminocarboxylate complex.

When electrochemically generated Eu\textsuperscript{2+} (reduction potential -0.36 V versus the standard hydrogen electrode (SHE) at pH 7.0) is mixed with a polyaminocarboxylate ligand (L) such as EGTA (ethylene glycol-bis(2-aminoethyl)-N,N',N'-tetraacetate-(4-)) or DTPA (diethylenetriamine-N,N',N'',N'''-pentaacetate(5-)), it instantly forms a strong 1:1 complex that is a powerful and reactive one-electron reductant; the Eu\textsuperscript{III}/II-EGTA and Eu\textsuperscript{III}/II-DTPA redox couples have reduction potentials of -0.88 V and -1.14 V, respectively (pH 8.0).\textsuperscript{7}

When the wild-type MoFe protein without Fe protein/ATP is mixed with Eu\textsuperscript{II}-EGTA or Eu\textsuperscript{II}-DTPA, no significant reduction of N\textsubscript{2} or hydrazine to ammonia is detected (Figure 6-2, panel A).\textsuperscript{8} We also examined a number of MoFe proteins with amino acid substitutions near the P cluster or FeMo-cofactor: α-70\textsuperscript{Val}→Ala, α-195\textsuperscript{His}→Gln, β-188\textsuperscript{Ser}→Cys, and α-70\textsuperscript{Val}→Ala/α-96\textsuperscript{Arg}→His (Figure 6-1). Each of these residues in the MoFe protein has been previously implicated in some aspect of electron transfer or substrate reduction by the MoFe protein.\textsuperscript{4} When any one of these MoFe protein variants was mixed with Eu\textsuperscript{II}-DTPA, no significant reduction of hydrazine (Figure 6-2, panel A) or N\textsubscript{2} to ammonia was detected.
Figure 6-2: Hydrazine reduction to ammonia is shown as a function of time for different MoFe proteins (panel A) and for the β-98His MoFe omitting each component of the assay (panel B). No Fe protein or ATP is added. Omission of Eu²⁺ resulted in no detection of ammonia. The concentration of hydrazine is 50 mM, the quantity of MoFe protein is 0.7 mg (2.8 nmol), and the concentration of Eu⁺⁻DTPA is 0.5 mM (panel A) and 1 mM (panel B), with the pH = 7.0 and temperature = 25°C.
A prior study of a MoFe protein with $\beta$-98$^{\text{Tyr}}$ substituted by histidine, which is located between the P cluster and the $R$-homocitrate end of FeMo-cofactor (Figure 6-1), implicated the tyrosine in electron transfer between the clusters.\(^9\) When the $\beta$-98$^{\text{His}}$ MoFe protein was mixed with Eu$^{\text{II}}$-DTPA and hydrazine, a time-dependent production of ammonia was observed (Figure 6-2, panel A). The production of ammonia was dependent on the presence of the $\beta$-98$^{\text{His}}$ MoFe protein, Eu$^{2+}$, L, and hydrazine (Figure 6-2, panel B). After one minute of assay, a specific activity for hydrazine reduction of $170 \pm 8$ nmols NH$_3$/min/mg MoFe protein was determined. This rate is 94% of the rate measured for a fully functioning wild-type or $\beta$-98$^{\text{His}}$ MoFe protein with Fe protein, MgATP and a MgATP regeneration system, and the reductant dithionite under the same conditions. The reduction of hydrazine by the $\beta$-98$^{\text{His}}$ MoFe protein without Fe protein/ATP was found to continue over extended times (tested to 25 min), although the specific activity at longer times was lower (50 nmol NH$_3$/min/mg MoFe protein at 20 min) than at earlier times, suggesting that a component of the reaction was being depleted. Addition of more Eu$^{\text{II}}$-DTPA to an assay that had run for 20 min reactivated the reaction to near the initial rates, indicating that depletion of the reductant was the cause of declining rates over time. The rate of reduction of hydrazine depended on the concentration of Eu$^{\text{II}}$-DTPA and hydrazine, in each case showing saturation kinetics, with maximal rates observed at 1 mM Eu$^{\text{II}}$-DTPA and 50 mM hydrazine. Doubling the amount of MoFe protein doubled the rate of hydrazine reduction. The milder reductant Eu$^{\text{II}}$-EGTA was also examined and was found to support hydrazine reduction, albeit at lower rates (17 nmol NH$_3$/min/mg MoFe protein) when compared to Eu$^{\text{II}}$-DTPA.
Reduction of N\textsubscript{2} to ammonia by the β-98\textsuperscript{His} MoFe protein was not detected with either reductant even after extended reaction times. The Eu\textsuperscript{II}-DTPA and Eu\textsuperscript{II}-EGTA complexes catalyze H\textsuperscript{+} and acetylene reduction in the absence of MoFe protein, so it was not possible to test these substrates in this reaction.

The reduction of N\textsubscript{2} by nitrogenase is a 6 e\textsuperscript{-}/6 H\textsuperscript{+} reaction that is likely to involve metal-bound partially reduced intermediates at the level of reduction of diazene (N\textsubscript{2}H\textsubscript{2}) and hydrazine.\textsuperscript{10} Hydrazine is reduced to ammonia by the fully functioning nitrogenase,\textsuperscript{11} so it is reasonable to conclude that hydrazine joins late in the N\textsubscript{2} reduction pathway, requiring only 2e\textsuperscript{-}/2H\textsuperscript{+} for reduction.\textsuperscript{12} Our observation of hydrazine reduction by the β-98\textsuperscript{His} MoFe protein, but not N\textsubscript{2} reduction, is consistent with the fact that hydrazine is easier to reduce than N\textsubscript{2}.\textsuperscript{13-14} The hydrazine reduction observed here cannot be catalyzed by FeMo-cofactor that has been released from the protein because isolated FeMo-cofactor does not catalyze this reaction\textsuperscript{15} and the β-98\textsuperscript{His} MoFe protein is stable.\textsuperscript{9} Further, we have determined that the β-98\textsuperscript{His} MoFe protein retains normal acetylene reduction activity\textsuperscript{16} when combined with Fe protein and ATP following removal of the Eu\textsuperscript{II}-L by gel filtration chromatography.

The results presented here demonstrate the catalytic reduction of hydrazine to ammonia by the β-98\textsuperscript{His} MoFe protein uncoupled from electron delivery by the Fe protein and ATP, instead using only a simple, but powerful, electron donor. How substitution of β-98\textsuperscript{Tyr} by histidine has activated the MoFe protein for hydrazine reduction without the Fe protein and ATP hydrolysis has yet to be established, but the results indicate that amino acid substitutions within the MoFe protein can partially mimic some of the conformational changes induced by the Fe protein/ATP that either activate electron
transfer from the P-cluster to FeMo-cofactor or allow substrate binding. The specific roles of the P-cluster and FeMo-cofactor in the Eu$^{II}$-L-dependent reduction of hydrazine observed here are under active investigation.

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(16) Acetylene reduction specific activity for the b-98$^{\text{His}}$ MoFe protein with Fe protein and ATP was similar to that reported earlier (9). Hydrazine reduction specific activity with Fe protein and ATP was 180 nmol NH$_3$/min/mg MoFe protein at pH 7.0.
CHAPTER 7

SUMMARY AND FUTURE DIRECTIONS

During biological nitrogen fixation reduced Fe protein (bound to two ATP molecules) binds to the MoFe protein and undergoes conformational changes (Chapter 2).¹ These conformational changes allow an electron to be transferred from the P cluster to the M cluster, thereby generating a deficit of an electron at the P cluster. This deficit is in turn fulfilled by the [4Fe-4S] cluster of the Fe protein (Chapter 3).² Analysis of the conformational gating at lower temperatures shows that the mechanism of the electron transfer event is unaffected by temperature (Chapter 4).³ After the oxidation of the Fe protein (140 s⁻¹), hydrolysis of the two bound ATP molecules takes place (70s⁻¹) followed by phosphate release and dissociation of the Fe protein from the MoFe protein (Chapter 5). The aforementioned conformational changes within the MoFe protein, induced by the Fe protein, serve as a gate to the electron transfer from the P to M cluster. Via site directed mutagenesis, this gate can be opened in variant forms of the MoFe protein allowing transfer of electrons to the M cluster from sources other than the Fe protein (Chapter 6).⁴

Based on research presented in this dissertation, the above statements can now be made about electron transfer and substrate reduction in nitrogenase. Though this knowledge increases our understanding of the mechanism of nitrogenase such as the order of events, the mechanism of ET, and the nature of conformational gating of intramolecular electron transfer within the MoFe protein, many core questions still remain. These will be discussed here in hopes for providing a guideline for future work.
Conformational Changes in Nitrogenase

Conformational changes occur in the two proteins in nitrogenase complex. Fe protein undergoes conformational changes upon the binding of two ATP molecules. This conformational change includes the movement of the [4Fe-4S] cluster of the Fe protein ~5 Å towards the surface of the protein. Upon complex formation, this conformational change aids in the electron transfer event by decreasing its distance from the P cluster.

Conformational changes in the Fe Protein of nitrogenase: In Chapters 2, 3, and 4 of this dissertation we have presented evidence for conformational changes in the Fe protein related to the electron transfer event. The use of osmotic pressure has allowed us not only to understand that only ~800 Å² from the suggested ~2000 Å² area of conformational changes is involved in electron transfer event, but that these changes are involved in the electron transfer from the P cluster to the M cluster upon Fe protein binding.

Questions still remain regarding the nature of the other events in the Fe protein cycle. Based on work presented in Chapter 5, we know that the ATP hydrolysis occurs after the electron transfer event. However, we are unaware of any conformational changes or gates between the electron transfer event and the ATP hydrolysis. Similar questions remain as gaps in our knowledge regarding both phosphate release and dissociation events. Studies conducted using the osmotic pressure effect of the viscogens used i.e. glycerol, sucrose, polyethylene glycol 300, 600, and glucose on the ATP hydrolysis, phosphate release and dissociation events might provide us insights into other conformational changes occurring in the Fe protein of nitrogenase.
Figure 7-1: Osmotic pressure effects on phosphate release. Phosphate release is measured as an increase in fluorescence due to the binding of phosphate (Pi) to MDCC labeled phosphate binding protein. Each trace represents varying osmotic pressure (0-0.9 molal (m)) conditions in the system. The rate for each trace was calculated using the model described in Chapter 5.

Initial work towards understanding these conformational changes has already begun. Conformational changes related to the release of phosphate and dissociation are being studied. Based on initial results (Figure 7-1) we can say that the osmolytes e.g. sucrose shows that the phosphate release event is conformationaly gated. The rates of the phosphate release event under differing osmotic pressure conditions have been analyzed (Figure 7-2) using the methodology described in Chapter 2. The slope of the change in rate of the phosphate release event is exactly double the slope of the change in rate of the
electron transfer event (Chapter 2). Based on these two osmolyte studies conducted thus far, we can say that ~1600 Å² area is involved from the complex formation to the release of phosphate from the Fe protein of nitrogenase.

**Figure 7-2: logk vs. molality of 1°ET vs. Pi release (logk_{ET} vs. logk_{Pi}).** Percent maximum of logk vs. molality for phosphate release and 1° ET using polyethylene glycol and sucrose as osmolytes. The slope of logk vs. molality doubles in the case of each osmolyte for phosphate release indicating doubling of area for conformational change for the system.

The area involved in the complex formation to the primary electron transfer event (~800 Å²) includes any conformational changes that occur prior to the release of an electron from the Fe protein. Similarly, the area of conformational changes involved in the release of phosphate from the Fe protein involves all conformational changes prior to it. This includes the conformational changes involved in the electron transfer event. Thus, the area involved in the release of phosphate after the electron transfer event is ~ 700-800 Å².
The ability to differentiate the conformational changes occurring from one step in the Fe protein cycle to another highlights the importance of these osmolyte studies. Based on conformational changes from the electron transfer event to the phosphate release event we can deduce that the protein is undergoing a certain amount of changes after each event. This data is in agreement with the earlier crystallographic studies conducted with ATP analogues (Figure 7-3).\textsuperscript{12,13}

![Protein-Protein docking geometries in AMPPCP and ADP bound Fe:MoFe complex. AMPPCP bound complex (left) (PDB ID 2AFK)\textsuperscript{12} shows the interactions between the Fe protein (magenta and yellow) and the MoFe protein (grey and green) resulting in the smallest distance between the [4Fe-4S] cluster of the Fe protein and the P cluster of the MoFe protein. ADP bound complex (right) (PDB ID 1M34) shows an increased distance between the two clusters due to the motion of the Fe protein away from the MoFe protein.](image)

Figure 7-3: Protein-Protein docking geometries in AMPPCP and ADP bound Fe:MoFe complex. AMPPCP bound complex (left) (PDB ID 2AFK)\textsuperscript{12} shows the interactions between the Fe protein (magenta and yellow) and the MoFe protein (grey and green) resulting in the smallest distance between the [4Fe-4S] cluster of the Fe protein and the P cluster of the MoFe protein. ADP bound complex (right) (PDB ID 1M34) shows an increased distance between the two clusters due to the motion of the Fe protein away from the MoFe protein.

The crystallographic studies on nitrogenase provide a picture of the different binding forms of the Fe protein to MoFe protein as analogues to different stages in the ATP hydrolysis events bound to the Fe protein.\textsuperscript{12} However, these studies provide no
information on the dissociation event itself. Osmolytes could be used to determine the area of conformational change involved in the dissociation event. Two approaches, in conjunction with the osmolytes, could be used. Firstly flavodoxin (a biological reductant of the Fe protein) oxidation could be used to monitor the dissociation of the Fe protein from the MoFe protein. Secondly, since dissociation of the Fe protein from the MoFe protein is the rate limiting step, rate of substrate reduction (H\(^+\) reduction to hydrogen) could be monitored with varying osmotic pressure to understand the conformational changes in the dissociation event. Both strategies have limitations and require proper controls. The use of flavodoxin as a reductant requires that its reduction of oxidized Fe protein is either not conformationally gated or is negligibly so. Similar questions need to be answered about product release.

In the crystallographic studies, no changes have been observed in the pre and post hydrolysis analogue (AMPPCP and ADP-AlF\(_4\)) bound complex of the Fe and MoFe protein. Moreover, though changes have been observed in the ADP bound complex, it is not known whether it is a state near or far from the phosphate release event. By performing osmotic pressure studies on the ATP hydrolysis event we can ascertain conformational changes that take place after the ATP hydrolysis event but prior to phosphate release. These studies could provide us with the complete picture of the different motions of the Fe protein on the surface of the MoFe protein during the Fe protein cycle.

Many surface residues on both Fe protein and MoFe protein have been studied in the past for their role in complex formation, electron transfer, and dissociation (e.g 100\(^{\text{Arg}}\), 140\(^{\text{Arg}}\) on the Fe protein and \(\alpha\) and \(\beta\)-125\(^{\text{Phe}}\) on the MoFe protein) (Figure 7-4).
Osmotic pressure studies in combination with site directed mutagenesis could also be used in further understanding the role of these residues in each event during the Fe protein cycle.

Figure 7-4: Interactions of surface residues between Fe and MoFe protein. Surface residues in the Fe protein (100 and 140Arg) and α and β-125Phe in the MoFe protein (β-subunit grey, α-subunit green) are shown in the AMPPCP complex crystal structure (PDB ID 2AFK).

Conformational changes in the MoFe protein: Conformational changes on the MoFe protein are much harder to visualize since no crystallographic evidence exists. In fact, crystal structures of the nitrogenase complex using ATP analogues i.e. AMPPCP, ADP+Pi analogue i.e. ADP+AlF$_4^-$, and ADP have been used to argue against significant conformational changes in the MoFe protein. These arguments have been aided by the fact that all but one (2MIN: oxidized MoFe protein) of the crystal structures of the MoFe protein show no significant conformational changes.
A major evidence supporting the role of conformational changes in the MoFe protein in its reduction of substrate is presented in this dissertation (Chapter 7). This initial study suggests not only the occurrence of conformational changes in the MoFe protein, but also its importance in electron transfer to the M cluster and substrate reduction. This study shows that by making a single mutation on the MoFe protein between the P and M cluster, the MoFe protein is no longer dependent on the Fe protein as its sole reductant. The β-98Tyr-His variant of the MoFe protein is capable of accepting electrons from a low potential reductant, polyaminocarboxylate ligated EuII (Em -1.2 V vs. NHE), and of using these electrons to reduce hydrazine to ammonia.4

This initial result provides us evidence for the role of protein in facilitating electron transfer to the FeMo-cofactor. However, this does not provide us with the understanding of how the protein is facilitating this electron transfer. In order to understand this phenomenon, further analysis is needed. Survey studies using the EuII-ligand reductant need to be performed over a wide range of MoFe protein variants. Substrates other than hydrazine need to be tested for reduction using β-98Tyr-His and EuII-L reductant. Crystallographic studies on the β-98Tyr-His variant need to be performed. EPR analysis data needs to be gathered to further understand the electronic structure of FeMo-cofactor of the β-98Tyr-His variant during a EuII-L based turnover event. More MoFe protein variants need to be made with mutations in the region between the P and M cluster.

Some initial work towards the goals described above has been accomplished. A small survey on some of the variant MoFe proteins available has been conducted and two other mutations lying between the P and M cluster have already been identified as similar
to β-98Tyr-His variant of MoFe protein (Figure 7-5). These other two variant MoFe proteins α-64Tyr-His and β-99Phe-His have activities similar to that of the β-98Tyr-His variant (Figure 7-6).

Figure 7-5: α-64Tyr, β-98Tyr, and β-99Phe in mesh representation. The three amino acid residues relevant to the Fe independent reduction of the substrates (N₂H₄, N₃⁻ and H₂) by the MoFe protein are highlighted in Mesh representation (PDB ID 3U7Q).

The specific activities, Vmax, and Km of all three variant MoFe proteins with Eu⁺⁻L as the source of electron have been determined. The specific activities for the three MoFe variants at 100 mM hydrazine were 295 +- 15, 198 +- 3, and 110 +- 2 nmols of ammonia per mg of protein per minute for α-64Tyr-His, β-98Tyr-His, and β-99Phe-His, respectively. Varying hydrazine concentration was used to determine Km and Vmax for the three MoFe variants with Eu−L as the electron source. The data was fit to Michaelis Menten equation and Km values of ~35, 21, and 89 mM hydrazine were found for α-64Tyr-His, β-98Tyr-His, and β-99Phe-His, respectively. Vmax values for α-64Tyr-His, β-98Tyr-His,
and $\beta$-99$^{\text{Phe-His}}$ were found to be ~424, 248, and 316 nmols of $\text{NH}_3/\text{min/mg}$ of protein respectively (Figure 7-7).

![Figure 7-6: Time course analysis of the reduction of hydrazine to ammonia by ▲ $\alpha$-64$^{\text{His}}$, ● $\beta$-98$^{\text{His}}$, and ■ $\beta$-99$^{\text{His}}$ variant of the MoFe protein.](image)

The assays were performed by adding 1mM Eu$^{\text{II}}$DTPA to a solution containing 0.7 mg of MoFe protein and 100 mM hydrazine.

These initial results, though encouraging, are far from a complete picture of events inside the MoFe protein. The kinetic data highlights the need of further studies on variants between the P and the M cluster. It also highlights the need for combinations of mutations for $\alpha$-64$^{\text{Tyr}}$, $\beta$-98$^{\text{Tyr}}$, and $\beta$-99$^{\text{Phe}}$ residues.

Investigations into other substrates have yielded azide as a substrate for the $\alpha$-64$^{\text{Tyr-His}}$, $\beta$-98$^{\text{Tyr-His}}$, and $\beta$-99$^{\text{Phe-His}}$ proteins. Other substrates of nitrogenase are problematic with the Eu$^{\text{II}}$-L system since the reductant alone can reduce $\text{H}^+$, acetylene and $\text{CN}^-$ to hydrogen, ethylene and ammonia, respectively. Proton reduction has been
analyzed in greater detail by varying the polyaminocarboxylate ligand. Eu\textsuperscript{II}-EDTA does not generate detectable quantities of hydrogen in an aqueous system, however, it does generate some hydrogen with $\beta$-98\textsuperscript{His} variant of the MoFe protein. This opens up new avenues of future research in which reductants with more positive potentials than Eu\textsuperscript{II}-DTPA in combination with different and combinations of different mutations could be used to investigate these substrates.

**Figure 7-7:** Reduction of hydrazine to ammonia by $\triangle$ $\alpha$-64\textsuperscript{His}, $\bullet$ $\beta$-98\textsuperscript{His}, and $\blacksquare$ $\beta$-99\textsuperscript{His} variant of the MoFe protein with increasing hydrazine concentration. All assays were conducted for two minutes with 1mM Eu\textsuperscript{II}-DTPA and 100mM hydrazine concentration. The data was fit to Michaelis Menten equation.

**Electron Transfer Events in Nitrogenase**

Many advances in understanding the electron transfer event have been made in recent years and some are presented in this dissertation in Chapters 2,3,4, and 5 with the
The electron transfer event in nitrogenase is a two-step process that is initiated after an iron protein, bound to two MgATP molecules, binds to the MoFe protein. The first step is the electron transfer event from the P cluster to the M cluster of the MoFe protein. This event is triggered by the binding of the iron protein to the MoFe protein, which results in an electron transfer from the P cluster to the M cluster which is conformationally gated ($k_{3-MET} = 140 \text{ s}^{-1}$). This electron transfer results in a deficit of electron at the P cluster ($k_{3-PET} = 1700 \text{ s}^{-1}$).

The studies discussed in Chapter 3 show the electron transfer event from the Fe protein to the β-188 Ser-Cys variant of the MoFe protein studied using stopped flow UV-Vis spectrophotometry. This technique allows us to study the Fe protein without any interference from the MoFe protein. However, it also limits us to events occurring on the Fe protein to the P cluster ($k_{3-PET} = 1700 \text{ s}^{-1}$).

Rapid freeze quench EPR (RFQ) is a technique in which protein samples are mixed rapidly and quenched using a liquid nitrogen cooled isopentane bath. The frozen liquid is then packed inside an EPR tube and analyzed using an EPR spectrometer. This method is used for early time point analysis of electron transfer events. RFQ represents an attractive technique for studying the events on the P and M cluster of nitrogenase. However, this method is affected by a few problems. A major issue is the packing of the protein sample and quantification of this packed sample. In the past, RFQ
studies with nitrogenase have yielded some results though have failed to answer any of the major questions regarding electron transfer and substrate reduction in the different E states of nitrogenase.\textsuperscript{21,22} In order to overcome these problems, a novel technique for freeze quenching the nitrogenase enzyme in early time points after mixing was developed.

**Viscogen freeze quench:** Based on work conducted for chapters two and four of this dissertation we knew that osmolytes/viscogens could be used to slow down the electron transfer event in nitrogenase. We also knew that the addition of viscogens and lowering the temperature did not change the mechanism of electron transfer in nitrogenase. Based on this, a freeze quench technique was developed that takes into account the effect of high viscosity and low temperature in combination. At a temperature of \( \sim 4^\circ\text{C} \) EPR samples of \( \beta-188^{\text{Cys}} \) MoFe protein were made in highly viscous buffer (\( \eta>8 \)). A time course analysis was conducted by making individual samples at each time point.

Utilizing VFQ allowed us to study the EPR spectrum of the entire complex which in \( \beta-188^{\text{Ser-Cys}} \) variant of the MoFe protein means being able to study all three clusters of nitrogenase.\textsuperscript{23} The electron transfer from the F cluster of the Fe protein to the P cluster of the MoFe protein was thus observed. The resting state FeMo-cofactor EPR signal remained undisturbed during this event (Figure 7-9). If the electron had traveled to the resting state M cluster its signal would have begun to disappear. Instead, the EPR signal of the P cluster changed. The change in the EPR signal of the P cluster was found to be identical to the change in the signal for the Fe protein [4Fe-4S] cluster in the same
samples. This serves as evidence for the electron from the Fe protein being transferred to the P cluster of the MoFe protein in accordance with the deficit spending model.

Figure 7-8: Percent maximum EPR signal intensity comparison of P cluster ●, M cluster ♦, and the [4Fe-4S] cluster of the Fe protein▲. The data shows that the Fe protein oxidation is concurrent with the reduction of the P cluster.

However, since the β-188Ser-Cys variant of the MoFe protein possesses a Pox cluster in the dithionite reduced state, it serves as evidence for electron transfer to an artificially generated Pox cluster. For deficit spending model of electron transfer to be proven, the Pox species would have to be discovered during the primary electron transfer event. VFQ conducted with wild type MoFe protein with a PN cluster does not show any evidence for a Pox cluster. This is understandable due to the high rate of electron transfer from the F cluster to the Pox cluster. Whereas in the β-188Ser-Cys, with its Pox cluster pre oxidized, binding of the iron protein is rate limiting during a VFQ experiment, in the wild type
protein \( P^{\text{ox}} \) is generated after the binding of the Fe protein and is thus unaffected by conditions generated due to high viscosity and lower temperature.

This emphasizes the need for further study in understanding the order of the electron transfer events in nitrogenase. Iron protein variants that can trigger the P to M electron transfer without allowing the backfill to \( P^{\text{ox}} \) are the most attractive avenues of future research since they allow the use of wild type MoFe protein. Variant forms of MoFe protein that are not in immediate vicinity of either the P or the M cluster but have an effect on the electron transfer and substrate reduction behavior e.g. \( \beta-98^{\text{Tyr-His}}, \alpha\beta-125^{\text{Phe-Ala}}, \alpha-281^{\text{Tyr-His}} \) could be good candidates for further exploration.\(^{18,24}\)

*Exploration of substrate binding and E-states of FeMo-cofactor using VFQ:* VFQ is an exciting new development in observing electron transfer to the MoFe protein. MoFe protein goes through eight successive electron transfer states in which it reduces substrates at varying E states (Figure 7-10).\(^{25-27}\)

![Diagram](image)

**Figure 7-9:** Lowe-Thorneley model of electron transfer and substrate binding in nitrogenase. Each successive state of FeMo-cofactor shows an E state.
VFQ can be used to study more than one electron transfer to the MoFe protein. Initial studies for these electron transfer events have already been conducted. Using $\alpha$-195$^{\text{His-Gln}}$ variant of the MoFe protein, the binding of acetylene to the cofactor has been studied (Figure 7-11)$^{28,29}$. In Figure 7-11 (A) we can see the resting state ($E_0$) signal of the MoFe protein disappear over time into $E_1$ (EPR silent) followed by the $2e^-$ state $E_2$ in (B) which is identified by its unique EPR signal with this variant of the MoFe protein. Thus, here, VFQ allows us to follow the disappearance of $E_0$ and appearance of $E_2$. The data presented in Figure 7-11(C) show earlier E states during nitrogenase reduction due to the high viscosity of the system. By varying the viscosity of the system, rate of accumulation of latter E states in nitrogenase can be enhanced allowing us to study different trapped states of nitrogenase. VFQ can thus be used to study the $N_2$, propargyl alcohol, hydrazine, proton, and methylidiazene trapped states allowing us to map the route of the enzyme to a certain E state.$^{30-33}$ The technique can further be used to study FeMo-cofactor interactions with multiple substrates. One of the most promising avenue of investigation here is understanding the relationship between $N_2$ and hydrazine (i.e. are they inhibitory towards each other).

**Kinetic Isotope Effect in the Fe Protein cycle**

Kinetic assays for the observation of product formation were conducted under H$_2$O and D$_2$O as a solvent for the enzyme. A kinetic isotope effect was observed in D$_2$O. The activity of the enzyme in D$_2$O dropped to 50% of the activity in H$_2$O.

In order to determine which step in the Fe protein cycle is affected by the change in solvent, electron transfer, ATP hydrolysis, phosphate release and dissociation of the Fe
Figure 7-10: VFQ analysis of the early E states in α-195Gln MoFe protein. (A) shows the disappearance of the resting state EPR signal into an EPR silent E\textsubscript{1} state. (B) shows the appearance of the E\textsubscript{2} state (EPR active, acetylene bound) from the EPR silent E\textsubscript{1} state. (C) shows the percent maximum EPR signal of each E state fitted with a sequential model developed in Kintek Global Explorer. The model can be used to simulate the amount of E\textsubscript{1} signal that accumulated between E\textsubscript{0} and E\textsubscript{1}.
protein from the MoFe protein was analyzed in D\textsubscript{2}O (Figure 7-12). The results of these experiments indicate that the phosphate release event slows down to $\sim 5$ s\textsuperscript{-1} in D\textsubscript{2}O from 22 s\textsuperscript{-1} in H\textsubscript{2}O. Since the rate of phosphate release in D\textsubscript{2}O is nearly identical to the rate of dissociation, which is rate limiting at 6 s\textsuperscript{-1} and is unaffected by D\textsubscript{2}O, it is not enough to account for the 50\% decrease in rate of substrate reduction. The most likely explanation is that the Fe protein undergoes conformational changes after the release of phosphate but prior to the dissociation of Fe protein and these two steps (i.e. Pi release and dissociation) in combination result in the overall slowing of the enzyme in D\textsubscript{2}O.

The use of VFQ in exploring the KIE: VFQ analysis on the accumulation of the acetylene trapped state was conducted with D\textsubscript{2}O as a solvent. The kinetics of the first electron transfer event from the Fe protein to the MoFe protein in D\textsubscript{2}O were identical to these kinetics in H\textsubscript{2}O. However, marked differences were observed in the accumulation of the E\textsubscript{2} (2e\textsuperscript{-} transferred to FeMo-cofactor) state of the cofactor. In D\textsubscript{2}O the E\textsubscript{2} state was not observed. After the electron transfer event from the Fe protein to the MoFe protein, ATP hydrolysis, phosphate release and dissociation occur. However, in D\textsubscript{2}O, one of these events slows down to where the accumulation of E\textsubscript{2} in D\textsubscript{2}O nearly halts (Figure 7-13). The VFQ data, thus, supports the hypothesis presented earlier in that the phosphate release event itself might not be the only step slowing the accumulation of E\textsubscript{2} and that it might be multiple steps that contribute to generate the D\textsubscript{2}O effect.

The dissociation event in Figure 7-12 (D) is studied by poising the Fe protein in its oxidized state after phosphate release has occurred. The un-dissociated complex is then mixed with dithionite and the dissociation of Fe protein from the MoFe protein is
Figure 7-11: Kinetics of key nitrogenase catalytic steps in H$_2$O and D$_2$O at 25°C. (A) The transfer of an electron is monitored as the change in absorbance at 430 nm as a function of time (Blue in H$_2$O and red in D$_2$O). (B) ATP hydrolysis is measured by monitoring the formation of $\alpha$-32P-ADP as a function of time. The data were fit to the kinetic model described in chapter 5 with a calculated $k_{ATP}$ of $\sim$70 s$^{-1}$ for both H$_2$O and D$_2$O. (C) Phosphate (Pi) release is monitored by the increase in fluorescence from Pi binding to MDCC-PBP as a function of time. (D) Dissociation of Fe from the MoFe protein is monitored by the decrease in absorbance at 430 nm as a function of time.
Figure 7-12: VFQ analysis of α-195$^{Gln}$ MoFe protein with acetylene in D$_2$O. (A) shows the disappearance of the resting state EPR signal into an EPR silent E$_1$ state. (B) shows the lack of appearance of the E$_2$ state (EPR active, acetylene bound) from the EPR silent E$_1$ state. (C) shows the comparison between the E$_2$ states of H$_2$O vs. D$_2$O.

observed. This specific part of the dissociation at the end of a Fe protein cycle is unaffected by D$_2$O and no current method exists to study real time dissociation of the Fe protein from the MoFe protein (from complex formation to dissociation).
Studying the effect of osmotic pressure in the dissociation event as described earlier could provide an answer regarding the role of conformational changes in this D$_2$O effect. However, why the phosphate release event from the Fe protein slows down in D$_2$O remains unknown. Future phosphate release studies by utilizing variant proteins on both the MoFe protein (e.g. $\beta$-98$^{\text{Tyr}}$-$\text{His}$, $\beta$-188$^{\text{Ser}}$-$\text{Cys}$ etc.) and the Fe protein (e.g. variants in the switch I and II region of the Fe protein)$^{11,34–36}$ might be able to provide an insight into the effect of D$_2$O on the phosphate release event in the Fe protein cycle.

REFERENCES


APPENDIX
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