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# X-ray Crystallography and Vibrational Spectroscopy Reveal the Key Determinants of Biocatalytic Dihydrogen Cycling by [NiFe] Hydrogenases

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Abstract: [NiFe] hydrogenases are complex model enzymes for the reversible cleavage of dihydrogen  $(H_2)$ . However, structural determinants of efficient H<sub>2</sub> binding to their [NiFe] active site are not properly understood. Here, we present crystallographic and vibrational-spectroscopic insights into the unexplored structure of the  $H_2$ -binding [NiFe] intermediate. Using an F<sub>420</sub>-reducing [NiFe]-hydrogenase from Methanosarcina barkeri as a model enzyme, we show that the protein backbone provides a strained chelating scaffold that tunes the [NiFe] active site for efficient  $H_2$  binding and conversion. The protein matrix also directs  $H_2$  diffusion to the [NiFe] site via two gas channels and allows the distribution of electrons between functional protomers through a subunit-bridging FeS cluster. Our findings emphasize the relevance of an atypical Ni coordination, thereby providing a blueprint for the design of bio-inspired  $H_2$ -conversion catalysts.

#### Introduction

Mapping out strategies for future energy storage and conversion represents one of the central challenges of the  $21^{st}$  century. Molecular hydrogen (H<sub>2</sub>) is an ideally clean fuel

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whose combustion releases large amounts of free energy but no greenhouse gases. To exploit it to its full potential, however, we require efficient and sustainable approaches for catalytic H<sub>2</sub> cleavage and formation. [NiFe] hydrogenases are valuable model enzymes that catalyze H<sub>2</sub> conversion at rates comparable to platinum electrodes by using a heterobimetallic active site containing cheap and earth-abundant base metals only.<sup>[1]</sup> Their rational utilization as biotechnological targets or blueprints for bio-inspired catalysts, however, requires a thorough understanding of the structural and mechanistic determinants of their reactivity. Here, we employ a bidirectional F<sub>420</sub>-reducing [NiFe] hydrogenase from the archaeon Methanosarcina barkeri MS (MbFRH) as a unique model system to yield structural and spectroscopic insights into a scarcely explored reaction intermediate that is the initial target for  $H_2$  binding to the active site of the enzyme. The structure of this central catalytic species is analyzed in detail to explore its relevance for the mechanism and performance of [NiFe] hydrogenases.

## **Results and Discussion**

The crystal structure of MbFRH was refined using reflections up to  $d_{\min} = 1.84 \text{ Å}$  (Supporting Information, Table S1). Each asymmetric unit contains three subunits, FRH-A, FRH-B, and FRH-G, and a total of four [4Fe4S] clusters as well as a flavin adenine dinucleotide (FAD) cofactor and the [NiFe] active site (detailed below). The FAD and the heterobimetallic [NiFe] center enable the redox conversion of the two substrates, coenzyme  $F_{420}$  and  $H_2$ , respectively, while the [4Fe4S] clusters mediate intramolecular electron transfer (ET) between the two reaction sites (Figures 1 A and S1). These features are shared with the related [NiFe] hydrogenase from Methanothermobacter marburgensis (Mm), and both enzymes exhibit a dodecameric overall architecture in a spherical shape (Figure S1).<sup>[2]</sup> Compared to the latter enzyme, however, MbFRH contains two additional cofactors that are both ET-accessible and solventexposed: a [2Fe2S] cluster bridging two FRH-G subunits (Figures 1 A and S2) and a mononuclear Fe site in FRH-A (Figures 1 A and S3).

Based on element-specific anomalous scattering, the single Fe ion has also been assigned in [NiFeSe] hydrogenases,<sup>[3]</sup> while  $Ca^{2+[4]}$  or  $Mg^{2+[5]}$  ions are typically found at the same position in other [NiFe] hydrogenases (Table S2). The [2Fe2S] cluster likely contributes to the biological

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**Figure 1.** A) Reversible ET (indicated by arrows) between the FAD and the [NiFe] active site is enabled by a chain of [4Fe4S] clusters. The [NiFe] center could also exchange electrons with a mononuclear Fe site (upper inset), and the presence of a [2Fe2S] cluster (lower inset) allows electrons to commute between antiparallel ET chains of two heterotrimers. Distances given in Å; selected atoms and amino acids shown as spheres (Fe: orange, S: yellow, Ni: green) and sticks, respectively. B) EPR spectra of an *Mb*FRH solution recorded at 80 K with 1 mW microwave power and 9.3 GHz microwave frequency. Signals from FAD and a [2Fe2S] cluster are highlighted in dark cyan and violet, respectively.

function of *Mb*FRH by enabling the distribution of electrons among individual protomers of the dodecameric enzyme. In line with this proposal, solution-phase electron paramagnetic resonance (EPR) experiments reveal signals from FAD, [2Fe2S] clusters (Figures 1B and S4), and [4Fe4S] clusters (Figure S4), corroborating the extended electron-transfer pathway of native *Mb*FRH.

In the following, we will focus on the [NiFe] active site of MbFRH, which exhibits the consensus structural properties observed for other [NiFe] hydrogenases.<sup>[6]</sup> Specifically, this heterobimetallic cofactor features two metal ions, Ni and Fe, that are coordinated by four strictly conserved cysteinate (Cys) residues and three Fe-bound diatomic ligands. Based on infrared (IR) spectroscopic analyses, the latter constituents are generally assigned to one CO and two CN<sup>-</sup> ligands.<sup>[7]</sup> Consistently, one CO and two CN stretching bands can be observed in IR spectra of MbFRH crystals, which confirms the presence of a standard set of inorganic ligands (Figure 2A, black trace). Since these stretching vibrations are

highly sensitive towards structural and electronic changes at the [NiFe] center,<sup>[8]</sup> the observation of a single set of three IR absorption bands in the relevant spectral range also demonstrates that the active site resides in a homogenous redoxstructural state. In contrast to crystal structures of many other [NiFe] hydrogenases, no electron density can be detected in the third bridging position between the two metals for MbFRH (Figure 2B).<sup>[6a-c]</sup> This excludes the presence of oxygen-containing ligands that would be indicative of inactive enzyme residing, for example, in the Ni<sub>u</sub>-A or Ni<sub>r</sub>-B states.<sup>[6a,9]</sup> Notably, a vacant third bridging site between the two metals has long been assumed to be a key feature of Nia-S (also termed Ni-SIa), the H<sub>2</sub>-binding intermediate of [NiFe] hydrogenases.<sup>[6a, 10]</sup> This assumption has been recently supported by spectroscopic analyses,<sup>[11]</sup> but detailed structural insights into this central catalytic intermediate have been so far unavailable. In the following, we will shed light on these catalytic key aspects by using a joint approach of crystallographic analysis and vibrational spectroscopy.

While the presence or absence of bridging oxygen ligands can be clearly established from crystal-structure analyses, H<sub>2</sub>derived (hydride) ligands are only observable in sub-atomicresolution structures,<sup>[12]</sup> and available crystal structures lacking detectable bridging ligands most likely represent hydride species.<sup>[9e, 13]</sup> Since crystals of *Mb*FRH have been prepared in a mildly reducing atmosphere containing up to 5% H<sub>2</sub>, the derived structural data could reflect Ni<sub>a</sub>-S or a hydride species. Thus, before analyzing the active-site structure of *Mb*FRH in more detail, we studied the underlying crystals using different vibrational-spectroscopic techniques to firmly identify the [NiFe] redox-structural state.

IR spectra of MbFRH crystals reveal CO and CN stretching bands at 1945 and 2065/2080 cm<sup>-1</sup>, respectively, which resemble the IR fingerprint of the Nia-S state of several [NiFe] hydrogenases (Figure 2A).<sup>[6a]</sup> In particular, these vibrational frequencies are close to those observed for the as-isolated regulatory hydrogenase from Ralstonia eutropha (ReRH), which is a spectroscopically valuable yet structurally unexplored reference system for the Ni<sub>2</sub>-S state.<sup>[11]</sup> Based on this finding, contributions from the Ni<sub>2</sub>-C hydride intermediate and its photo-inducible congener, Ni<sub>a</sub>-L, can be excluded since these catalytic species would give rise to higher and lower CO stretching frequencies, respectively.<sup>[6a]</sup> The fully reduced Ni<sub>a</sub>-SR state, however, cannot be excluded on the basis of IR-spectroscopic data alone since its dominating subspecies exhibits an IR fingerprint similar to that of Ni<sub>a</sub>-S in several [NiFe] hydrogenases.<sup>[6a]</sup> Thus, we next recorded resonance Raman (RR) spectra of MbFRH crystals to probe Fe-CO and Fe-CN metal-ligand vibrations as structural markers of the [NiFe] active site (Figure 2C).<sup>[11b,14]</sup> In these measurements, Ni<sub>a</sub>-SR would be photo-converted to Ni<sub>a</sub>-L, while Ni<sub>a</sub>-S would remain unaffected in terms of structural and electronic properties.<sup>[11b,14]</sup> Again, the obtained vibrational signature is very similar to that of the Ni<sub>a</sub>-S state as observed for ReRH and a membrane-bound hydrogenase from the same organism (ReMBH).[11b,14b] In particular, a structurally sensitive vibrational mode with considerable Fe–CO stretching character can be detected at 596–598 cm<sup>-1</sup>, which excludes contributions from Ni<sub>a</sub>-SR, since its photo-

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**Figure 2.** A) IR spectra of an *Mb*FRH single crystal (at 80 K, black) and a protein solution of *Re*RH (10 °C, gray). Spectra were normalized with respect to the CO stretching-band intensity. B) Crystal structure of the [NiFe] active site, exhibiting a distorted seesaw coordination geometry of the Ni(Cys)<sub>4</sub> moiety and a vacant coordination site between the Ni and Fe ions. The  $2F_{obs}-F_{calc}$  electron density map after full refinement is shown as a gray mesh (1.8  $\sigma$ ). C) RR spectrum of an *Mb*FRH single crystal (black) compared to that of a protein solution of *Re*RH.<sup>[11b]</sup> RR spectra were recorded at 80 K using 568-nm laser excitation and normalized with respect to the most intense signal at 551/553 cm<sup>-1</sup>. D) [NiFe] active-site crystal structure. Selected interatomic distances are given in Å. E) IR absorbance spectrum of an *Mb*FRH single crystal, recorded at 80 K. Bands corresponding to the intrinsic, Fe-bound diatomic ligands and the extrinsic, Ni-bound CO of the Ni-SCO redox-structural state are highlighted in brown and orange, respectively. F) X-ray structure of the CO-inhibited [NiFe] active site. The  $2F_{obs}-F_{calc}$  electron density map after full refinement (1  $\sigma$ ) and the residual  $F_{obs}-F_{calc}$  map (5.5  $\sigma$ ) before CO-modeling are shown as a gray mesh and green surface, respectively. Selected atoms and amino acid residues are shown as spheres (Ni in green, Fe in gray (B) and orange (D and F), S in yellow) and sticks, respectively.

product Ni<sub>a</sub>-L would give rise to an Fe–CO stretching frequency above 600 cm<sup>-1</sup>.<sup>[11b,14a]</sup> Moreover, the intensity of the RR spectrum was found to increase with the excitation wavelength of the Raman probe laser, which is in line with previous observations for Ni<sub>a</sub>-S<sup>[11b,14]</sup> and contrary to expectations for reduced [NiFe] hydrogenases.<sup>[11b]</sup>

Finally, we also explored the effect of treating *Mb*FRH crystals with CO, a typical inhibitor of [NiFe] hydrogenases.<sup>[6a,15]</sup> We performed these experiments to check for interaction of CO with the [NiFe] active site, which is expected for Ni<sub>a</sub>-S (yielding Ni-SCO) but not for Ni<sub>a</sub>-C or Ni<sub>a</sub>-SR.<sup>[15d]</sup> Binding of extrinsic CO to the terminal vacant coordination site at the Ni ion of *Mb*FRH is evident from the electron density at this position (Figures 2F and S6) and a high-frequency CO stretching band at 2048 cm<sup>-1</sup> in the corresponding IR spectrum (Figure 2E), as also observed for other [NiFe] hydrogenases in the Ni-SCO state.<sup>[15a,c,d]</sup> Both structural and spectroscopic data show that this inhibited species accumulated to at least 50% (Figure S6), while the remainder can be assigned to the Ni<sub>a</sub>-S parent state. In total, the above experiments show that the crystal structure of untreated *Mb*FRH reflects a homogenous Ni<sub>a</sub>-S state, allowing a detailed analysis of this  $H_2$ -binding intermediate.

Structural and electronic properties of Ni<sub>a</sub>-S have been proposed to be essential for efficient H<sub>2</sub> binding and hydride formation in [NiFe] hydrogenases.<sup>[11b,16]</sup> In the following, we will revisit these proposals to evaluate their validity based on the crystal structure. While there is wide agreement regarding the overall catalytic mechanism of [NiFe] hydrogenases, details about the central steps of H<sub>2</sub> binding and activation are, so far, elusive. In particular, the site of initial H<sub>2</sub> binding is not known, and either of the two metal ions may be involved in the formation of a (side-on) H<sub>2</sub>  $\sigma$ -bond complex from Ni<sub>a</sub>-S. Experimental data on this first catalytic step are not yet available, but recent theoretical studies favor the Ni ion as the initial site of H<sub>2</sub> binding.<sup>[16-17]</sup> According to these studies, the coordination geometry of this metal ion represents a key to the energetically favorable interaction of H<sub>2</sub> with the [NiFe]

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active site. Specifically, a peculiar seesaw-shaped geometry with *trans* S–Ni–S angles approaching 120° and 180° was postulated to be mandatory for thermodynamically favorable binding of H<sub>2</sub> to Ni<sub>a</sub>-S.<sup>[16]</sup> In line with this proposal, the [NiFe] active site of *Mb*FRH exhibits *trans* S–Ni–S angles of 107° and 171°, thereby structurally confirming this unusual geometry of the Ni<sub>a</sub>-S state (Figure 2B). This finding also agrees with previous RR studies on *Re*RH,<sup>[11b]</sup> highlighting the merit of combining crystallographic, spectroscopic, and theoretical methods.

Notably, four-coordinate Ni<sup>II</sup> sites, as found in Ni<sub>a</sub>-S, typically exhibit (distorted) square-planar or tetrahedral coordination geometries. This indicates that the atypical seesaw geometry of this [NiFe] intermediate is dictated by the four-cysteinate coordination pattern and the protein matrix of [NiFe] hydrogenases.<sup>[16,18]</sup> Remarkably, trans S-Ni-S angles in the Ni<sub>a</sub>-S state of MbFRH resemble those found in computationally optimized Nia-S geometries of previous theoretical studies (124° and 151°), but comparison with the underlying crystal-structure-derived starting geometries (109° and 166°) yields an even better agreement.<sup>[16]</sup> This indicates that the magnitude and relevance of the structural constraints imposed by the protein matrix is even more pronounced than previously anticipated. Further evidence for the relevance of structural constraints comes from comparing the experimental Ni-Fe distance of Ni<sub>a</sub>-S to values obtained in theoretical studies. Calculations on small- to medium-size models typically report Ni-Fe distances of up to 3.3 Å, [6a, 11b, 19] while models including larger parts of the protein matrix yield smaller values,<sup>[19]</sup> close to those we observe in the crystal structure (2.7 Å; Figure 2D). This indicates that the protein matrix compresses the Ni-Fe distance to values close to those observed for other catalytic intermediates, including hydride species, Ni<sub>a</sub>-C and Ni<sub>a</sub>-SR (2.6 Å),<sup>[12]</sup> and presumed metal-H<sub>2</sub> adducts (2.6-2.8 Å).<sup>[16,17b]</sup> This effect likely minimizes structural reorganization during H2 turnover, thereby adding to the remarkable catalytic efficiency of [NiFe] hydrogenases.[11b] Additionally, a short Ni-Fe distance may also be relevant for metal-metal bond formation, as previously proposed for catalytic intermediates of [NiFe] hydrogenases.<sup>[20]</sup> Remarkably, Ni-Fe distances obtained from large computational models only reproduce the experimental value observed for MbFRH if a low-spin (S=0) configuration of the [NiFe] active site is assumed in the calculations,[16,19] supporting a singlet ground state of Ni<sub>a</sub>-S in *Mb*FRH.

To further explore the initial interaction of *Mb*FRH with  $H_2$ , we also investigated possible intramolecular  $H_2$  transfer pathways. To this end, *Mb*FRH crystals were derivatized with xenon to explore hydrophobic gas channels connecting the exterior of the enzyme with the [NiFe] active site (Figures 3 and S7 A). These experiments revealed a second  $H_2$ -transfer channel that has not been observed in similar studies on other [NiFe] hydrogenases.<sup>[21]</sup> Surprisingly, the unrelated [NiFeSe] hydrogenase from *Desulfovibrio vulgaris* Hildenborough features a similar hydrophobic tunnel (Figure S7B), indicating analogous developments in certain hydrogenases that operate bidirectionally in vivo.<sup>[22]</sup> While this narrow channel appears obstructed in crystal structures of other [NiFe] hydrogenases (Figure S7B),  $^{[21n,b]}$  dynamically enhanced H<sub>2</sub>



*Figure 3.* Seven Xe atoms (shown as red spheres) were detected within the noncanonical channel cutting through the FRH-G subunit. FRH-A, FRH-B, and FRH-G are represented by surfaces and colored navy blue, violet, and green, respectively.

transfer via this route remains as a far-reaching possibility and a target for future studies.

#### Conclusion

In the current account we have experimentally explored the unique structural properties of the H<sub>2</sub>-binding catalytic intermediate of [NiFe] hydrogenases. Besides revealing additional electron and H<sub>2</sub> pathways, the spectroscopically validated crystal structure of a pure Nia-S state unraveled two key determinants of efficient H<sub>2</sub> cycling: a peculiar seesaw-shaped coordination geometry of the Ni ion and a short Ni-Fe distance that is indicative of a low-spin electronic ground state. Both structural observations contradict expectations for unconstrained low-molecular-weight transition-metal compounds, thereby illustrating the central role of chelating ligand scaffolds and outer coordination layers in biocatalytic H<sub>2</sub> cycling. These findings expand our understanding of [NiFe] hydrogenases and, thus, provide valuable guidance for the future design of bio-inspired catalysts for H<sub>2</sub>-based energy-conversion approaches.

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## Conflict of interest

The authors declare no conflict of interest.

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## **Research Articles**

## Biocatalysis

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X-ray Crystallography and Vibrational Spectroscopy Reveal the Key Determinants of Biocatalytic Dihydrogen Cycling by [NiFe] Hydrogenases



Where does the  $H_2$  go? Using an  $F_{420}$ reducing [NiFe]-hydrogenase from *Methanosarcina barkeri* as a model enzyme, it is shown that the protein backbone provides a strained chelating scaffold that tunes the [NiFe] active site for efficient  $H_2$ binding and conversion. The protein matrix also directs  $H_2$  diffusion to the [NiFe] site via two gas channels and allows the distribution of electrons between functional protomers through a subunit-bridging FeS cluster.