Turnover-Dependent Inactivation of the Nitrogenase MoFe-Protein at High pH

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Supporting Information

ABSTRACT: Proton uptake accompanies the reduction of all known substrates by nitrogenase. As a consequence, a higher pH should limit the availability of protons as a substrate essential for turnover, thereby increasing the proportion of more highly reduced forms of the enzyme for further study. The utility of the high-pH approach would appear to be problematic in view of the observation reported by Pham and Burgess [(1993) Biochemistry 32, 13725–13731] that the MoFe-protein undergoes irreversible protein denaturation above pH 8.65. In contrast, we found by both enzyme activity and crystallographic analyses that the MoFe-protein is stable when incubated at pH 9.5. We did observe, however, that at higher pHs and under turnover conditions, the MoFe-protein is slowly inactivated. While a normal, albeit low, level of substrate reduction occurs under these conditions, the MoFe-protein undergoes a complex transformation; initially, the enzyme is reversibly inhibited for substrate reduction at pH 9.5, yet in a second, slower process, the MoFe-protein becomes irreversibly inactivated as measured by substrate reduction activity at the optimal pH of 7.8. The final inactivated MoFe-protein has an increased hydrodynamic radius compared to that of the native MoFe-protein, yet it has a full complement of iron and molybdenum. Significantly, the modified MoFe-protein retains the ability to specifically interact with its nitrogenase partner, the Fe-protein, as judged by the support of ATP hydrolysis and by formation of a tight complex with the Fe-protein in the presence of ATP and aluminum fluoride. The turnover-dependent inactivation coupled to conformational change suggests a mechanism-based transformation that may provide a new probe of nitrogenase catalysis.

Substrate reduction by the molybdenum-dependent nitrogenase involves two protein components, the molybdenum iron protein (MoFe-protein), containing the active site for substrate reduction, and the iron protein (Fe-protein), serving as the unique ATP-dependent reductant for the MoFe-protein. A striking feature of the nitrogenase-catalyzed reaction is that the electron flux through the system is independent of the substrate that is being reduced, under a given set of conditions, the number of electrons transferred to substrate per active site per unit time is the same for the reduction of dinitrogen to ammonia (the physiological reaction), the reduction of acetylene to ethylene (commonly used to assay nitrogenase activity), or the reduction of protons to dihydrogen, which occurs in the absence (or sufficiently low concentrations) of other reducible substrates. An important characteristic of the nitrogenase reaction is that substrates can bind only to forms of the MoFe-protein reduced by two or three electrons relative to the “as-isolated” form, which can be efficiently generated only in the presence of the reduced Fe-protein and ATP. Efforts to generate significant populations of more highly reduced forms of the MoFe-protein for biophysical or structural characterizations are confounded by the ubiquitous presence of protons that are reduced to dihydrogen with the concomitant return to the initial stages of the catalytic cycle.

While protons cannot be eliminated from the aqueous solution environment, their concentration can be reduced by working at higher pHs. The pH dependence of nitrogenase activity may be described as a bell-shaped curve with optimal activity occurring around pH 7–8 and decreasing at both lower and higher pHs (half-maximal pHs of ~6.5 and 8.5, respectively). While it is not surprising that nitrogenase activity decreases at high pH, Pham and Burgess reported that the MoFe-protein is “irreversibly damaged by preincubation above pH 8.65” in a 3 min incubation, through a process suggested to arise from “a critical group on the MoFe protein with a pK in that range whose deprotonation leads either to cluster destruction or to an irreversible change in the structure of some critical part of the protein.” Indeed, the idea that the MoFe-protein denatures at pHs outside the optimal range has
been reflected in the subsequent design of pH-dependent experiments of nitrogenase activities (see ref 11). However, while attempting to trap substrates and inhibitors of nitrogenase for crystallization studies, we observed a distinct type of high-pH inactivation of the MoFe-protein that was not simple denaturation. As we report here, inactivation of the MoFe-protein at higher pHs is not due to protein instability but is rather the consequence of a complex, mechanism-based reaction with the potential to provide insight into the mechanism(s) of nitrogenase.

## Materials and Methods

### Preparation of Nitrogenase Proteins.

The MoFe-protein and Fe-protein from *Azotobacter vinelandii* (designated Av1 and Av2, respectively) were isolated under anaerobic conditions as previously described.12 The specific activities for acetylene reduction were ∼2200 nmol of ethylene min⁻¹ mg⁻¹ for Av1 and ∼1800 nmol of ethylene min⁻¹ mg⁻¹ for Av2. Unless otherwise noted, all operations were conducted anaerobically by appropriate manipulations using a Schlenk line connected to oxygen-scrubbed argon or in an anaerobic chamber.

The component ratio (CR) of Fe-protein to MoFe-protein is defined as the moles of Av2 per mole of Av1 active site (with two active sites per Av1 tetramer). The CR may be calculated using extinction coefficients of 76 and 9.4 mM⁻¹ cm⁻¹ for Av1 and Av2, respectively. The protein concentration was determined by absorbance at 410 nm using extinction coefficients of 76 and 9.4 mM⁻¹ cm⁻¹ for Av1 and Av2, respectively.

### Tribuffer Systems.

Because the kinetics of reduction of the substrate by nitrogenase are sensitive to both pHs13−10 and ionic strength,13,14 it is critical to use solution conditions under which the two parameters may be varied independently. For the pH range of 7.8–10 utilized in this study, a tribuffer system developed by Ellis and Morrison15 was employed, composed of three components: 100 mM N-(2-acetamido)-2-aminoethanesulfonic acid (ACES), 50 mM tris(hydroxymethyl)aminomethane (Tris), and 50 mM ethanolamine, with pH values of 6.67, 8.0, and 9.5 at 30 °C, respectively. Over the pH range from 8 to 10, the ionic strength of this buffer ranges from 0.096 to 0.100 M. We note that this is a range from 8 to 10, the ionic strength of this buffer system di-

### ATP Hydrolysis As Measured by Creatine.

Because 1 mol of creatine is produced by the creatine kinase-catalyzed phosphorylation of 1 mol of ADP by creatine phosphate, the amount of ATP hydrolyzed was established by measuring the change in creatine concentration in the reaction mixture.16,17

### Formation of the Av1−Av2−ADP-AlF4⁻ Complex.

The ADP-AlF4⁻-stabilized Av1−Av2 complex was prepared following the protocols of Renner and Howard.17 Inactive Av1 at a final concentration of 0.066 mg/mL and Av2 at a CR of ∼4 were incubated in 100 mM MOPS, 50 mM Tris, and 100 mM NaCl (pH 7.3) with 10 mM sodium dithionite, 4 mM NaF, and 0.2 mM AlCl₃. For the reaction in the presence of ATP, 1 mM ATP, 8 mM MgCl₂, and the creatine kinase-based ATP regenerating system were added. For reactions with ADP, 1 mM ADP and 8 mM MgCl₂ were added, without an ATP regenerating system.

The reactions were allowed to proceed for 30 min at 30 °C and the products analyzed by analytical column chromatography.

### Nitrogenase Activity Assays.

The activity of the nitrogenase proteins was determined by the reduction of acetylene to ethylene as expressed in the head space gas equilibrated with the reaction solution. The assay contained 20 mM sodium dithionite (the source of reducing equivalents and also to maintain anaerobic conditions) and an ATP regeneration system (5 mM MgCl₂, 5 mM ATP, 20 mM creatine phosphate, and 23 units/mL creatine phosphokinase).12 For the standard assay, the reaction mixture was buffered with 50 mM Tris-HCl (pH 7.8), while for reactions above pH 8, the tribuffer system described above was utilized. The pH of the tribuffer with ATP and creatine phosphate was adjusted by adding 6 M NaOH to a value higher than the target pH, so that the final pH was 9.5 (or other target pH) after processing, in order, addition of the kinase and MgCl₂, multiple cycles of degassing, addition of dithionite, and finally introduction of the component proteins. Because these common buffers are strongly dependent on temperature, the final pH was verified by measurement at 30 °C for a test vial with all components. No inhibitory effects on acetylene reduction by nitrogenase were observed for the tribuffer when compared to the standard assay. For the kinetic studies, the reduction of acetylene was initiated by the addition to the reaction assay of premixed Av1 and Av2 to achieve the desired concentrations and component ratio. The volume of the added Av1/Av2 mixture was kept constant by addition of 50 mM Tris-HCl (pH 7.75), 200 mM NaCl, and 5 mM sodium dithionite, so that the final ionic strength was independent of the CR. The anaerobic assay reactions were conducted at 30 °C and for 10 min in 8.9 mL serum vials with a 1.0 mL assay solution and an argon atmosphere containing 1.0 mL of acetylene gas. The reaction was terminated by the addition of 0.25 mL of 1 M citric acid, and ethylene in the gas phase was quantified by gas chromatography.

In addition to the standard assay described above, two types of activity measurements were used: the “head space” and “specific activity” assays. In the head space assay, the activity was monitored by the rate of appearance of ethylene in the gas phase of the reaction vial while allowing the reaction to continue. Ethylene and acetylene were quantified by gas chromatography of gas aliquots (50 μL) removed from the head space at the designed time points. The specific activity of Av1 in the reaction vial was monitored by removing liquid samples from the assay mixture at specified time points, for immediate transfer into a pH 7.8 standard assay solution containing excess additional Av2 (CR of ∼60). The transfer time was <10 s, and the transfer initiated the standard 10 min, pH 7.8 assay. SigmaPlot version 11.0 (Systat Software, Inc.) was used for curve fitting and display of the experimental data.

### Analytical Column Chromatography.

The proteins in a reaction mixture were analyzed by size exclusion chromatography on a 1 cm × 30 cm column of Superdex S-200, run under anaerobic conditions. The elution profile was monitored by the absorbance at either 390 or 410 nm, as indicated.
Metal Analysis. The molybdenum and iron contents of Av1 were determined by inductively coupled plasma mass spectrometry (ICP-MS) using a Hewlett-Packard 4500 ICP-MS system. Iron and molybdenum standards (Ultra Scientific Analytical solutions for ICP-MS) over concentration ranges comparable to and overlapping with those of the protein were analyzed in tandem with the protein samples. Proteins and standards were prepared in ultrapure H2O containing 1% nitric acid (ICP-MS grade, Fluka) with appropriate reagent and buffer blanks. The protein concentration was determined by absorbance at 410 nm prior to 1:400 dilution in nitric acid to give a concentration of 35–50 μg/mL for aspiration by the ICP-MS instrument. The Fe and Mo concentrations were determined using 56Fe and 96Mo isotopes and normalized for the Av1 concentration. The values were verified using the other natural abundance isotopes for both elements.

Determination of the Crystal Structure for Native Av1 at pH 9.5. Av1 was crystallized at 22 °C in an anaerobic chamber using the sitting drop method with mixed drops containing 2μL of a 42 mg/mL Av1 solution and 4μL of the precipitant. The precipitant contained 150 mM ACES, 75 mM Tris, 75 mM ethanolamine (pH 9.79), 17–18% PEG 3350, 0.8 M NaCl, and 1 mM sodium dithionite. Upon addition of the protein, the final pH of the mixed drop was 9.5. For cryoprotection, crystals of native Av1 grown at pH 9.5 were successively transferred into the reservoir solution with increasing concentrations of 2-methyl-2,4-pentanediol (MPD) up to 20%. Diffraction data were collected on beamline 12-2 at the Stanford Synchrotron Radiation Lightsource (SSRL), processed with XDS, and scaled with SCALA. The structure was determined by molecular replacement with MOLREP, using as a search model the Av1 structure determined at 1.16 Å resolution [Protein Data Bank (PDB) entry 1M1N]. The refinement was performed with REFMAC and PHENIX, while graphics program COOT was used for displaying maps and rebuilding the model. Data and refinement statistics are summarized in Table 1.

Residue 440 of the α-chain was corrected to Gln rather than the Glu reported in all previous structures to reflect the correct A. vinelandii nifD gene sequence; this discrepancy was highlighted by a recent analysis of 95 MoFe-protein sequences demonstrating that all but two had Gln and Asn at this position, with no examples of Glu.

| RESULTS |

Time Course of Acetylene Reduction at High pH. The initial test of nitrogenase activity well above its pH optimum was conducted at pH 9.5 using the head space measure of ethylene formation with time as shown in Figure 1A. The rate curve indicated a progressive loss of product formation with complete cessation well before depletion of any component of the assay. Supplemenating the reaction mixture at longer times with fresh aliquots of dithionite, Av2, or components of the ATP regenerating system did not result in further substrate reduction, which confirmed that the loss of acetylene reduction activity was not due to depletion of any of these components. Together, these studies implicated progressive inhibition of the Av1 that can be described by a first-order process characteristic of slow inhibitors such as the aluminum fluoride inhibition of nitrogenase activity:

\[ P(t) = \frac{P_0}{k_i} (1 - e^{-k_i t}) \]  

| Table 1. Data and Refinement Statistics for the Determination of the Crystal Structure of Native Av1 at pH 9.5 |

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*Rmerge(I) = \sum_{hkl} (\sum I_{hkl} - \langle I_{hkl} \rangle)/\sum I_{hkl}, where I_{hkl} is the intensity of an individual measurement of the reflection with indices hkl and \langle I_{hkl} \rangle is the mean intensity of that reflection.|

where \( P_0 \), \( k_i \), and \( P(t) \) are the initial rate of product formation (the enzyme catalytic activity expressed in nanomoles per minute), the observed first-order inactivation rate constant (inverse minutes) of Av1, and the time-dependent amount of ethylene in the head space (nanomoles), respectively. \( v_0/k_i \) is the total amount of product produced at the limit \( t = \infty \). The solid lines in Figure 1 represent the nonlinear curve fitting of the data points based on eq 1. For this type of inhibition, the cessation of product formation at long time points (the plateau in Figure 1) implies a quasi-irreversible state of inhibition of the enzyme, at least under the pH 9.5 conditions of the assay. For a slowly reversible reaction, a flat line plateau would not be observed and the kinetic expression would have additional elements to include the slower back reaction.

The generality of the progressive inhibition as a consequence of pH was demonstrated by a series of experiments using the head space assay in the upper range of pH 8.8–9.8 that has been previously reported for various nitrogenase studies.

The data were fit using eq 1 shown as the solid lines in Figure 1B, with the calculated \( k_i \) and \( v_0 \) values given in Figure 1C. Consistent with the accepted nitrogenase pH activity dependence, the initial velocity, \( v_0 \), decreases with an increased pH. More significantly, the observed inactivation rate, \( k_i \), increases with an increased pH, suggesting that a deprotonated state of the enzyme is a precursor for the inhibition. This clearly separates the two phenomena: the enzymatic activity follows a pH-dependent decline, while the inhibition process increases with a higher pH.

Turnover Conditions Are Required for High-pH Inactivation of Av1. Inhibition of the enzymatic activity as a consequence of Av1 inactivation was established by the specific activity assay that determines the amount of active Av1 at a saturating Av2 concentration in the pH 7.8 assay. Following
Initiation of the reaction at pH 9.5, aliquots of the reaction mixture were sampled at designated time points and the Av1 specific activity was determined. As shown in Figure 2, Av1 from the pH 9.5 incubation was progressively inactivated as indicated by the loss of specific activity measured at pH 7.8. The time-dependent loss of Av1 specific activity at pH 9.5 required turnover of the complete system that was established by a series of changes in the pH 9.5 reaction conditions. If Av2 or the ATP regenerating system was omitted or if AMP-PNP, a nonhydrolyzable ATP analogue, was used, no acetylene reduction was detected at pH 9.5 (as expected), and through the first 2 h of incubation at pH 9.5, there was an only ~10% decrease in Av1 specific activity as determined at pH 7.8, a time in which there was ~90% inactivation under the full turnover conditions. Most importantly, incubation of Av1 or Av2 alone at pH 9.5 showed only minimal activity loss even after 4 h. Although acetylene reduction was the most convenient way to monitor enzyme activity, the inactivation proceeded equally well when dinitrogen replaced the acetylene or when there was no added substrate beyond the protons in the solution (see Figure S1 of the Supporting Information). Furthermore, the inhibitor of all substrates except protons, carbon monoxide, did not block the inactivation of Av1.

**Component Ratio and Protein Concentration Effects on the pH 9.5 Reaction.** Central to the understanding of the initiation of the reaction at pH 9.5, aliquots of the reaction mixture were sampled at designated time points and the Av1 specific activity was determined. As shown in Figure 2, Av1 from the pH 9.5 incubation was progressively inactivated as indicated by the loss of specific activity measured at pH 7.8. The time-dependent loss of Av1 specific activity at pH 9.5 required turnover of the complete system that was established by a series of changes in the pH 9.5 reaction conditions. If Av2 or the ATP regenerating system was omitted or if AMP-PNP, a nonhydrolyzable ATP analogue, was used, no acetylene reduction was detected at pH 9.5 (as expected), and through the first 2 h of incubation at pH 9.5, there was an only ~10% decrease in Av1 specific activity as determined at pH 7.8, a time in which there was ~90% inactivation under the full turnover conditions. Most importantly, incubation of Av1 or Av2 alone at pH 9.5 showed only minimal activity loss even after 4 h. Although acetylene reduction was the most convenient way to monitor enzyme activity, the inactivation proceeded equally well when dinitrogen replaced the acetylene or when there was no added substrate beyond the protons in the solution (see Figure S1 of the Supporting Information). Furthermore, the inhibitor of all substrates except protons, carbon monoxide, did not block the inactivation of Av1.

**Component Ratio and Protein Concentration Effects on the pH 9.5 Reaction.** Central to the understanding of the
complex kinetic properties of the two-protein component system of nitrogenase is that the activity is dependent on the total protein concentration and component ratio as important factors controlling the electron flux during turnover.27,28 As one example, for a fixed concentration of the MoFe-protein, the initial velocity for acetylene reduction increases with an increasing CR until a plateau level of activity is obtained, indicative of saturation kinetics; the specific activity of the MoFe-protein is determined from the plateau level of activity upon titration with the Fe-protein at pH 7.8. The same general behavior was also observed upon titration of a fixed amount of Av1 with different CRs of Av2 at pH 9.5, although the amounts of product formed per time are much smaller than at pH 7.8 (Figure 3). By fitting the head space data (Figure 3A) to eq 1, we can determine initial velocity $v_0$ as a function of CR, and $v_0$ is presented in the form of an Av1 titration curve. As shown in Figure 3B, the initial velocity exhibits a strong CR dependence similar in form to that observed at pH 7.8, suggesting the initial rate correctly reflects the nitrogenase reaction and its pH dependence. At the saturation of Av1 by Av2, the specific activity of Av1 at pH 9.5 was estimated to be 100 nmol min$^{-1}$ mg$^{-1}$, or ~5% of the value at pH 7.8. As also observed at the optimal pH,29–31 the Av1 titration curve at pH 9.5 is not fully hyperbolic and has a "lag" at low concentrations of Av2 (CR < ~1). This is consistent with a similar process of interactions between the MoFe-protein and Fe-protein that is observed at the optimal pH and is maintained at the higher pH. One measure of the association of Av2 and Av1 is the CR at half-saturation that at pH 9.5 is ~4 and similar to that at pH 7.8 (an absolute comparison is obviated by the necessary protein concentration differences with the approximately 2 order of magnitude difference in $v_0$). An additional test that shows that the enzymatic substrate reduction at pH 9.5 reflects a turnover mechanism similar to that at pH 7.8 is the protein concentration dependence of the reaction rate. As shown in Figure 3C, over an 8-fold protein concentration range and at a constant CR, $v_0$ increased ~6-fold, a relative increase comparable to that observed at pH 7.8.32–34

In contrast to the clear similarity of the patterns in turnover parameters at pH 9.5 and at the optimal pH, the inactivation rate constant, $k_i$, showed a fully different pattern with these variables (Figure 3B,C). Namely, $k_i$ is substantially independent of CR and protein concentration and was calculated to be ~0.07 min$^{-1}$ (corresponding to a half-life of ~10 min). The small variance of ~20% for $k_i$ is in contrast to ≥1 order of magnitude changes in $v_0$ values for the same CR. The implication is that whatever mechanism of inhibition is occurring, it is a first-order process that is independent of the overall nitrogenase turnover rate as measured by product formed.

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**Figure 3.** Component ratio and protein concentration dependence of Av1 inactivation monitored by the head space assay at pH 9.5 and 30 °C. (A) Time-dependent ethylene formation at various CRs with the Av1 concentration fixed at 0.25 mg per reaction. Solid lines are nonlinear curve fits to the data points by eq 1. (B) Initial rate ($v_0$) and inhibition rate ($k_i$) calculated according to eq 1. Error bars indicate standard errors from the curve fitting. (C) Initial rate ($v_0$) and inhibition rate ($k_i$) determined from progress curves at various protein concentrations and at a constant component ratio of 2.5. The data are plotted in terms of the Av1 concentration using an average of two determinations for each data point.
Effects of the Component Ratio on Specific Activity.
In contrast to the apparent first-order inhibition kinetics observed for the head space assay at pH 9.5, the inactivation kinetics monitored by the specific activity measured at pH 7.8 cannot be modeled by a single kinetic phase. As shown in Figure 4, the kinetics of inactivation measured by specific activity are quite sensitive to the CR used in the pH 9.5 reaction; e.g., for a CR of 0.15, incubation for ~3 h is required to eliminate 50% of the specific activity, while for CRs of >1, the corresponding time is ~30 min. Most importantly, the loss of specific activity is significantly slower than the inhibition observed at pH 9.5 based on the direct measure of product in the head space. As seen via comparison of Figures 3 and 4, for all CRs at 60 min, acetylene reduction at pH 9.5 has stopped while the specific activity ranges from 25 to 75%. This strongly implies that the inhibited state at pH 9.5 is partially reversible at pH 7.8. The time of incubation at pH 7.8 before initiating the specific activity assay showed no effect on the measured value, which suggests the reversible step must be fast compared to the minimal time for sampling of the pH 9.5 reaction into the pH 7.8 assay.

ATP Hydrolysis during Turnover. The kinetic properties of the reaction at pH 9.5 versus the loss of specific activity measured at pH 7.8 suggested that some level of interactions between Av1 and Av2 proceeds after cessation of the observed product formation in the head space gas. ATP hydrolysis occurs only in the complex between Av1 and Av2, although ATP hydrolysis is independent of the oxidation state of the two proteins. Indeed, while Av2 can bind MgATP, a conformational change stabilized by complex formation with Av1 is required for hydrolysis. The number of ATP equivalents turned over by the enzyme can be conveniently measured by following the formation of creatine in the ATP regenerating system. The results for the correlation of ATP equivalents used and product formation at pH 9.5 are plotted in Figure 5A for CRs of 0.3 and 1.0. The clear result is that ATP hydrolysis initially tracks the substrate reduction yet continues long after product formation has reached completion. Even for those low component ratios (e.g., CR = 0.3) where the level of ethylene production is below the threshold of detection at pH 9.5, ATP hydrolysis readily occurs long into the time course for inactivation and provides evidence that the Av2 is still active. Indeed, the ability of the various forms of Av1 generated during turnover at pH 9.5 to support ATP hydrolysis is unmistakably shown in Figure 5B for the changes in specific activity over the time course of inactivation. For a CR of 1.0 where at 240 min the Av1 is nearly fully inactivated, ATP hydrolysis is still supported at ~40% of the rate of native Av1 at saturation with Av2. The initial time points show an ATP:ethylene ratio of ~5, in keeping with the accepted value of 4−5 with dithionite as the terminal reducing agent, but the ratio rapidly increases to >50 for the later time points as the Av1 is inactivated. These high ratios represent the small fraction of active enzyme plus the increasing amount of inactive enzyme that is still capable of supporting substantial ATP hydrolysis.

Characterization of Native Av1 Incubated at pH 9.5.
The activity studies clearly indicated that Av1 was not denatured with loss of activity at pH 9.5. To directly assess the structural consequences of incubation of Av1 at pH 9.5 (in the absence of turnover) and to provide a control structure for future work at higher pHs, the crystal structure at pH 9.5 was...
Figure 6. Electron density map around the FeMo-cofactor and homocitrate of the A. vinelandii MoFe-protein crystallized at pH 9.5. The map is calculated at 2.0 Å resolution and contoured at 1.5 times the standard deviation. The yellow bonds and colored atoms represent the Av1 structure at pH 9.5, while the black bonds and atoms indicate the structure of Av1 at pH 8.0 as determined by Spatzal et al. at 1.0 Å resolution (PDB entry 3U7Q39). Overall, the two structures are quite similar (rmsd of 0.3 Å), although a displacement of the C1 carboxyl arm of homocitrate is indicated in the pH 9.5 structure compared to the pH 8.0 structure.

Characterization of Modified (inactive) Av1. To characterize the inactive protein, we isolated Av1 from the pH 9.5 reaction mixture after incubation for 4 h (specific activity of <10%) using the size exclusion chromatography column at pH 7.3. Intriguingly, as shown in Figure 7A, the peak position of inactive Av1 shifted to an earlier elution time, relative to the position observed for native Av1 or for Av1 incubated at pH 9.5 in the absence of turnover components. The shift in elution position indicates that the hydrodynamic radius of inactive Av1 has increased relative to that of native Av1. The observed shift in the hydrodynamic radius was a consequence of the pH 9.5 turnover as shown in Figure 7A; incubation of Av1 alone or in combination with Av2 without ATP did not cause a shift in the elution position. Indeed, the complete recovery of both Av1 and Av2 (as measured by the absorbance at 410 nm) from a 4 h incubation at pH 9.5, either alone or under turnover conditions, further substantiates the stability of the proteins at pHs as high as 9.5. This form of Av1, hereafter termed Av1-mod, is stable to multiple cycles of rechromatography at pH 7.3–9.0 in 100–200 mM NaCl.

The absorption at 410 nm arises from the metalloclusters, and the integrated peak areas for Av1 were similar for all samples, irrespective of pH and precise conditions of the turnover (the same total amounts of Av1 were initially used in the compared chromatographs). This observation implies that the metalloclusters in Av1-mod are not significantly changed through either loss of iron, rearrangement of clusters, or other changes in the metal environment. This latter conclusion was substantiated by the results of Fe and Mo analysis of native and Av1-mod as given in Table 2. Within the experimental uncertainty of the analysis, there are two important findings: incubation of native Av1 at pH 9.5 does not lead to metal loss as might be expected for a denatured protein, and likewise, inactivation of Av1 is not due to general denaturation with the loss of either iron or molybdenum.

The results of monitoring ATP hydrolysis during the two assay procedures (see the preceding section) clearly demonstrated that the inactive Av1 retained the ability to interact with Av2 in a way that induces the Av2 conformation necessary for nucleotide hydrolysis. To further evaluate the integrity of Av1-mod with respect to binding of ATP-bound Av2, the established ability of Av1 and Av2 to form a complex stabilized by ADP-AlF4− was investigated.17,42 For native Av1 and Av2, the complex can be formed by reaction of either MgADP or MgATP as the nucleotide component with AlF4−.17 The reaction with MgATP is faster than with MgADP but leads to the same final complex.17,43,44 As shown in Figure 7B, Av1-mod forms a faster-migrating species when it is incubated with Av2, ATP, and AlF4−. On the basis of this earlier elution time compared to that of Av1-mod, a stable complex is inferred. This peak contained Av2 (by sodium dodecyl sulfate–polyacrylamide gel electrophoresis) and is estimated to be a 2:1 complex (assuming the tetramer of Av1-mod). Under similar conditions, MgADP also induced a stable complex that was intermediate in elution time between Av1-mod and the putative 2:1 Av1-mod–2Av2[ADP-AlF4−] complex. However, in contrast to the similar study with the native Av1, formation of the complex with MgADP appears to give only a 1:1 Av2–Av1 complex (based upon elution position). The assumption for the native proteins is that AlF4− stabilizes a hypothetical transition state in electron transfer between Av2 and Av1 and that ATP hydrolysis leads to this state; by microscopic reversibility, ADP reaches the same state. For Av1-mod, the path by microscopic reversibility for ADP appears to be much slower, and only partial conversion to the complex is observed even after reaction for 30 min.

DISCUSSION

Protons have multiple roles in the catalytic mechanism of enzymes as reflected in the variation of activity with pH. Among these roles for nitrogenase are the proper protonation state of catalytic residues, the ionic state of residues in the nucleotide binding site, ionic states of residues involved in binding the Fe-protein to the MoFe-protein, and, most
(A) Reaction mixtures (1.0 mL) containing 1.0 mg of Av1, with or without Av2 at a CR of 3.4 and with or without the ATP regenerating system, were incubated at pH 9.5 in tribuffer and 50 mM NaCl at 30 °C for 3.5 h; 0.8 mL of the reaction mixture was injected from the reaction mixture onto the column that was equilibrated with 50 mM Tris buffer (pH 7.5) containing 200 mM NaCl and 5 mM sodium dithionite: elution line 1 (red), Av1 and Av2 incubated without the ATP regenerating system; elution line 2 (blue), Av1 alone; elution line 3 (black), complete turnover conditions, Av1 and Av2 with the ATP regenerating system. Elution was monitored at 410 nm. (B) Same column as in panel A equilibrated with 50 mM MOPS buffer (pH 7.3) containing 100 mM NaCl, 1 mM MgCl₂, 5 mM NaF, 0.25 mM AlCl₃, and 5 mM sodium dithionite. Av1-mod was prepared as described for line 3 of panel A and re-isolated in MOPS buffer (pH 7.3) without AlCl₃ or NaF. Av1-mod (~0.2 mg) was incubated with Av2 (CR = 4.6) in 2.4 mL of MOPS buffer containing 5 mM NaF, 0.25 mM AlCl₃, and either no nucleotide [elution line 1 (black)], 1 mM ATP regenerating system [elution line 2 (purple)], or 1 mM ADP [elution line 3 (green)]. After incubation at 25 °C for 30 min to form the complex, 0.9 mL of each reaction mixture was applied to the column. The absorption of the eluate was monitored at 390 nm.

Figure 7. Anaerobic size exclusion chromatography of Av1 on a 1 cm × 30 cm column of Superdex S-200. The flow rate was 0.5 mL/min at 22 °C. (A) Reaction mixtures (1.0 mL) containing 1.0 mg of Av1, with or without Av2 at a CR of 3.4 and with or without the ATP regenerating system, were incubated at pH 9.5 in tribuffer and 50 mM NaCl at 30 °C for 3.5 h; 0.8 mL of the reaction mixture was injected from the reaction mixture onto the column that was equilibrated with 50 mM Tris buffer (pH 7.5) containing 200 mM NaCl and 5 mM sodium dithionite: elution line 1 (red), Av1 and Av2 incubated without the ATP regenerating system; elution line 2 (blue), Av1 alone; elution line 3 (black), complete turnover conditions, Av1 and Av2 with the ATP regenerating system. Elution was monitored at 410 nm. (B) Same column as in panel A equilibrated with 50 mM MOPS buffer (pH 7.3) containing 100 mM NaCl, 1 mM MgCl₂, 5 mM NaF, 0.25 mM AlCl₃, and 5 mM sodium dithionite. Av1-mod was prepared as described for line 3 of panel A and re-isolated in MOPS buffer (pH 7.3) without AlCl₃ or NaF. Av1-mod (~0.2 mg) was incubated with Av2 (CR = 4.6) in 2.4 mL of MOPS buffer containing 5 mM NaF, 0.25 mM AlCl₃, and either no nucleotide [elution line 1 (black)], 1 mM ATP regenerating system [elution line 2 (purple)], or 1 mM ADP [elution line 3 (green)]. After incubation at 25 °C for 30 min to form the complex, 0.9 mL of each reaction mixture was applied to the column. The absorption of the eluate was monitored at 390 nm.

Table 2. Fe and Mo Contents of Native and Modified Av1

<table>
<thead>
<tr>
<th>sample</th>
<th>Fe (mg)</th>
<th>Mo (mg)</th>
<th>Fe:Mo ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>native Av1, 1</td>
<td>29.7 ± 0.5</td>
<td>1.99 ± 0.03</td>
<td>14.9 ± 0.5</td>
</tr>
<tr>
<td>native Av1, 2</td>
<td>29.7 ± 0.5</td>
<td>1.98 ± 0.03</td>
<td>15.0 ± 0.5</td>
</tr>
<tr>
<td>Av1-mod, 1</td>
<td>29.1 ± 0.5</td>
<td>1.99 ± 0.03</td>
<td>14.6 ± 0.5</td>
</tr>
<tr>
<td>Av1-mod, 2</td>
<td>29.0 ± 0.5</td>
<td>1.99 ± 0.04</td>
<td>14.6 ± 0.5</td>
</tr>
</tbody>
</table>

“Av1-mod was generated at pH 9.5 by incubation of Av1 and Av2 (CR = 4.0) under turnover conditions at 30 °C for 4 h. The protein was isolated by size exclusion chromatography at pH 9.5 using the same tribuffer that was used for the inactivation. The control sample of Av1 alone was isolated by chromatography at pH 7.5. Full duplicate reactions and isolations were prepared for both proteins. The metal content was determined by ICP-MS performed in triplicate for each sample. The standard deviation for ^56Fe was ±0.56% and for ^96Mo was ±0.22%, while for the protein, the concentration varied from ±1.5 to ±1.8%. The results are normalized to the protein concentration for each individual sample and given with the standard deviation based upon multiple determinations of an individual sample.”

While the inhibition at pH 9.5 requires enzyme turnover and, hence, is related to turnover, the inactivation was distinctly irreversibly inactivated by protein denaturation at higher pHs would preclude this potentially promising approach.

We have reinvestigated the effect of increased pH on Av1 with significantly different results and conclusions. First among our results is the finding that Av1 is neither denatured nor inactivated when it is incubated alone at pHs as high as 9.5. Av1 exhibited only minimal change in specific activity after incubation for several hours at pH 9.5 and 30 °C and could be crystallized at pH 9.5 with no important changes in the structure even after the several days needed to form the crystals. As Pham and Burgess found, we did observe a pH-dependent inactivation of Av1, but in contrast to their conclusions, we found that the inactivation is a consequence of catalytic turnover resulting in a modified but structurally intact protein; inactivation of Av1 requires both Av2 and ATP hydrolysis. At pH 9.5, the model condition studied here, the enzyme activity, while turning over, decreased with time with a pattern similar to that of "slow inhibitors" such that both the initial rate, v₀, and the inhibition rate constant, kᵢ, could be determined by following the product in the gas phase of the reaction. Evaluation of the changes in v₀ with protein concentration and component ratio indicated this rate was measuring the true enzyme reaction and was consistent with a decrease in catalytic activity expected over a measured pH range. Hence, we find that some forms of activity can be reliably studied at higher pHs when they are limited to appropriately determined initial rates, a condition generally applicable to all steady state enzyme kinetics.
different from the enzyme activity. The inhibition rate increased as the rate of enzyme turnover decreased; that is, the inhibition rate increases with an increase in pH as activity decreases. The following characteristics summarize the observations with some of the implications.

(i) The inactivation reaction, being turnover-dependent, suggests a process involving states of the MoFe-protein that are distinct from the as-isolated form, mostly likely a different redox state. Because the inactivation is independent of which added substrate is present, one of the first enzymatically reduced states of the MoFe-protein may be sufficient to lead to inactivation.

(ii) The inhibition at pH 9.5 appears to be first-order when it is monitored by the decrease in the rate of ethylene formation with time at pH 9.5 in the head space assay.

(iii) The rate of inhibition, $k_i$ (eq 1), appears to be independent of the component ratio, protein concentration, and the amount of product produced (at least for CRs above $\sim 1$). This indicates that the probability of inhibition is distinct from the probability of product formation.

(iv) Although there are insufficient data to assign reliable apparent $pK_a$ values for the groups associated with the $v_0$ and the $k_i$, the results in Figure 1C clearly show them to be different and, hence, derived from different functional groups.

(v) The decrease in specific activity upon turnover at pH 9.5 as measured at pH 7.8 follows a more complex kinetic pattern involving multiple phases and requiring significantly longer times (hours) for completion versus the time needed for the pH 9.5 inhibition.

(vi) The observed difference in the two inactivation/inhibition rates suggests that the inhibited state of Av1 at pH 9.5 is different from, although likely on the path to, the ultimately inactive Av1-mod. The first state must be somewhat reversible for activity to be seen at pH 7.8. For any CR used in Figure 3 at pH 9.5 and at 60 min, product formation has ceased, while at the same time point and pH 7.8, the specific activity retains 25–75% of the initial value, depending on the CR at pH 9.5. The final state, however, is irreversibly inactive in substrate reduction.

(vii) Turnover at pH 9.5 continues past the cessation of product formation as determined by continued ATP hydrolysis. For the fully inactive Av1-mod, the rate of associated ATP hydrolysis is $\sim 40\%$ of that of the native protein with a saturating Av2 concentration.

Av1-mod has an expanded conformation (increased hydrodynamic radius) that maintains structural integrity. The new conformation is stable over time, retains metal content within the limits set by the analytical methods, and has the ability to associate with Av2. The latter is demonstrated by supporting ATP hydrolysis and the formation of a tight complex mediated by nucleotide-AIF₄⁻. Together, these observations suggest a mechanism-based reaction with multiple forms of Av1 that can interconvert during the turnover at pH 9.5, some of which lead to inactivation. Although a comprehensive model that describes the kinetic behavior of this system under all these conditions would be desirable, we have demurred in such a construction because there are an enormous number of potential pH-dependent ionic states relating the multiple components. To assign $pK_a$ activity levels, inhibition rates, and association constants for the two components and for the nucleotides would be outside the number of parameters that we have collected. Indeed, although the pH 9.5 inhibition rate is first-order and follows "slow inhibitor" rate law, we have not found a simple, rational kinetic measure of the specific activity decrease that would follow from the pH 9.5 inhibition rate.

Most importantly, our work identifies some of the conditions promoting a mechanism-based inactivation of the MoFe-protein. This should provide both a new direction for probing the catalytic mechanism(s) and the conditions to evaluate studies at pH >8.5, e.g., the reported changes in electron allocation with pH or the trapping of intermediates. While the focus of this paper has been on the inactivation reaction at pH 9.5, the process carries to lower pHs; it is only that the inhibition at the higher pH is sufficiently fast to facilitate kinetic characterization. One may anticipate that the phenomenon of inactivation would also take place at more physiologically relevant pHs, albeit with a slower rate. For example, the inactivation process may be particularly sensitive to amino acid residues in the cofactor environment such that differences in apparent substrate reduction rates for mutant nitrogenase with amino acid substitutions may be a consequence of changes in inactivation rates even when they are studied at the native protein pH optimum. Just as the generation of highly oxidizing species in photosystem II during water oxidation leads to protein inactivation with subsequent adaptations for protein turnover, the generation of highly reducing species in nitrogenase during substrate reduction is also associated with protein inactivation, which may require protein turnover adaptations in vivo.

**ASSOCIATED CONTENT**

Supporting Information
Inactivation rates for Av1 with different substrates and inhibitors at pH 9.5 (Figure S1). This material is available free of charge via the Internet at http://pubs.acs.org.

Accession Codes
Coordinates and structure factors have been deposited in the Protein Data Bank as entry 4ND8.

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Notes
The authors declare no competing financial interest.

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**ABBREVIATIONS**

Av1 and Av2, MoFe-protein and Fe-protein from *A. vinelandii*, respectively; Av1-mod, stable form of the inactive MoFe-protein that can be isolated after incubation under turnover conditions at pH 9.5; ACES, N-(2-acetamido)-2-aminoethanesulfonic acid; Bis-Tris, [bis(2-hydroxyethyl)amino]tris(hydroxymethyl)methane; CHES, 2-(cyclohexylamino)-ethanesulfonic acid; CR, component ratio or molar ratio of active sites in Fe-protein and MoFe-protein; HEPPS, N-(2-hydroxyethyl)piperazine-N′-propanesulfonic acid; ICP-MS, inductively coupled plasma mass spectrometry; MPD, 2-methyl-2,4-pentanediol; Tris, tris(hydroxymethyl)aminomethane.

**REFERENCES**


