Substrate Interactions with the Nitrogenase Active Site

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ABSTRACT

The chemical mechanism for biological cleavage of the N_2 triple bond at ambient pressure and temperature has been the subject of intense study for many years. The site of substrate activation and reduction has been localized to a complex cofactor, called FeMo cofactor, yet until now the complexity of the system has denied information concerning exactly where and how substrates interact with the metal—sulfur framework of the active site. In this Account, we describe a combined genetic, biophysical, and biochemical approach that was used to provide direct and detailed information concerning where alternative alkyne substrates interact with FeMo cofactor during catalysis. The relevance and limitations of this work with respect to N_2 binding and reduction also are discussed.

Nitrogenase Catalyzes Biological Nitrogen

The agronomic and economic significance of nitrogen fixation, reduction of N_2 to yield $2NH_3$, can be appreciated through the perspective of an estimate that suggests that more than a third of today's human population would not

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exist without application of the synthetic Haber–Bosch process. Biological nitrogen fixation, which utilizes energy in the form of MgATP, is catalyzed by a complex metal-loenzyme called nitrogenase (see ref 2 for a recent review).

$$N_2 + 8e^- + 8H^+ + 16MgATP \Rightarrow$$

 $2NH_3 + H_2 + 16MgADP + 16Pi$

Whether the biological process can be more effectively exploited for agronomic benefit remains an open question. Nevertheless, nitrogen fixation is necessary to sustain life on earth, and how organisms manage to activate and cleave N₂ at ambient temperature and pressure remains a fascinating and unsolved chemical problem. There are three different types of nitrogenases,3 distinguished by the metal composition of their respective active site metalloclusters, but all of them share common structural and mechanistic features. The best studied of these is the Modependent enzyme that contains an active-site organometallic cluster called FeMo cofactor.⁴ All nitrogenases comprise two catalytic components, and in the case of the Mo-dependent enzyme, these are designated as the Fe protein and the MoFe protein. The Fe protein is a specific Fe₄S₄ cluster-containing nucleotide-dependent reductant of the MoFe protein, which contains the active site FeMo cofactor. Complex formation and electron transfer between the Fe protein and MoFe protein is controlled by MgATP binding and hydrolysis.⁵ The interaction of the Fe protein and MoFe protein can be considered to involve two coupled cycles. During catalysis the Fe protein Fe₄S₄ cluster cycles between 1⁺ (with 3Fe²⁺, $1Fe^{3+}$, and $4S^{2-}$) and 2^+ (with $2Fe^{2+}$, $2Fe^{3+}$, and $4S^{2-}$) states as single electrons are delivered to the MoFe protein, whereas the MoFe protein cycle involves the accumulation of multiple electrons necessary for substrate binding and reduction. Because multiple electrons are required for substrate reduction, multiple Fe protein cycles are required to complete a single MoFe protein cycle.

Nitrogenase also catalyzes the reduction of a variety of triply bonded substrates other than N_2 , the most familiar being acetylene.⁶

$$C_2H_2 + 2e^- + 2H^+ \rightarrow C_2H_4$$

In its resting state the MoFe protein is not capable of reducing or even covalently binding any substrate. Rather, substrate interaction requires the prior activation of the MoFe protein by accumulation of electrons donated from the Fe protein.⁷ In the absence of other substrates, the activated MoFe protein reduces protons to yield H₂, thereby continuously cycling the protein back to its resting state. These properties, and the fact that accumulation of a different number of electrons is required to make the

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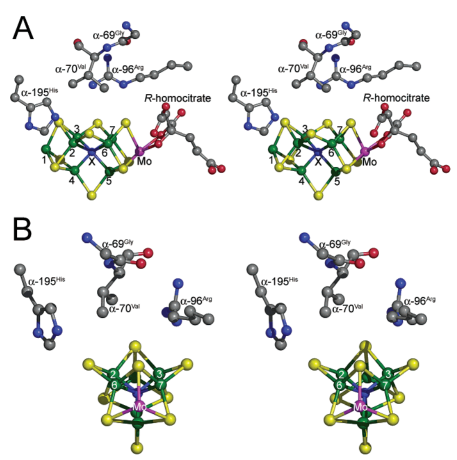


FIGURE 1. Two stereoviews of FeMo-cofactor and selected residues that provide the first shell of noncovalent interactions, a side-on stereoview (A) and a Mo end-on stereoview (with homocitrate removed) (B). FeMo cofactor is a [7Fe-8S-Mo-X-homocitrate] cluster, where X is an atom of unknown identity but is believed to be a nitride. The figure was generated from PDB file 1M1N using the program PyMOL. The color scheme is Fe in green, S in yellow, C in gray, N in blue, O in red, and Mo in magenta.

MoFe protein competent to bind different substrates, have enormously complicated analysis of nitrogenase catalysis. Perhaps the most important feature that has frustrated study of the chemical mechanism is an inability to capture a homogeneous form of the enzyme having substrate bound at high occupancy. Consequently there has been no direct information concerning where and how substrates interact with the active site. This Account describes a combination of genetic, biochemical, and biophysical strategies that were used to overcome this problem.

A Genetic Approach

The FeMo cofactor metal—sulfur core is constructed from Fe_4S_3 and Fe_4S_3Mo subclusters (Figure 1) joined by a shared central ligand, suspected to be a nitride, and linked by three bridging $S^{2-,8,9}$ An organic constituent, homocitrate, is attached to the Mo atom through its 2-hydroxyl and 2-carboxy groups, and the cofactor is covalently anchored to the MoFe protein through a N ligand provided by a histidine imidazole to the Mo atom and a cysteine thiolate ligand to an Fe atom at the opposite end. This unusual structure has invited considerable theoretical speculation concerning the identity of the substrate-binding site and description of possible mechanisms for substrate activation and also has inspired a variety of

model chemistries as approaches to understand biological $N_2\ activation.^{10}$

An initial approach to resolve the question of where substrates interact with the active site involved evaluation of the catalytic and biophysical consequences arising from substitution of the residues that provide the first shell of noncovalent interactions with FeMo cofactor.^{11–13} It soon became apparent, however, that the complexity of the system denied unequivocal interpretation of most information gained in this way. For example, for most substitutions, it was not possible to assign catalytic defects as specifically arising from electron transfer, chemical reactivity of the cofactor, or substrate access to the cofactor.

Acetylene is a potent growth inhibitor for nitrogen fixing microorganisms because N_2 and acetylene compete for the available reducing equivalents during nitrogenase catalysis. We therefore decided to use a genetic strategy to circumvent the above problems by isolation of mutant strains resistant to the inhibitory effects of acetylene. This approach was valuable for two important reasons. First, the method cannot be biased by an investigator's preference for the location of the substrate-binding site. Second, because N_2 reduction is thermodynamically more demanding and requires more electrons than acetylene reduction, a mutant strain impaired in acetylene reduction but not N_2 reduction cannot arise from defects in either

electron transfer or active site reactivity. Rather, this feature can only be assigned to an inability of acetylene to effectively access the active site.

MoFe protein purified from an acetylene-resistant strain exhibited a substantially lower affinity for acetylene binding/reduction, whereas N2 reduction parameters remained unaltered. Nucleotide sequence analysis of the genes encoding the MoFe protein from this strain showed that the MoFe protein α-subunit Gly⁶⁹ residue was substituted by serine, and inspection of the MoFe protein crystal structure revealed that α -Gly⁶⁹ is located on a short helix that skirts one of three Fe-S "faces" that compose the central portion of FeMo cofactor (Figure 1). Although the basis for acetylene resistance could be interpreted in a variety of different ways, we proposed a model where this substitution alters local conformational flexibility so that the side-chain of the adjacent α-Val⁷⁰ residue is locked into a conformation that impairs active site access by acetylene but does not affect access by the slightly smaller N₂ molecule. If this model is correct, we reasoned that substitution of α-Val⁷⁰ by residues having smaller side chains would expand the size of substrates that can access the substrate reduction site. This possibility was easily tested because previous studies had established that shortchain alkynes other than acetylene are not effectively reduced by nitrogenase, presumably because they are denied access to the activation site due to the larger size. 16,17 Indeed, shortening the side-chain at the α -Val 70 residue position, by substitution with either alanine or glycine, progressively expands the size of short-chain alkyne substrates, for example, propyne and 1-butyne, which can be accommodated at the nitrogenase active site.18

Biophysical Analysis of a Trapped Substrate Reduction Intermediate

The as-isolated form of the MoFe protein exhibits a characteristic $S = {}^{3}/{}_{2}$ electron paramagnetic resonance (EPR) spectrum that can be uniquely assigned to a semireduced state of FeMo cofactor. In this oxidation state of FeMo cofactor, termed the MN state, the formal oxidation states of the metals are not settled, results from Mossbauer spectroscopy¹⁹ indicating 1Mo⁴⁺, 3Fe³⁺, 4Fe²⁺, and 9S2- and results from electron nuclear double resonance (ENDOR) spectroscopy studies²⁰ indicating 1Mo⁴⁺, 1Fe³⁺, 6Fe²⁺, and 9S²⁻. When the MoFe protein is freezetrapped under turnover conditions, where electrons are being delivered from the Fe protein to the MoFe protein, a more reduced form of FeMo cofactor is accessed, which exists predominantly in an EPR-silent state (Figure 2). Attempts to freeze-quench the nitrogenase system in the presence of various substrates as a way to trap an EPRactive substrate/intermediate bound species has proven to be problematic. In particular, this approach has resulted in trapping bound species at only very low occupancy and often in multiple forms. For the inhibitor CO, two different CO bound states have been trapped at relatively high occupancy, one at lower CO concentrations (called lo-CO)

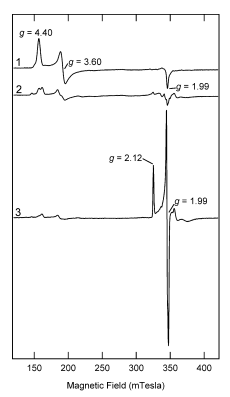


FIGURE 2. X-band EPR spectra of the MoFe protein. EPR spectra of the α -Ala⁷⁰ MoFe protein are shown for the resting state under argon (trace 1), the turnover state under argon (trace 2), and the turnover state in the presence 3 mM propargyl-OH at pH 6.7 (trace 3). The *g*-values for select inflections are shown.

and one at higher CO concentrations (called hi-CO). Characterization of these states by various spectroscopic methods has resulted in models where one or two CO molecules are bound to Fe atoms.^{20,21} Very low occupancy intermediates have also been trapped when nitrogenase is freeze-quenched during reduction of the substrates carbon disulfide and acetylene.^{22,23} The low occupancy of these intermediates has limited the information that could be deduced, although it was concluded that both substrates are likely bound to one or more Fe atoms.

Expansion of the active site of the MoFe protein so that larger substrates can be accommodated permitted the use of substrates that contain functional groups as a new strategy to trap a substrate reduction intermediate in an EPR-active form and at concentrations amenable to spectroscopic analyses.²⁴ During the course of evaluating various functionalized substrates, we discovered that when either propargyl alcohol (HC≡C-H2OH) or propargyl amine (HC=C-CH2NH2) is used as a substrate for the substituted α -Ala⁷⁰ MoFe protein, the resting state $S = \frac{3}{2}$ EPR spectrum is converted to an intense $S = \frac{1}{2}$ signal (Figure 2). In the context of earlier work identifying the S $= \frac{1}{2}$ intermediate formed during turnover in the presence of the inhibitor CO, the properties of the propargyl alcohol derived signal indicated that a single species was trapped in high occupancy.

These features provided an unprecedented opportunity to explore the properties of a bound nitrogenase substratereduction intermediate by EPR and ENDOR spectroscopies.

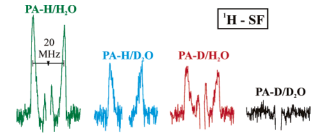


FIGURE 3. Quantitative stochastic field-modulated ENDOR spectra ($^{1}H-SF$) of the α -Ala 70 MoFe protein incubated with propargyl alcohol under turnover conditions. The deuteration patterns are indicated; spectra are centered at the ¹H frequency and split by the hyperfine coupling. The important observation from these spectra is that the intensity for the nondeuterated sample (green) is halved when either D₂O is used as solvent (blue) or the substrate is deuterated (red) and eliminated when deuterated substrate is used in D₂O (black). These results show that the bound intermediate contains two strongly coupled, magnetically identical protons. One of these protons must be from the substrate, whereas the other originates from the solvent. A third, weakly coupled proton is seen in the red and green spectra originating from the solvent. The structural interpretation of these data is shown in Figure 4.

By using uniformly ¹³C-labeled propargyl alcohol and ¹³C ENDOR spectroscopy, we could observe the coupling of three unique C atoms to FeMo cofactor.24 This result confirmed that the substrate-derived intermediate was covalently bound to FeMo cofactor and further indicated an asymmetric spin coupling of the propargyl alcohol carbon atoms (C3 > C2 > C1) to FeMo cofactor. The nature of the bound intermediate was revealed by examination of the ENDOR parameters when either H- or D-labeled propargyl alcohol (PA-H; PA-D) was used as substrate with turnover occurring in either H₂O or D₂O.²⁵ Key was the combined use of a new quantitative ¹H ENDOR technique, stochastic field-modulated (SF) EN-DOR, plus Mims-pulsed ²H ENDOR, to study strongly coupled protons (Ha) observed in the PA-H/H2O spectrum (hyperfine coupling of $A(^{1}\text{H}^{a}) \approx 20 \text{ MHz}$). This signal appears with half intensity in the spectra of the PA-H/ D₂O and PA-D/H₂O samples and is lost with the "doubly deuterated" PA-D/D₂O sample (Figure 3). These observations imply that the H^a doublet in the PA-H/H₂O spectrum is the superposition of doublets from two magnetically identical and hence symmetry-equivalent protons, one derived from propargyl alcohol (Hp) and the other acquired from solvent (H_s) during reduction. In addition, the experiments disclosed one weakly coupled proton (Hb) derived from solvent. Inspection of inorganic model compounds having similar compositions suggested two models that could explain the ENDOR observations, in particular, the two equivalent protons having different chemical origins (for a more complete discussion of the model compounds see ref 24). Both involve a threemembered ring that includes the propargyl alcohol C3 and C2 atoms and a single Fe atom. One of these is a one proton added, semireduced (ferracyclopropene) intermediate, and the other is a two proton added, further reduced (ferracylopropane) adduct of the allyl alcohol product. Between these we favor the latter (Figure 4), largely

$$H_s^b$$
 H_s^a
 H_p^a
 H_p^a
 H_p^a

FIGURE 4. Proposed structure of the trapped propargyl alcohol reduction intermediate. This structural model is based on the results of ENDOR parameters and consideration of model compounds. The figure indicates two magnetically identical protons (Ha) bound to C3, one derived from the solvent (H_s) and one derived from the substrate (H_n). The weakly coupled proton (H^b) derived from the solvent is assigned to that bound to C2. Asymmetric binding of the propargyl alcohol C3 and C2 atoms (C3 coupling > C2) as revealed by ¹³C ENDOR is indicated by a longer Fe—C2 bond than the Fe—C3 bond.

because it gives a more "natural" explanation for the presence of the solvent-derived Hb (see ref 25 for a more complete discussion). Density functional theory (DFT) calculations on FeMo cofactor subsequently showed that the proposed model is an energetically favorable one.²⁶

Elucidating the Location of the Bound Propargyl Alcohol Adduct

Genetics established the general location for binding of alkyne substrates, and advanced spectroscopic techniques provided insight on how the reduction intermediate is bound, but neither approach pinpointed which of the Fe atoms at the Fe 2, 3, 6, and 7 face is involved. To address this question, we sought to determine why a propargyl alcohol reduction intermediate becomes trapped at the active site. The important observation was that although a propargyl alcohol intermediate becomes trapped at the active site, a propyne reduction intermediate does not, even though both molecules are substrates. This suggested that the propargyl alcohol -OH group is likely to interact with a functional group provided by the protein to stabilize a reduction intermediate. Among the candidate amino acids that could provide this function, the α -His¹⁹⁵ residue was the most attractive because it is appropriately positioned within the identified binding region (Figure 1) and because previous work indicated a possible role for α -His¹⁹⁵ as an obligate proton donor during N_2 reduction. The possibility that α -His¹⁹⁵ stabilizes a propargyl alcohol reduction intermediate was tested and confirmed in two different ways.²⁷ First, it was shown that the appearance of the $S = \frac{1}{2}$ EPR signal under freeze-quench conditions when propargyl alcohol is used as substrate disappears when the α -His¹⁹⁵ residue is substituted by glutamine, even though propargyl alcohol retains an ability to interact with the active site of the substituted MoFe protein. Second, appearance of the characteristic EPR signal is pH-dependent when either propargyl alcohol or propargyl amine is used as substrate. The respective pH-dependence profiles are different and show that optimum population of the trapped propargyl alcohol intermediate occurs when the α -His¹⁹⁵ imidazole is protonated (pH 6.7, Figure 5A), whereas optimum population of the propargyl amine intermediate occurs when the imidazole group is deprotonated and propargyl amine is protonated (pH 8.0, Figure 5B).

FIGURE 5. Proposed H-bonding of the trapped intermediates. The proposed H-bonding (dashed lines) is shown between the protonated α -His¹⁹⁵ and propargyl alcohol (A) and between the deprotonated α -His¹⁹⁵ and protonated propargyl amine (B).

The above results indicated formation of a hydrogen bond between the -OH group of a propargyl alcohol reduction intermediate and the imidazole of α -His¹⁹⁵ thereby localizing the position of the -OH group within ~ 2 Å of the ϵN of α -His¹⁹⁵. Furthermore the previously described ENDOR studies showed that the bound adduct is allyl alcohol, which is proposed to interact with a single Fe atom in an η^2 configuration. Within these constraints, the location of the bound intermediate is largely defined. To further refine the likely binding site, DFT computational methods were used to suggest a detailed bonding geometry of the cofactor—reduction intermediate adduct, which was then fit into the α -Ala⁷⁰ MoFe protein using force field methods to give the model shown in Figure 6.

Relevance of the Identified Alkyne Binding Site to N_2 Binding

The results of experiments described so far provide compelling evidence that alkyne substrates can bind and be reduced at a specific FeS face of FeMo cofactor and perhaps at a single Fe atom within that face. A remaining question is whether N_2 is also activated and reduced at this same face. This question is an important one because acetylene is a noncompetitive inhibitor of N_2 reduction whereas N_2 is a weak competitive inhibitor of acetylene reduction. Also, there is abundant evidence that acetylene can bind to the MoFe protein with both high and low affinity. Reasonable interpretations of these observations are either that there are multiple and separate binding sites or that a single site can be accessed at

different redox states. Differentiation between these possibilities, which are not mutually exclusive, is experimentally challenging because the MoFe protein pool is populated by a variety of different redox states under turnover conditions.

Although not yet answered unequivocally, there are recent results that bear on the question of whether acetylene and N₂ share the same binding site. In experiments that have already been described, it was possible to expand the substrate reduction site by substitution of the α-Val⁷⁰ residue by amino acids having smaller side chains. We therefore reasoned that substitution of α-Val⁷⁰ by amino acids having larger side chains should compromise the reduction of all substrates that must access this face. To test this possibility, the $\alpha\text{-Val}^{70}$ residue was substituted by isoleucine, and the impact on acetylene and N₂ reduction was evaluated.²⁹ The results of these experiments showed that the substituted α-Ile⁷⁰ MoFe protein is severely compromised for both acetylene reduction (>120-fold increase in K_m) and N_2 reduction (>15-fold increase in $K_{\rm m}$) with no effect on proton reduction. The observations that proton reduction is not affected by the substitution and that acetylene and N2 reduction can still be detected, although at very low levels, are consistent with an interpretation that neither electron delivery to the active site nor the ability of the enzyme to access the redox state required for substrate binding have been affected. Although an effect of the substitution on N2 activation has not been ruled out, a logical interpretation of the results is that the α-Ile70 substitution denies access of both N2 and acetylene to the same or overlapping binding sites. Whatever the relationship between N₂ binding and acetylene binding, the effect of the α -Ile⁷⁰ substitution indicates that all acetylene reduction reactions must occur at the same face for several reasons. First, both acetylene and proton reduction involve the same redox states of the MoFe protein, both requiring two electrons. Second, proton reduction is unaffected by the α -Ile⁷⁰ substitution, which means that the capacity for the altered MoFe protein to access the redox states required for acetylene reduction is not affected by the substitution. Third, the ability of the substituted protein to catalyze acetylene reduction at substrate concentrations used to saturate the

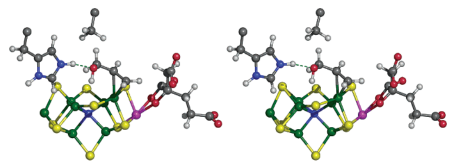


FIGURE 6. Stereoview of the proposed structure for the trapped propargyl alcohol reduction intermediate bound to FeMo cofactor. The most favorable state of the propargyl alcohol reduction intermediate bound to FeMo cofactor in the α -Ala⁷⁰ MoFe protein is shown. The alkane unit of allyl alcohol is bound to Fe6 of the FeMo cofactor. A H-bond is shown (dashed line) between the hydroxyl 0 of allyl alcohol and the ϵ N of α -His¹⁹⁵ in the α -Ala⁷⁰ MoFe protein. This model was deduced from experimental evidence for hydrogen bonding between the -OH group of the reduction intermediate and the α -His¹⁹⁵ side-chain imidazole group and a combination of density functional theory and molecular mechanics computational methods.

normal enzyme is virtually eliminated. Of course, the possibility that multiple substrate sites for acetylene binding exist within the same FeS face has not been ruled out.

A second line of experiments that indicates N2 and acetylene share the same or overlapping binding sites involves analysis of the semireduced form of N2, hydrazine, as a substrate.²⁹ It was previously shown that hydrazine is a very poor substrate for nitrogenase.³⁰ Further, acid quenching of nitrogenase under turnover conditions results in the release of small amounts of hydrazine when N₂ is used as substrate.³¹ Hydrazine is also a minor reduction product when N₂ is used as a substrate for the vanadium-dependent nitrogenase.32 In aggregate, these observations provide compelling evidence that a bound form of hydrazine is an intermediate during nitrogenasecatalyzed N₂ reduction. One possible explanation for the slow rate of hydrazine reduction when used as a substrate is that its relatively larger size, compared to N2, denies facile access to the active site. If this explanation is correct and N2 is reduced at the same FeMo cofactor face as acetylene reduction, then expansion of the active site by the α -Ala⁷⁰ substitution should increase the capacity for hydrazine reduction. This prediction was verified experimentally where it was shown that substitution of the α -Val⁷⁰ residue by alanine dramatically increases the capacity for hydrazine reduction (8-fold decrease in $K_{\rm m}$).²⁹

The Riddle of N2 Activation Is Not Solved

Despite nearly 40 years of intense investigation, the chemical mechanism of biological nitrogen reduction has remained enigmatic. In this Account, we have described a comprehensive approach that has been used to gain evidence that nitrogenase substrates bind and are reduced at a specific FeS face of FeMo cofactor. Work described here cannot be considered to have eliminated Mo as the activation site for N_2 reduction, but none of our results are compatible with that model. Perhaps the most important contribution of the current work is that it has provided a framework for future theoretical models and proposed chemical mechanisms that can be tested experimentally.

From the experimental perspective, there are two immediate goals of our future research efforts. There is now strong evidence that binding of hydrazine, a likely intermediate during N2 reduction, occurs at the same general site as that identified for propargyl alcohol. However, exactly where and how hydrazine is bound at the active site when used as a substrate is not known. To address this question, we are attempting to trap an EPRactive N₂ reduction intermediate in a highly populated state, using the same general approach described for detection and characterization of an alkyne reduction intermediate. Our second goal is related to the activation of N₂. Although nitrogenase is able to reduce a variety of substrates, N2 reduction is differentiated from other substrates in the following ways: (i) a more reduced state of the MoFe protein is required for N2 binding, (ii) N2 binding/reduction is associated with the obligate evolution of H_2 , (iii) H_2 is a competitive inhibitor of N_2 reduction, and (iv) in the presence of N_2 and D_2 nitrogenase catalyzes HD formation.⁴ These observations suggest that the N_2 activation, but not activation of other substrates, could involve the obligate displacement of hydrides bound to FeMo cofactor. Validation of this model requires elucidation of whether hydrides are bound to FeMo cofactor in the activated state and, if so, determination of where they are bound. Again we believe that a combined genetic and biochemical approach as a way to trap a highly populated, EPR-active state so that advanced spectroscopies can be applied provides the best opportunity for success. These experiments are in progress.

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