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Protonation/reduction dynamics at the [4Fe–4S] cluster of the hydrogen-forming cofactor in [FeFe]-hydrogenases†

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The [FeFe]-hydrogenases of bacteria and algae are the most efficient hydrogen conversion catalysts in nature. Their active-site cofactor (H-cluster) comprises a [4Fe–4S] cluster linked to a unique diiron site that binds three carbon monoxide (CO) and two cyanide (CN[−]) ligands. Understanding microbial hydrogen conversion requires elucidation of the interplay of proton and electron transfer events at the H-cluster. We performed real-time spectroscopy on [FeFe]-hydrogenase protein films under controlled variation of atmospheric gas composition, sample pH, and reductant concentration. Attenuated total reflection Fourier-transform infrared spectroscopy was used to monitor shifts of the CO/CN[−] vibrational bands in response to redox and protonation changes. Three different [FeFe]-hydrogenases and several protein and cofactor variants were compared, including element and isotopic exchange studies. A protonated equivalent (**HoxH**) of the oxidized state (**Hox**) was found, which preferentially accumulated at acidic pH and under reducing conditions. We show that the one-electron reduced state **Hred'** represents an intrinsically protonated species. Interestingly, the formation of **HoxH** and **Hred'** was independent of the established proton pathway to the diiron site. Quantum chemical calculations of the respective CO/CN[−] infrared band patterns favored a cysteine ligand of the [4Fe–4S] cluster as the protonation site in **HoxH** and **Hred'**. We propose that proton-coupled electron transfer facilitates reduction of the [4Fe–4S] cluster and prevents premature formation of a hydride at the catalytic diiron site. Our findings imply that protonation events both at the [4Fe–4S] cluster and at the diiron site of the H-cluster are important in the hydrogen conversion reaction of [FeFe]-hydrogenases.

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Introduction

Hydrogenases are enzymes that catalyze hydrogen conversion in archaea, bacteria, fungi, and algae.¹ Among the three types of hydrogenases only [FeFe]-hydrogenases are found in eukaryotes, for example HYDA1 of the unicellular green alga

Chlamydomonas reinhardtii.^{1–3} HYDA1 represents the ‘minimal unit’ of biological hydrogen conversion as it exclusively binds the catalytic cofactor.³ Prokaryotic [FeFe]-hydrogenases carry additional iron–sulfur clusters and typically stem from obligate anaerobic bacteria such as *Clostridium pasteurianum* (CPI) or *Desulfovibrio desulfuricans* (DDH).^{4–7} [FeFe]-hydrogenases are bidirectional enzymes that catalyze H₂ oxidation and H₂ release from proton reduction at negligible electrochemical overpotential.^{8–10} Hydrogen conversion in [FeFe]-hydrogenases was proposed to be limited by the diffusion velocity of reactants^{11–14} and understanding the molecular principles of hydrogen conversion in [FeFe]-hydrogenases will be an important step towards the design of bio-inspired catalysts with comparable H₂ release activity.^{15–17}

The active site cofactor of [FeFe]-hydrogenases is referred to as the H-cluster (Fig. 1). It comprises a [4Fe–4S] cluster covalently linked to a unique diiron site with a ‘proximal’ and ‘distal’ iron ion (Fe_p and Fe_d, respectively).⁶ An azadithiolate ligand (adt) bridges the two iron ions and introduces a

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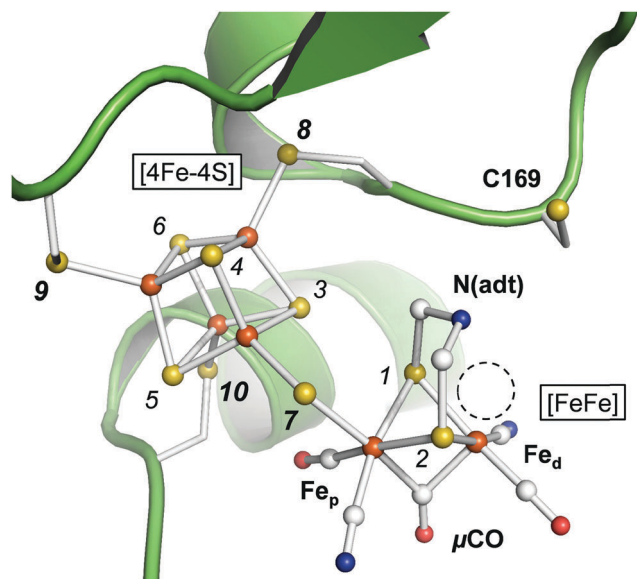


Fig. 1 Active site and catalytic cofactor in [FeFe]-hydrogenases. The crystal structure corresponds to the active-ready, oxidized state (**Hox**) of bacterial [FeFe]-hydrogenase CPI (PDB entry 4XDC).⁶ The catalytic cofactor comprises a [4Fe-4S] cluster and a diiron site, [FeFe]. Fe_p and Fe_d – proximal and distal iron ions. μCO–Fe–Fe bridging CO ligand. N(adt) – nitrogen headgroup of the adt ligand. The dashed circle highlights an apical vacancy at Fe_d. C169 is a conserved cysteine residue that couples the H-cluster to the proton transfer pathway. Sulfur atoms are numbered 1–10 according to the DFT model (see the ESI†).

potential nitrogen base in close proximity to Fe_d.^{18–20} The metal ligand sphere is complemented by a terminal carbonyl (CO) and cyanide (CN[−]) ligand at Fe_p and Fe_d. In the following, pCO/CN[−] and dCO/CN[−] refer to the terminal ligands in the proximal or distal position. A third carbonyl ligand is found in the Fe–Fe bridging position (μCO), for example in the active-ready, oxidized state of the cofactor (**Hox**).^{21–23} Crystal structures have shown that Fe_d exhibits a vacant, apical coordination site in **Hox** (Fig. 1). Binding of a fourth CO ligand in the CO-inhibited state (**Hox-CO**)^{24–27} has recently been proposed to result in an octahedral Fe_d site with an apical CN[−] ligand.²⁸ Several further redox species were identified (Table 1). Two distinct one-electron reduced states have been described, **Hred** and **Hred'**, which differ in the location of the surplus electron at the H-cluster.^{29–31} In the two-electron 'super-reduced' state, **Hsred**, μCO adopts a terminal, non-bridging position³² and a bridging hydride (μH)

was suggested.^{33,34} Reorientation of μCO has been discussed also for **Hred**.^{22,31,33–36} In amino acid variants of HYDA1, another two-electron reduced species has been observed. **Hhyd** carries a μCO ligand and was suggested to bind a terminal hydride in the apical position at Fe_d.^{37–40} It needs to be clarified whether and how the various redox species are involved in the catalytic cycle.^{41–45}

Exchange of protons between the [FeFe]-hydrogenase active site and the aqueous solvent is supported by a chain of phylogenetically conserved amino acids. In HYDA1, the putative proton transfer pathway comprises residues E144, S189, E141, C169, and at least one water molecule.^{50–52} Side-directed mutagenesis studies identified C169 as the primary proton relay to the diiron site of the H-cluster⁵¹ and proton transfer is further facilitated by the N(adt) group (Fig. 1).^{18–20,35} Theoretical studies support a rapid exchange of protons *via* this 'catalytic' pathway.^{53–55} However, the interplay between proton transfer and electron transfer events in the hydrogen conversion reaction of [FeFe]-hydrogenases remains largely unclear.

Here, we describe *in situ*, real-time spectroscopy on [FeFe]-hydrogenase protein films under controlled variation of gas composition, sample pH, and reductant concentration. Shifts of the CO/CN[−] vibrational bands were used to study electron and proton transfer at the active site *via* attenuated total reflection Fourier-transform infrared spectroscopy (ATR FTIR). Vibrational band patterns were characterized for three different [FeFe]-hydrogenases, proteins with amino acid exchanges in the proton path, and five cofactor variants comprising alternative dithiolate groups. Redox and pH titrations, element and isotope exchange studies, and density functional theory (DFT) calculations identified novel H-cluster species, most likely protonated at the [4Fe-4S] cluster.

Experimental

Protein sample preparation

[FeFe]-hydrogenase HYDA1 and CPI apo-proteins were over-expressed in *Escherichia coli*, purified, quantitatively reconstituted *in vitro* with synthetic diiron complexes (Fe₂(x)(CO)₄(CN)₂), and concentrated to 500 μM protein (~25 g L^{−1}).²⁰ The typical H₂ release activity was ~800 μmol H₂ mg^{−1} HYDA1^{adt} min^{−1} and ~2500 μmol H₂ mg^{−1} CPI^{adt} min^{−1}. Analyzed cofactor variants included adt = (SCH₂)₂NH, odt = (SCH₂)₂O, sdt = (SCH₂)₂S, pdt = S₂(CH₂)₃, or edt = S₂(CH₂)₂ as bridging dithiolate groups (x) at the diiron site. Site-directed mutagenesis of HYDA1 at position 169 yielded variants C169D and C169A as described earlier.⁵⁰ Selenium substitution at the [4Fe-4S] cluster was performed as recently published.⁵⁶ Wildtype DDH enzyme was purified from *D. desulfuricans* in a functional form.⁵⁷ All protein preparation and handling procedures were carried out at room temperature, under strictly anoxic conditions, and dim light.

Fourier-transform infrared spectroscopy

Fig. 2 shows a schematic drawing of the experimental ATR FTIR set-up for exposure of protein films to an aerosol with defined humidity, pH, and reactant concentration. In section (A), gasses

Table 1 Electronic configuration of different redox species

Redox species	Δe [−]	[4Fe-4S] cluster	Diiron site ^a	Alternative annotation	Ref.
Hox	—	+2	I/II	—	46–48
Hox-CO	—	+2	I/II	—	25–28
Hred'	1	+1	I/II	H'red', Hred	29, 31 and 49
Hhyd	2	+1	II/II	—	37–40
Hred	1	+2	I/I	HredH ⁺	21, 22 and 31
Hsred	2	+1	I/I	—	32 and 33

^a Formal charge is given for Fe_p/Fe_d. Only the mixed-valence state I/II is EPR-active.



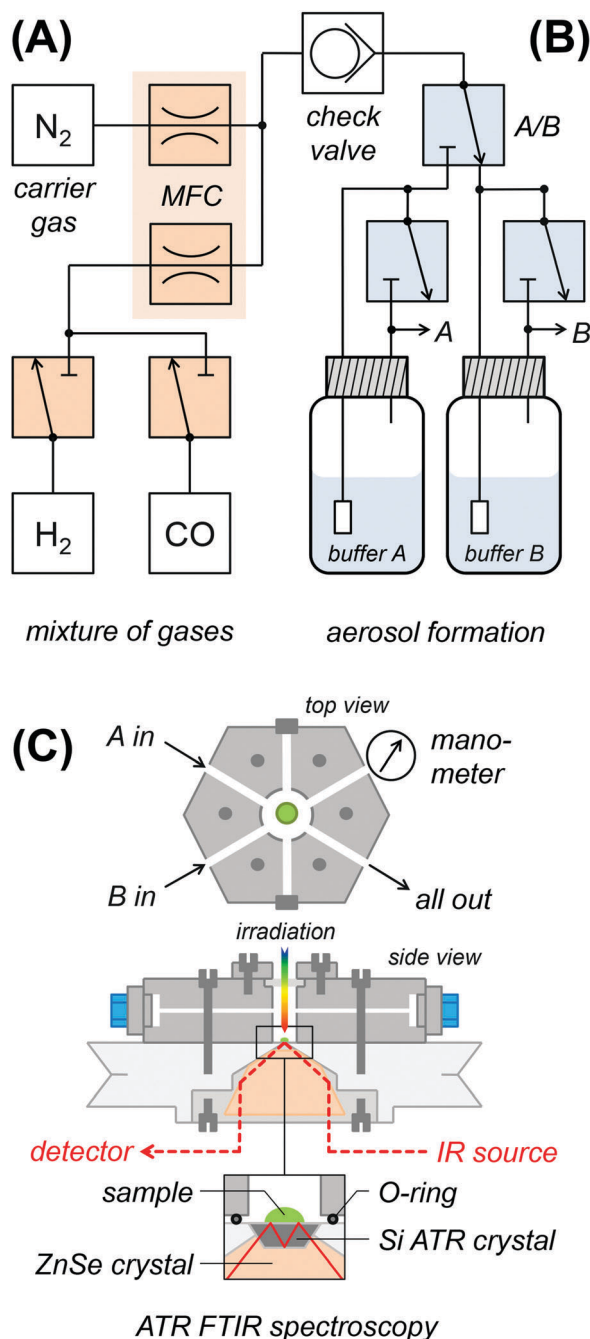


Fig. 2 Experimental set-up for ATR FTIR on protein films. (A) Various gases (e.g. H_2 , CO) were mixed into a constant stream of N_2 carrier gas. The gas flow was controlled by digital mass flow controllers (MFC). (B) Dry gas was passed through either wash bottle A or B, comprising different buffer compositions. (C) The thereby created aerosol was fed to the headspace above the protein film. On top of the ATR reflection unit, a custom-made PCTFE chamber with gas in- and outlets and manometric pressure control capped the silicon crystal and the protein film.

are mixed and the flow volume is adjusted. Dry N_2 carrier gas of ultra-high purity (5.0) was provided by a nitrogen generator (PN1450, Innatec, Germany). A constant gas stream (typically 1.5 L min^{-1}) was adjusted using digital mass flow controllers (MFC, SmartTrak, Sierra, USA). Reactant gases (^{12}CO , ^{13}CO , H_2 from Linde, Germany)

were added to the N_2 stream *via* separate flow controllers. A 200 mbar check valve separates the dry gas section (A) from the wet gas section (B). In order to create a reactive aerosol, the gas stream was passed through a wash bottle containing 150 mL mixed buffer (50 mM Tris, MES, PIPPS, pH 3–9) and sodium dithionite (DT) and/or methyl viologen (MV). In Fig. S1 (ESI[†]), we demonstrate how water-soluble ingredients are transported with the aerosol stream, *i.e.* to reduce MV *via* DT enrichment. Section (C) shows the spectroscopic set-up. The aerosol was fed into a gas-tight polychlorotrifluoroethylene (PCTFE) compartment, attached on top of the ATR crystal plate and equipped with six optional gas inlets, a manometer for pressure control, and a transparent glass window for UV/vis irradiation.²⁸ The FTIR spectrometer (Tensor27, Bruker Optik, Germany) was equipped with a triple-reflection ZnSe/Si crystal ATR cell (Smith Detection, USA) and placed in an anaerobic chamber (Coy Laboratories, USA). Infrared spectra were recorded with 80 kHz scanning velocity at a spectral resolution of 1 cm^{-1} from 3800 to 1200 cm^{-1} . Under these conditions, the time-resolution of data acquisition is in the range of 1 s (one interferometer scan in the forward/backward direction). In order to determine the timescale of equilibration in the aerosol set-up, we probed a fast process, *i.e.* CO inhibition.²⁸ Fig. S2 (ESI[†]) shows the accumulation of **Hox**-CO upon exchange of N_2 to CO gas. Here, the **Hox** $\rightarrow$ **Hox**-CO conversion rate determined by ATR FTIR largely exceeds the value suggested by flash-photolysis IR spectroscopy ($t_{1/2} \approx 3 \text{ s}$ and 15 ms , respectively).^{58–60} The former represents the timescale of equilibration and defines the temporal resolution of the aerosol set-up. Note that all analyzed titrations proceeded by far slowly and were therefore not limited by the experimental time resolution. ATR FTIR measurements were performed at 25°C air-conditioning and on hydrogenase films derived by controlled dehydration and rehydration of $1 \mu\text{L}$ protein sample as reported earlier ($500 \mu\text{M}$ hydrogenase in 100 mM Tris/HCl pH 8, 100 mM NaCl, and 2 mM DT if not stated otherwise). Samples of anaerobically purified and activated [FeFe]-hydrogenase typically contained the H-cluster in a mixture of the **Hox**, **Hox**-CO, and **Hred** states. **Hox** was enriched in the film under a constant stream of N_2 aerosol for 30 minutes prior to the pH or redox titrations.

Infrared data evaluation

A quantitative analysis of infrared spectra was performed using in-house software tools (Fig. S2, ESI[†]). Spectra in the $1750\text{--}2150 \text{ cm}^{-1}$ frequency range were corrected for background contributions (liquid H_2O) by subtraction of polynomial spline functions. The resulting spectra were normalized to unity area in the CO/CN^- spectral region and the CO/CN^- bands were fitted by Voigt functions (50% Gaussian and Lorentzian characters) for frequency, peak width, and relative intensity determination. A global analysis approach was used (simultaneous fitting of time-series of spectra with a common set of frequencies and relative intensities for each H-cluster species). The relative population of a specific H-cluster species at a given time point was derived as the fraction of its summed CO/CN^- band intensities of the total (normalized) spectral intensity. We measured an experimental



uncertainty of <25% for the determination of rate constants in pH and redox titrations (Fig. S2, ESI†).

Density functional theory calculations

DFT calculations on model structures of the H-cluster were performed at the TPSSH functional and TZVP basis-set level of theory using Gaussian09.^{61–64} Starting models comprised the complete H-cluster and were constructed on the basis of the crystal structure (PDB entry 4XDC) of CPI [FeFe]-hydrogenase as described previously.²⁸ In some cases, adjacent amino acids and water molecules were included (see below). Anti-ferromagnetic coupling was assigned to the H-cluster prior to geometry optimization by proper definition of molecular fragments (broken symmetry approach). Calculations involved the COSMO solvation model ($\epsilon = 4$). H-cluster structures with varying protonation sites were calculated. During structural relaxation, the positions of the four carbon atoms of the S-CH₃ groups (representing the truncated terminal cysteine ligands of the H-cluster) were restraint at their crystallographic positions.⁶ A maximum deviation of $\sim 3 \text{ cm}^{-1}$ was observed for the CO and CN[−] band frequencies between calculations on constraint or unconstraint model structures (average deviation of $\sim 1.5 \text{ cm}^{-1}$). Vibrational frequencies for the relaxed molecular structures were derived from normal mode analysis as implemented in Gaussian09. Summed band intensities of CO/CN[−] ligands from the calculations were normalized to unity for comparison with experimental data. For quantification of the correlation between experimental (exp) and calculated (cal) frequencies (F) and relative band intensities (I), the root-mean-square-deviation (rmsd) was calculated (eqn (1), n = number of CO/CN[−] bands). For band intensities, rmsd_I values were derived using the uncorrected calculated intensities (an almost ideally linear correlation was observed for **Hox**: $I_{\text{ox}}^{\text{cal}} = 0.2\% + 0.99I_{\text{ox}}^{\text{exp}}$). For band frequencies, rmsd_F values were calculated after correction of the calculated frequencies for a systematic deviation from ideal correlation due to the DFT theory level.⁶¹ Calculated frequencies for **Hox** were plotted *versus* the experimental **Hox** frequencies, a linear fit was performed yielding slope ($s_{\text{ox}} = 1.15$) and offset ($o_{\text{ox}} = -243 \text{ cm}^{-1}$) values for **Hox** (eqn (2)). The corrected (cor) frequencies for all H-cluster species (i) were calculated using eqn (3) (n = number of bands, *i.e.* 2 CN[−] plus 3 CO = 5 bands for, *e.g.*, **Hox**) and the s_{ox} and o_{ox} values (meaning that rmsd_F values for all species were directly comparable to the rmsd_F for **Hox**).

$$\text{rmsd} = \sqrt{\sum_{i=1}^n (F_i^{\text{exp}} - F_i^{\text{cor}})^2 / n} \quad (1)$$

$$F_{\text{ox}}^{\text{cal}} = o_{\text{ox}} + s_{\text{ox}} F_{\text{ox}}^{\text{exp}} \quad (2)$$

$$F_i^{\text{cor}} = (F_i^{\text{cal}} - o_{\text{ox}}) / s_{\text{ox}} \quad (3)$$

Results and discussion

IR band shifts upon acidification of [FeFe]-hydrogenase

Fully active [FeFe]-hydrogenase HYDA1 as reconstituted with the native adt cofactor^{19,20} was used to form a protein film on

the ATR silicon crystal. The active-ready, oxidized state **Hox** was enriched to purity under a stream of humidified N₂ at pH 8 and 2 mM sodium dithionite (DT).²⁸ In contrast to earlier studies^{23,32} no build-up of reduced H-cluster species was observed, most likely due to release of H₂ from the protein film. It can be assumed that the decay rate of reduced species exceeds their formation rate and the removal of H₂ *via* the N₂ carrier gas precludes a re-reduction of enzyme. When the film was exposed to an acidified aerosol (pH 4, 2 mM DT), the IR band pattern of **Hox** was lost in favor of a novel spectrum (Fig. 3A and B). We note that the indicated pH values refer to the pH of the respective buffer used in the experiments. Based on calibration experiments, a difference of approximately ~ 0.25 pH units between the buffer and the sample film was estimated (Fig. S3, ESI†). The novel pattern showed an up-shift of the CO/CN[−] bands by 4–6 cm^{−1} with similar relative intensities as in the **Hox** spectrum. Furthermore, the pattern comprised the characteristic low-frequency band of the μCO ligand, up-shifted by 10 cm^{−1}. The conversion was found to be fully reversible when changing the pH in the aerosol back to alkaline values (Fig. S4 and S5, ESI†). Although the HYDA1 H₂ release activity was increasingly inhibited from pH 7 to 5, no structural changes were observed in the amide region when titrating the film from pH 8 to 4 (Fig. S6, ESI†). Note that extreme conditions (pH 3, 500 mM DT) led to irreversible loss of cofactor bands (Fig. S4, ESI†). Frequency shifts by 4–10 cm^{−1} are not compatible with redox changes at the diiron site and the shift to higher frequencies excludes a reduction of the H-cluster, *e.g.* by DT. Dithionite is commonly used as a reductant to promote H₂ evolution from [FeFe]-hydrogenases in solution.⁶⁵ Accordingly, we attribute the observed ‘blue shift’ to a protonation event and denote the novel species **HoxH** (Table 2). It is important to note that the **Hox** → **HoxH** conversion was not observed in the absence of DT (Fig. S4, ESI†). This result suggests that reducing conditions are required to induce the protonation event and the shift of cofactor bands is unrelated to protonation of amino acid residues.

Exposure of bacterial [FeFe]-hydrogenases CPI^{adt} and native DDH to pH 4 and 2 mM DT caused a similar ‘blue shift’ of the **Hox** band pattern (Fig. 3C). Formation of **HoxH** was not affected by the accessory iron–sulfur clusters in the prokaryotic enzymes or amino acid differences near the H-cluster. We note that HYDA1 and CPI apo-enzymes were artificially matured^{19,20} whereas DDH was isolated from *D. desulfuricans* in the native form.⁵⁷ The formation of **HoxH** hence is unrelated to the maturation procedure. Besides HYDA1^{adt}, we investigated cofactor variants with a less basic (odt, sdt) or hydrophobic (pdt, edt) bridgehead group at the diiron site. For all modified enzymes, a similar **Hox** → **HoxH** conversion at pH 4 and 2 mM DT was observed (Fig. 3D). This clearly proves that the bridgehead atom does not correspond to the protonation site. However, the cofactor variant HYDA1^{pdt} showed an approximately 20-fold slower formation of **HoxH** in comparison to HYDA1^{adt} (Fig. S5, ESI†), presumably related to the much smaller H₂ release activity of the pdt variant.⁶⁶ Protein variants with amino acid exchanges in the proton pathway towards the diiron site were studied (C169D and C169A). Although typical IR spectra and quantitative formation of **HoxH** were



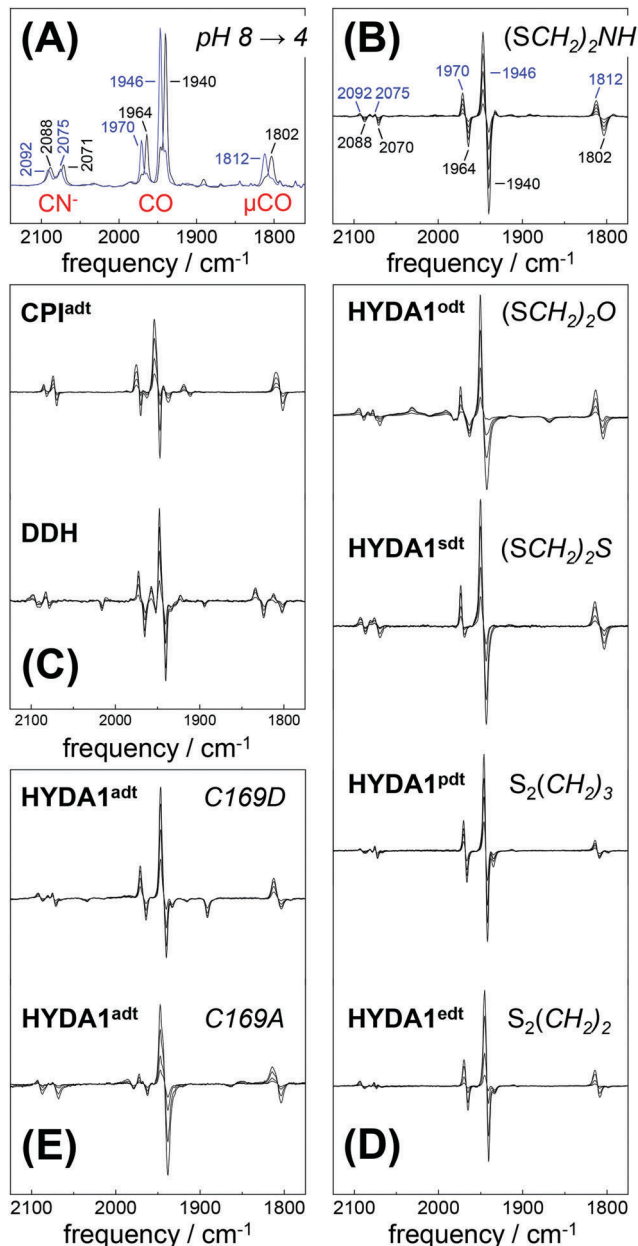


Fig. 3 FTIR difference spectra for the **Hox** → **HoxH** conversion ('blue shift'). The CO/CN[−] spectral region is shown. (A) Absolute spectra for the **Hox** → **HoxH** conversion in native HYDA1^{adt} obtained at pH 8 (black trace) and pH 4 in the presence of 2 mM DT (blue trace). Assignment of bands to vibrational modes was derived from DFT calculations (CN[−] and CO/μCO marked in red). (B) Difference spectra obtained during **Hox** (negative bands) to **HoxH** (positive bands) conversion in HYDA1^{adt} from the same data set as in (A). (C) Difference spectra for bacterial [FeFe]-hydrogenases CPI^{adt} and DDH^{adt}. (D) Difference spectra for HYDA1 cofactor variants odt, sdt, pdt, and edt. (E) Difference spectra for HYDA1^{adt} amino acid variants C169A and C169D. Table S1 (ESI[†]) lists the vibrational frequencies for all analyzed variants.

observed in both variants (Fig. 3E), the rate of **HoxH** formation in the C169A variant was smaller than in wild-type HYDA1^{adt} or in the C169D variant (Fig. S5, ESI[†]), ruling out that cysteine 169 functions as the protonation site in **HoxH**. The amino acid variant HYDA1 C169D showed H₂ release activity significantly

Table 2 IR band frequencies of CO and CN[−] ligands in HYDA1^{adt}

H-Cluster species	IR frequency/cm ^{−1}					Additional reference
	CN [−] bands		CO bands			
Hox	2088	2070	1964	1940	1802	21, 22 and 28
HoxH	2092	2074	1970	1946	1812	This study
Hred'	2084	2066	1962	1933	1792	29, 31 and 49
Hred'/H	2086	2068	1966	1938	1800	This study
Hhyd	2088	2076	1980	1960	1860	40
Hred	2070	2033	1961 ^a	1915	1891	21, 22 and 31
Hsred	2068	2026	1953 ^a	1918	1882	32
Hox-CO	2091	2081	1968	1962 ^b	1808	25–28
HoxH-CO	2094	2086	1972	1966 ^b	1816	This study

^a The bridging carbonyl (μCO) is moved to a terminal position at Fe₄ in **Hred** and **Hsred**. ^b In the CO-inhibited states, the coupled pCO/dCO stretching vibration gives rise to an additional IR band (**Hox-CO**: 2012 cm^{−1}, **HoxH-CO**: 2006 cm^{−1}). IR frequencies were determined from the spectra shown in Fig. 3, 6 and Fig. S7, S10 (ESI).

similar to native enzyme (up to 80%) while HYDA1 C169A did not evolve H₂ in the presence of DT and MV. We therefore assume that H₂ release activity is correlated with the formation rate of **HoxH**.

Proton and reductant concentration effects on HoxH formation

Protein concentration, hydration level, and temperature affected the **Hox** → **HoxH** conversion rate (Fig. S6, ESI[†]). Accordingly, a quantitative comparison demanded tight control of all experimental parameters. We determined an experimental uncertainty in the determination of conversion rates of less than 25% (Fig. S2, ESI[†]) and observed negligible differences in the hydration level (Fig. S6, ESI[†]). Accordingly, it was possible to investigate the influence of proton and DT concentration on the yield and formation rate of **HoxH**. Fig. 4A shows the **Hox** → **HoxH** conversion with individual protein films after a change of the aerosol from pH 8 to pH 6–4 with 10 mM DT. To elucidate the influence of DT, **HoxH** formation was monitored in a concentration range of 0–1000 mM DT at pH 6 (Fig. 4B). Aqueous solutions of DT were freshly prepared and the pH was checked before and after each experiment, which excluded any decomposition of DT.^{67,68} Clearly, the formation rate of **HoxH** depends on the DT concentration as well. The sigmoidal time course of the **Hox** → **HoxH** transition suggested that at least two consecutive processes are involved, *i.e.* proton transfer and electron transfer. In contrast, back conversion to **Hox** showed exponential behavior and the rate was found to be virtually independent of pH and DT (Fig. S5, ESI[†]). These observations might suggest that the divergent velocity of the **Hox** ↔ **HoxH** conversion is related to protein structural rearrangements in the 'blue shift' reaction. However, the formation of **HoxH** was not associated with changes in the protein structure or denaturation (Fig. S6, ESI[†]). Despite the high concentrations of DT used in some experiments, no enrichment of reduced H-cluster species was observed.

The formation rate of **HoxH** increased linearly with proton concentration in a pH range of 4–7 at 10 mM DT (Fig. 5A). A pH titration with a 50-fold increase of DT (500 mM) showed an overall faster **Hox** → **HoxH** conversion and linear regression



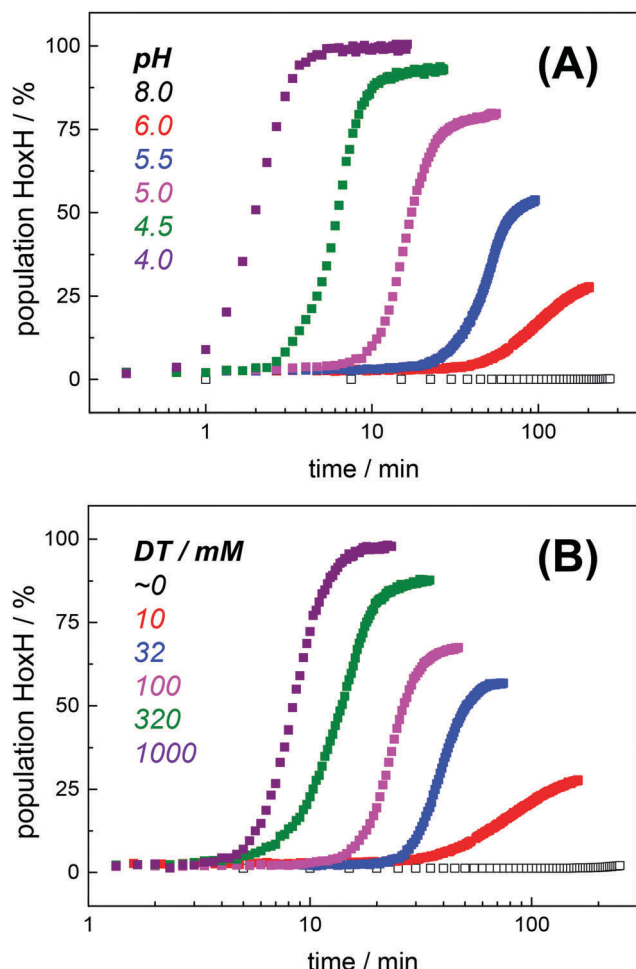


Fig. 4 **Hox** → **HoxH** conversion in **HYDA1^{adt}** as a function of pH and dithionite concentration. (A) Increase of the **HoxH** population at a constant DT concentration of 10 mM and pH 6–4 in the aerosol. No **HoxH** is observed at pH 8 (open squares). (B) Increase of the **HoxH** population at a constant pH of 6 and varying DT concentration (10–1000 mM) in the aerosol. No **HoxH** is observed in the absence of DT (open squares, residual DT might still be present in the preparation). Each data point in (A) and (B) corresponds to a normalized IR spectrum for which the relative **HoxH** population (in %) was calculated from the respective sum of CO/CN[−] peak areas.

from pH 3 to 7 (Fig. S4, ESI[†]). In contrast, saturation behavior of the **HoxH** formation rate was observed at excess DT concentrations (Fig. 5B). Accelerated formation of **HoxH** was also observed in the presence of the redox mediator MV (Fig. S5, ESI[†]) which is often used to optimize electron transfer from DT to hydrogenase enzymes.⁶⁵ Fig. 5 compares the **HoxH** yield for pH titrations at 10 mM or 500 mM DT, respectively. Significant formation of **HoxH** was observed even at alkaline pH, and the apparent p*K* shifted from ~5.5 to ~7.3 for higher DT concentrations. The titration behavior of a protonatable group should be independent of reductant concentration. The variable, apparent p*K* value thus excludes a specific amino acid residue as the protonation site in **HoxH**.

Ligand exchange and isotope editing at the H-cluster

Further information on the formation of **HoxH** was obtained from CO-inhibition and isotopic exchange studies. **HYDA1^{adt}**

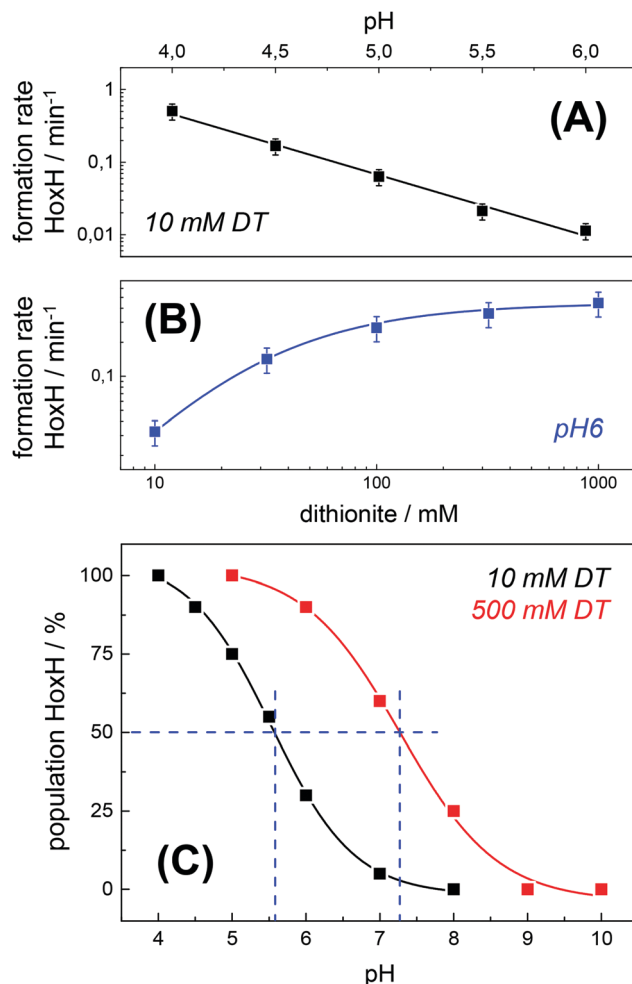


Fig. 5 Rate and yield of **Hox** → **HoxH** conversion in **HYDA1^{adt}**. From the time points (*t*) of half-maximal **HoxH** formation, apparent rate constants (1/*t*) for the **Hox** → **HoxH** conversion were derived (see Fig. 4 for corresponding data). Error bars were estimated from multiple experiments under the same conditions (Fig. S2, ESI[†]). (A) **HoxH** formation rate as a function of pH (10 mM DT). Note the logarithmic y-axis, the lines show linear regressions. (B) **HoxH** formation rate as a function of DT concentration (pH 6) (the line shows a saturation curve). (C) **HoxH** yield as a function of pH for 10 mM (black) and 500 mM DT (red, see Fig. S4, ESI[†] for corresponding data). The apparent p*K* (blue dashes) differs by about two units.

was quantitatively inhibited with 1% CO gas, as indicated by the typical IR bands of **Hox**-CO (Table 2).²⁸ Acidification in the presence of 2 mM DT under 1% CO induced an overall similar 'blue shift' of the CO/CN[−] bands as observed for **Hox** (Fig. S7, ESI[†]). We attribute the shifted spectrum to a protonated, CO-inhibited H-cluster species, **HoxH**-CO. Formation of **HoxH**-CO implies that protonation cannot occur at the distal iron ion because its ligand sphere was completed by the additional CO ligand already.^{24–28} Similar to the **Hox** → **HoxH** conversion, DT was required for the formation of **HoxH**-CO. Exchange of the bridging sulfur (μS) against selenium (μSe) in the [4Fe-4S] cluster of the catalytic cofactor has been shown to barely affect catalytic activity.⁵⁶ Selenium-substituted **HYDA1^{adt}** showed unchanged **Hox** and **HoxH** IR-band patterns. Although the p*K* values of thiols and selenols can differ by up to three units^{69–71}



and a protonation event at μSe of the $[\text{4Fe-4Se}]$ cluster should influence the formation rate of **HoxH**, the latter was not found to be affected (Fig. S8, ESI†). Therefore, direct protonation of the bridging sulfur atoms in the $[\text{4Fe-4S}]$ cluster seems to be less likely. Exposure of $[\text{FeFe}]$ -hydrogenase to an aerosol containing D_2O instead of H_2O facilitated quantitative deuteration of the protein film (Fig. S9, ESI†). For diiron model complexes, protonation at the bridging thiolate sulfur atoms has been considered.^{72–74} The frequency down-shift of the S–H vibration upon deuteration⁷⁵ from ~ 2550 to $\sim 1850\text{ cm}^{-1}$ could help in identifying protonation of the dithiolate ligand at the diiron site due to vibrational coupling with the CO/CN^- ligands. However, formation of **HoxH** in the presence of 0.5% D_2SO_4 and 2 mM DT did not result in altered **Hox** and **HoxH** IR-band patterns (Fig. S9, ESI†), even when HYDA1 was isolated and activated in the presence of D_2O before film formation and acidification. A direct detection of SH/SD bands in the sample is unrelated to vibrational coupling with the cofactor ligands and was not attempted. These results render protonation of the thiolate sulfur atoms at the diiron site in **HoxH** less likely.

Reduction of HYDA1 with H_2

The H_2 uptake activity of $\text{HYDA1}^{\text{adt}}$ was exploited to enrich reduced H-cluster species in the protein film. In the presence of 10% H_2 the IR signature of **Hox** was lost in favor of band patterns mainly attributable to reduced species (Fig. 6 and Table 2). In the H_2 – N_2 difference spectrum at pH 8, prominent CO bands at 1933 cm^{-1} and 1792 cm^{-1} resembled a pattern previously observed in reduced $\text{HYDA1}^{\text{pdt}}$ enzyme.²⁹ These bands have been assigned to **Hred'** in $\text{HYDA1}^{\text{adt}}$ recently.^{31,49} Interestingly, we found that **Hred'** was favored over **Hred** when progressing from acidic to alkaline pH, and *vice versa* (Fig. S10, ESI†). This can be explained by the PCET reactivity in the formation of **Hred** and destabilization of the reduced diiron site at alkaline pH.³¹ For a distinct assignment of IR band patterns, we exchanged the bridging and distal ^{12}CO ligands in $\text{HYDA1}^{\text{adt}}$ to ^{13}CO , following a previously established protocol.²⁸ Reduction of the isotope-edited enzyme with H_2 facilitated direct assignment of the bands at 1891 cm^{-1} and 1933 cm^{-1} to dCO of **Hred** and **Hred'**, respectively (Fig. 6). Whether the μCO band at 1792 cm^{-1} belonged to **Hred** or **Hred'** still remains unclear. To refine the assignment, HYDA^{adt} was reduced under 10% H_2 and titrated from pH 8 to pH 6 (Fig. 6 and Fig. S10, ESI†). Although the H_2 concentration was kept constant, acidification caused pronounced changes in the relative population of redox species and **Hsred** and **Hred'** were lost in favor of **Hred** and **Hox**. Due to the concomitant decrease of bands at 1933 cm^{-1} (dCO) and 1793 cm^{-1} (μCO), the experiment facilitated a clear assignment of these bands to **Hred'** (Table 2). The deviating pH-dependence of bands at 1933 cm^{-1} and 1891 cm^{-1} further distinguishes **Hred'** from **Hred** (Fig. S10, ESI†). In agreement with former observations,⁴⁰ an increase of **Hhyd** was observed when decreasing the pH from 8 to 6 (Fig. 6). We observed no 'blue shift' of the marker bands of **Hred** (1891 cm^{-1}) and **Hsred** (1882 cm^{-1}) at acidic pH.

To investigate **Hred'** in greater detail, we focused on HYDA1 cofactor variant pdt. This variant does not form **Hred** and

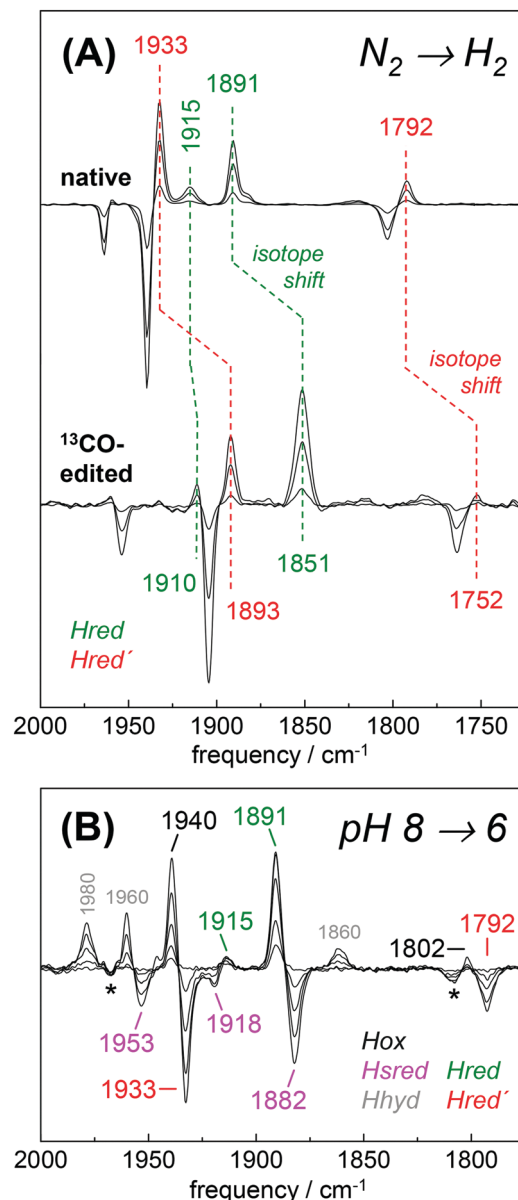


Fig. 6 Reduction of $\text{HYDA1}^{\text{adt}}$ in the presence of H_2 . (A) IR difference spectra in the CO region collected during a change from 100% N_2 to 10% H_2 on native (top) and ^{13}CO -edited $\text{HYDA1}^{\text{adt}}$ (bottom). Spectra show an increase of **Hred** (green) and **Hred'** (red) at the expense of **Hox** (black). Isotope editing of the μCO and dCO ligands leads to $\sim 40\text{ cm}^{-1}$ frequency down-shifts in **Hox**, **Hred**, and **Hred'**. (B) Difference spectra for changing the pH from 8 to 6 in the presence of 10% H_2 (for absolute spectra see Fig. S10, ESI†). Here, **Hred** (green) and **Hox** (black) are populated at the expense of **Hred'** (red) and **Hsred** (magenta). Note the additional formation of **Hhyd** (grey) and decrease of the minor **Hox-CO** population (asterisks).

Hsred and therefore represents an excellent model to study the relation between **Hox** and **Hred'**.²⁹ Although $\text{HYDA1}^{\text{pdt}}$ has been reported to show only minimal H_2 uptake activity,⁶⁶ we observed quantitative **Hox** $\rightarrow$ **Hred'** conversion under 100% H_2 (Fig. 7). At pH 8, both states showed similar IR band patterns as in $\text{HYDA1}^{\text{adt}}$ with relative frequency shifts of less than 5 cm^{-1} . When $\text{HYDA1}^{\text{pdt}}$ was purged with N_2 at pH 4 and in the presence of 2 mM DT, an IR spectrum similar to the one of



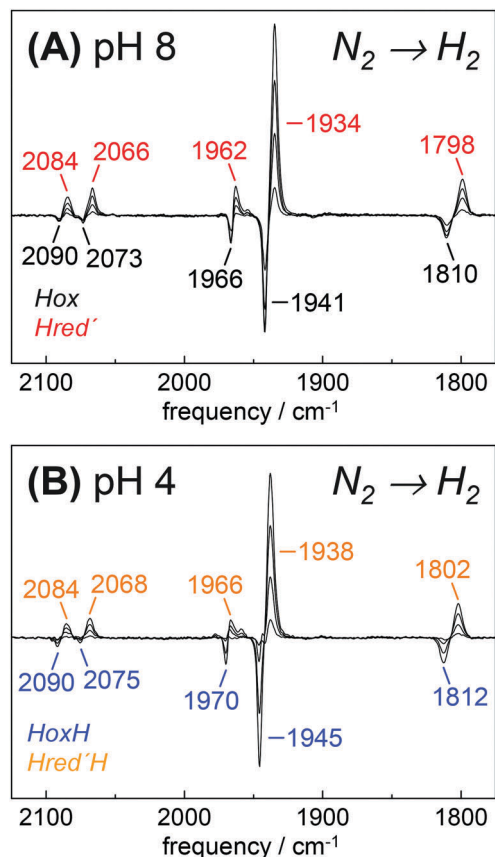


Fig. 7 Reduction of HYDA1^{Pdt} in the presence of H₂. IR difference spectra collected during a change in the aerosol from N₂ to H₂. (A) At pH 8 and 2 mM DT, **Hox** (black) was found to be de-populated in favor of **Hred'** (red). (B) At pH 4 and 2 mM DT, **HoxH** (blue) converted into **Hred'H** (orange).

HoxH was observed (compare Fig. 3). Exposure to 100% H₂ under acidic conditions resulted in a minor 'blue shift' of all CO/CN⁻ bands with respect to **Hred'** (Fig. 7). We denote the species responsible for this shifted spectrum as **Hred'H** (Table 2). Redox titrations indicated the presence of **Hred'H** in HYDA1^{adt} as well, although this state was more difficult to detect in native enzyme due to competing population of **Hred'**, **Hsred** and **Hhyd** (Fig. S10, ESI†).⁴⁰

When the hydrogenase concentration in the film was decreased by dilution with bovine serum albumin protein (BSA), the **Hox** → **Hred'** conversion rate drastically dropped for relative concentrations of HYDA1^{Pdt} below 25% and no formation of reduced states was observed with 5% HYDA1 (Fig. S11, ESI†). This behavior suggests that the inter-molecular electron transfer between [FeFe]-hydrogenase molecules (disproportionation) was involved in the formation of one-electron reduced **Hred'** from oxidized **Hox** via two-electron reduced species transiently formed upon reduction with H₂. Inefficient inter-molecular electron transfer may explain the low rates of H₂ uptake that have been reported for HYDA^{Pdt} in solution.⁶⁶ In contrast, the formation rate of **HoxH** was only slightly affected by dilution of HYDA1^{Pdt} with BSA (Fig. S11, ESI†). This observation rules out inter-molecular electron transfer as an explanation for the

apparent absence of reduced species in the presence of high DT concentrations.

Density functional theory on H-cluster protonation

DFT calculations on H-cluster model structures (Fig. S12, ESI†) were used to explore conceivable protonation sites. Respective positions included N(adt), the μS atoms in the diiron site (S1, S2) and in the [4Fe-4S] cluster (S3–S6), as well as the sulfur atoms of the three terminal cysteine ligands (S8–S10). Good agreement between the experimental and calculated CO/CN⁻ band patterns of **Hox** was obtained, as judged on the basis of the small root-mean-square-deviations (rmsd) derived from experimental and calculated band frequencies (*F*) and intensities (*I*), i.e. rmsd_F ~ 8 cm⁻¹ and rmsd_I < 3% (Fig. 8).²⁸ We note that relative band positions and intensities of all investigated H-cluster species were largely independent of the size of the model structure and geometry-optimization restraints that were

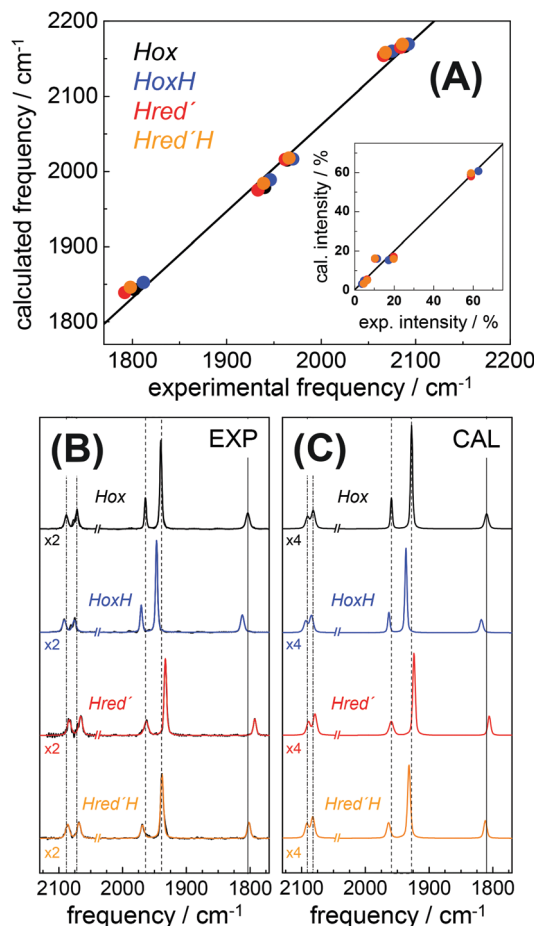


Fig. 8 Comparison of IR spectra from experiment and theory. (A) Plots of calculated (cal) vs. experimental (exp) IR frequencies and intensities (inset) for **Hox**, **HoxH**, **Hred'**, and **Hred'H**. (B) Experimental normalized IR spectra (black thin lines) and fit curves (smooth lines). (C) Calculated normalized IR spectra (from DFT at the TPSSH/TZVP level and after correction of band frequencies for systematic effects, eqn (1)–(3)). Dashed lines in (B) and (C) emphasize band shifts relative to **Hox**; CN⁻ bands were scaled by factors of 2 (exp) or 4 (cal) for clarity. Experimental and calculated IR data of **Hox**-CO and **HoxH**-CO are shown in Fig. S15 (ESI†).

used in the calculations (Fig. S13, ESI†). Protonation at N(adt) or S1/S2 at the diiron site caused frequency up-shifts three to four fold larger than experimentally observed for **HoxH** (compare Table 1). The large rmsd_F values ($15\text{--}25\text{ cm}^{-1}$) suggested that these groups are unlikely protonation sites (Fig. S14, ESI†). Protonation at the μS atoms of the $[\text{4Fe-4S}]$ cluster caused band shifts in better agreement with the experiment (rmsd_F $8\text{--}11\text{ cm}^{-1}$). However, protonation of S3 or S5 produced a downshift of the pCO band and intensity changes less compatible with the experimental observations. Furthermore, protonation at S4 or S6 caused considerable distortion of the $[\text{4Fe-4S}]$ cluster (e.g. $>0.5\text{ \AA}$ elongation of the S4/S6-Fe bonds), which rendered μS atom protonation at the $[\text{4Fe-4S}]$ cluster less likely. This interpretation is in line with previous theoretical results on $[\text{4Fe-4S}]$ clusters.⁷⁶ Protonation at S10 of a terminal cysteine ligand yielded a frequency downshift of the pCO band not observed in the experiment. In contrast, good agreement was found for protonation at S8 or S9 ($\text{rmsd}_F \sim 7\text{ cm}^{-1}$). Best overall agreement with the **HoxH** band pattern was observed for protonation at S9 (Fig. 8). Protonation at water molecules in the vicinity of the $[\text{4Fe-4S}]$ cluster or at the bridging cysteine S7 caused significantly larger IR band shifts than protonation at S9 (Fig. S14, ESI†). Notably, deuteration at S9 caused negligible CO/CN[−] band shifts ($<1\text{ cm}^{-1}$), in good agreement with the experimental data of **HoxH** in D₂O buffer and **HoxH-CO** (Fig. S14 and S15, ESI†).

Infrared spectra were calculated for H-cluster structures of the one-electron reduced **Hred'** state. A **Hox**-like structure with a μCO ligand and apical vacancy at the distal iron ion yielded a mean frequency down-shift of $\sim 35\text{ cm}^{-1}$ of the CO/CN[−] bands, far larger than experimentally observed (Fig. S16, ESI†). Excess frequency down-shifts ($\text{rmsd}_F > 30\text{ cm}^{-1}$) were also observed for protonation of the reduced H-cluster at N(adt) or S1/S2. Inclusion of an apical hydride at the distal iron ion produced excess frequency up-shifts ($\text{rmsd}_F \sim 40\text{ cm}^{-1}$), in particular for the μCO ligand, resembling the μCO shift in **Hhyd**.^{37–40} Protonation at one of the sulfur atoms of the $[\text{4Fe-4S}]$ cluster yielded small IR band down-shifts in good agreement with the experiment, but led to structural distortions, whereas protonation at a cysteine sulfur preserved the cubane geometry in **Hred'**. Best agreement between calculated and experimental IR frequencies of **Hred'** was obtained for protonation at S9 ($\text{rmsd}_F \sim 9\text{ cm}^{-1}$, see Fig. 8). These results showed that the IR spectrum of **Hred'** is well compatible with a one-electron reduced H-cluster, which carries an additional proton and adopts a similar geometry to **Hox**. Structures for **Hred'H** were calculated comprising a second-site protonation. Similar trends with respect to structural changes and IR patterns in response to protonation at various sites as for **HoxH** and **Hred'** were observed (Fig. S16, ESI†). Protonation at two cysteine S-atoms of the $[\text{4Fe-4S}]$ cluster (i.e., S8 and S9) yielded IR frequencies in good agreement with the **Hred'H** spectrum (Fig. 8). Accordingly, **Hred'H** was attributed to a double-protonated H-cluster species with the same electron count and cofactor geometry as **Hred'**.

Good agreement between calculated and experimental IR band patterns for **Hox/HoxH**, and **Hred'/Hred'H** facilitated assignment of the IR bands of the CO/CN[−] ligands to individual

vibrational modes. In addition, our simulations provided access to the spin and charge distribution of the cofactor (Fig. S17, ESI†). Protonation (i.e. at S9) and protonation/reduction affected the largely uncoupled behavior of the diatomic ligand vibrations only marginally. In the four states, the spin density distribution at the diiron site was similar, showing that the apparent mixed-valence Fe(I)Fe(II) configuration was preserved. The net spin thus resided largely on the diiron site in **Hox/HoxH**, in agreement with EPR data.^{46–48} For **Hred'/Hred'H**, the spin density implied that the surplus electron was localized mainly at the $[\text{4Fe-4S}]$ cluster, as supported by the increased negative charge density on the $[\text{4Fe-4S}]$ cluster. Anti-ferromagnetic coupling between the unpaired spins at the diiron site and at the reduced $[\text{4Fe-4S}]$ cluster was expected to render these states EPR-silent.²⁹ The minor charge variations at the iron ions of the diiron site correlate well with the small experimental IR band frequency differences of the CO ligands in the four states (Fig. S17, ESI†).

Protonation at the H-cluster

Real-time ATR FTIR spectroscopy on $[\text{FeFe}]$ -hydrogenase protein films identified four states of the H-cluster that show similar CO/CN[−] band patterns but differ in the redox and/or protonation state (**Hox/HoxH** and **Hred'/Hred'H**). The oxidized state, **Hox**, has been characterized extensively. In agreement with crystallographic assignments^{4–7} and previous FTIR studies,^{21–28} DFT favored a μCO ligand and an apical vacancy at the distal iron ion in **Hox**.⁷⁷ Our calculations suggest similar cofactor structures in all four species. Furthermore, both **Hox** and **HoxH** can be assigned to a mix-valence diiron site with an oxidized $[\text{4Fe-4S}]$ cluster^{46–48} while in **Hred'** and **Hred'H** a reduced $[\text{4Fe-4S}]$ cluster was identified.²⁹ The mixed-valence state clearly distinguishes **Hred'** from **Hred** and **Hsred** (see Table 1). While **Hox** represents the unprotonated H-cluster, protonation at the diiron site has been suggested for **Hred** and **Hsred**.^{31–34,38} We found that a satisfactory calculation of the **Hred'** band pattern also requires addition of a proton. This view is supported by the experimentally observed pH-dependence of **Hred'** formation in both HYDA1^{pd} and HYDA1^{adt}. However, protonation of the diiron site or at surrounding amino acid residues cannot explain the observed IR band shifts. Our computational results favor a terminal cysteine ligand of the $[\text{4Fe-4S}]$ cluster as a common protonation site in **HoxH** and **Hred'/Hred'H**. In HYDA1, cysteine 417 (S9) is the most likely candidate. Additional protonation occurs in **Hred'H**, presumably at C170 (S8). Iron-sulfur clusters are typically involved in electron transfer processes, but their participation in proton transfer has been documented as well.^{78–82} More specifically, protonation at a terminal cysteine ligand has been proposed to be involved in the catalytic mechanism of $[\text{NiFe}]$ -hydrogenases.^{83–86}

Formation of **HoxH** and **Hred'/Hred'H** was found to be independent of the proton pathway to the active site that includes C169, N(adt), and Fe_d. The crystal structure of the HYDA1 apo-protein⁸⁷ suggests that the $[\text{4Fe-4S}]$ cluster in HYDA1 is in direct contact with bulk water while in bacterial $[\text{FeFe}]$ -hydrogenases, the H-cluster is buried deeper in the catalytic domain.^{4–7} However, a chain of water molecules extends from the surface close to the cysteine that corresponds to our favored protonation site at the



[4Fe–4S] cluster (Fig. S18, ESI†). Two proton trajectories thus may be present in [FeFe]-hydrogenases: the ‘catalytic’ proton pathway that shuttles protons to the diiron site^{50–52} and a pathway that facilitates protonation at the [4Fe–4S] cluster.

Reaction sequence of electron and proton transfer

While the exponential **HoxH** → **Hox** conversion implies a direct deprotonation of the [4Fe–4S] cluster, the sigmoidal time course of **HoxH** formation suggests at least two successive reaction steps to be involved in the ‘blue shift’ reaction. **HoxH** was found to accumulate under H₂ release conditions, that is, for increasing concentrations of the reductant (such as the one-electron donor sodium dithionite, DT). This places **HoxH** towards the end of a multi-step reaction cycle (Fig. S19, ESI†). Under conditions of maximal H₂ release (e.g. pH 7–8, 200 mM DT as in Fig. S6, ESI†) the prevalence of **Hred'** over **Hred** suggests **Hred'** as an intermediate in the **Hox** → **HoxH** conversion. **Hox** dominates at alkaline pH so that we assume that deprotonation of **HoxH** is required to regain **Hox**. In contrast, the quantitative accumulation of **HoxH** suggests that this deprotonation was slow or impaired at low pH. Numerical simulations show that the above considerations can qualitatively reproduce the pH and DT dependence of **HoxH** formation (Fig. S20, ESI†). This model also consistently explains the formation of **HoxH**-CO due to reductive conversion of **Hox**-CO into **Hred'**, H₂ release, and subsequent rebinding of CO to **HoxH** (Fig. S7, ESI†). It has been shown that reduction of the [4Fe–4S] cluster weakens the binding of external CO.^{13,29}

Our data provide evidence that protonation at the same cysteine ligand of the [4Fe–4S] cluster occurred in both **HoxH** and **Hred'**. We assume that this protonation occurs from a putative donor group (BH⁺), such as a water molecule consistently observed in CPI and DDH structures (Fig. S18, ESI†).^{4–7} Deprotonation of **HoxH** likely involves the same molecule, now serving as a proton acceptor (B). Re-protonation of this group at acidic pH (*i.e.* via the adjacent chain of water molecules) would impair the deprotonation reaction so that **HoxH** accumulates (Fig. S19, ESI†). **Hred'H** is populated upon reduction and protonation of **HoxH** at low pH, but seems not to advance further in the reaction cycle, possibly because second-site protonation interferes with the H₂ forming reactions. Further studies are required to clarify this point. Accumulation of **Hred'** under H₂ likely involves inter-molecular electron transfer between oxidized and (transiently formed) two-electron reduced [FeFe]-hydrogenase proteins (either in the **Hsred** or **Hhyd** states). We note that inter-molecular electron transfer seems to require a densely packed protein film and is unlikely to occur under physiological conditions.

Conclusions

In contrast to **Hox** and **Hred'**, **Hred** and **Hsred** carry an additional electron at the diiron site and lack the Fe–Fe bridging CO ligand.^{31–34,38} Respective ligand rotation into a terminal or “semi-bridging” position has been considered difficult to reconcile with the large hydrogen turnover rates of [FeFe]-hydrogenases.^{41–43}

An involvement of **Hred** and **Hsred** in hydrogen turnover is certainly not excluded at present.^{31,38,60} However based on the differences in ligand orientation (bridging *vs.* terminal), these states can be considered unlikely intermediates in the reaction cycle of rapid H₂ formation.³⁴ In HYDA1^{adt}, formation of **Hred** was suppressed at alkaline pH in favor of **Hred'** while for acidic pH values **Hred** dominated over **Hred'**. Thus, the relative population of **Hred** and **Hred'** in HYDA1^{adt} critically depends on pH. This suggests that the rates of proton delivery either to the [4Fe–4S] cluster or to the diiron site may be decisive for the location of electrons at the H-cluster. In addition to the ‘catalytic’ proton pathway towards the diiron site,^{50–55} we propose a ‘regulatory’ proton pathway to the [4Fe–4S] cluster. Concomitant protonation and reduction at the [4Fe–4S] cluster in **Hred'** stabilize the mixed-valence state and prevent ligand rotation and premature protonation at the diiron site. Whether this is a sequential or concerted process remains to be elucidated.^{88–90}

Conflicts of interest

There are no conflicts to declare.

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