Hydride state accumulation in native [FeFe]-hydrogenase with the physiological reductant H₂ supports its catalytic relevance†

Moritz Senger, † Tobias Kernmayr, † Marco Lorenzi, ‡ Holly J. Redman † and Gustav Berggren †*‡

Small molecules in solution may interfere with mechanistic investigations, as they can affect the stability of catalytic states and produce off-cycle states that can be mistaken for catalytically relevant species. Here we show that the hydride state (H_{hyd}), a proposed central intermediate in the catalytic cycle of [FeFe]-hydrogenase, can be formed in wild-type [FeFe]-hydrogenase treated with H₂ in absence of other, non-biological, reductants. Moreover, we reveal a new state with unclear role in catalysis induced by common low pH buffers.

Hydrogenases are redox enzymes catalysing molecular hydrogen (H₂) uptake and production. The high H₂ evolution frequency of prototypical [FeFe]-hydrogenases motivates basic research on the reaction mechanism and inspires catalyst design for a hydrogen economy.¹−³ In these enzymes, hydrogen catalysis takes place at a hexa-iron cofactor, called the H-cluster. It is composed of a [4Fe4S]-cluster linked via a cysteine residue to a unique diiron site ([2Fe]) that binds two cyanide (CN⁻) and three carbonyl (CO) ligands. These ligands serve as intrinsic probes sensitive to infrared spectroscopy and report on reduction and protonation events during catalysis. The diiron site is bridged by an azadithiolate ligand (SCH₂NHCH₂S⁻, ADT) that is proposed to shuttle protons via its amine group to the apical vacancy, the site of hydrogen catalysis, located at the Fe ion (Feₕ) distal to the [4Fe4S] cluster (Fig. 1). The most studied group of [FeFe]-hydrogenases (Group A) transfers protons to the active site via a conserved Proton Transfer Pathway (PTP) composed of amino acid residues and water molecules.⁴−⁶ Recently described [FeFe]-hydrogenases in group D lack this PTP.⁷,⁸ Several redox and protonation states have been characterised and proposed as catalytic intermediates.¹,² The starting point of catalysis is the oxidised state H_{ox}. Under inert atmosphere conditions at low pH values and in presence of chemical reductant a blue-shifted variant of H_{ox} is formed, commonly denoted H_{oxH}.⁵,¹⁰ One electron reduction of H_{ox} results in population of two singly reduced states: H_{red} or H_{redH}. H_{red} features a reduced [4Fe4S] cluster while the diiron site is still in an oxidised configuration. Formation of the diiron site reduced state H_{red} is coupled to a proton uptake event, although the site of protonation is under discussion.¹¹,¹² A second reduction step yields the super-reduced state (H_{redH}) or a hydride state (H_{hyd}). H_{redH} is characterised by a reduction of both the [4Fe4S] cluster and the diiron site.¹³ In contrast, H_{hyd} features a terminal hydride at Feₕ, yielding a

Fig. 1  The calculated hydride state structure of the H-cluster in [FeFe]-hydrogenases. The H-cluster consists of a protein bound [4Fe4S] cluster that is covalently linked via a cysteine thiolate ligand to the diiron site ([2Fe]), further ligated by three carbonyl (CO) and two cyanide (CN⁻) ligands. A terminal hydride (H⁻) at the distal iron ion is proposed to combine with a proton delivered by the ADT ligand to yield molecular hydrogen in the last step of hydrogen catalysis. Colour code: orange-iron, yellow-sulphur, green-carbon, red-oxygen, blue-nitrogen, white-hydrogen. The structure is drawn after the PDB coordinates 4XDC optimized for the H_{hyd} state by DFT calculations. (ref. 33.)
formally oxidized diiron site (Fig. 1). As the last step in hydrogen turnover, the terminal hydride of H$_{\text{hyd}}$ is proposed to combine with a proton delivered via the PTP to generate molecular hydrogen. A hydride state similar to H$_{\text{hyd}}$ but with an additionally protonated ADT bridge has been proposed as the last intermediate in hydrogen evolution.

A terminal-hydride state of the H-cluster, considered to reflect the H$_{\text{hyd}}$ state, can be stabilised by impairing the enzyme’s catalytic function via amino acid variations in the PTP or cofactor alteration. This has enabled its characterization using a range of techniques, including NMR, NRVS, Mössbauer, EPR and FTIR spectroscopy. In native, fully functional, [FeFe]-hydrogenases the accumulation of a highly similar H$_{\text{hyd}}$ state is commonly achieved by exposure to H$_2$ at low pH values in the presence of the non-physiological reductant sodium dithionite (NaDT, Table S1, ESI†) (with exceptions). Based on chronoamperometry experiments it was recently proposed that [FeFe]-hydrogenases are inhibited by NaDT at low pH values (or rather SO$_2$, one of its oxidized by-products), i.e. conditions similar to H$_{\text{hyd}}$ accumulation. Considering its in-depth spectroscopic characterization, and proposed central importance in the catalytic cycle, the possibility that the to-date characterized H$_{\text{hyd}}$-state reflects an inhibited “artefact-state” would represent a significant setback in our mechanistic understanding of [FeFe]-hydrogenase.

Here, we selectively enrich the double reduced states, H$_{\text{hyd}}$ and H$_{\text{sred}}$, by exposure to molecular hydrogen and follow the respective absorbance changes of cofactor ligand bands by Attenuated Total Reflection Fourier-transform Infrared (ATR-FTIR) Spectroscopy. Data is collected at different pH values and buffers. This study reveals that small carboxylic acids, often used in low pH buffers in enzyme electrochemistry experiments, are non-innocent in formation of a new H-cluster species similar to H$_{\text{ox}}$. At present it is not clear if this is an artefactual off-cycle state, or in fact a central intermediate that previously has escaped detection. Moreover, we show the accumulation of H$_{\text{hyd}}$ in native [FeFe]-hydrogenase, HydA1 from Chlamydomonas reinhardtii, at mildly basic and low pH values regardless of buffer choice and more importantly in the absence of NaDT.9,10,16 However, the preference for H$_{\text{hyd}}$ accumulation at low pH is clearly independent of the presence of NaDT. As H$_{\text{hyd}}$ and H$_{\text{sred}}$ are generally considered to reflect two different tautomers, their relative stability as a function of pH value is arguably a consequence of the protonation state of the H-cluster species denoted H$_{\text{oxc}}$. Green bands belong to H$_{\text{oxc}}$. The band positions are indicated by bars.

A solution of the HydA1 [FeFe]-hydrogenase was applied to the crystal surface of the ATR-FTIR setup, following removal of any NaDT contaminations by chromatography and buffer exchange. The enzyme solution was subsequently dried and rehydrated under N$_2$ atmosphere to form an auto-oxidised protein film.28,29 The hydrogenase adopted the expected oxidised active-ready state H$_{\text{ox}}$ and was subsequently exposed to H$_2$ gas. The resulting difference spectra are displayed in Fig. 2. As a consequence of H$_2$ uptake, a mixture of the hydride state H$_{\text{hyd}}$ (blue positive bands, 2081, 2066, 1979, 1960, 1860 cm$^{-1}$) and the super-reduced state H$_{\text{sred}}$ (red positive bands, 2070, 2025, 1954, 1918, 1882 cm$^{-1}$) was populated while the oxidised state H$_{\text{ox}}$ (grey negative bands, 2089, 2071, 1965, 1940, 1802 cm$^{-1}$) was depopulated. This holds true for different pH values, in a range of different buffers. Fig. 2 displays the ATR-FTIR difference spectra of HydA1 samples adjusted to pH 8 or pH 4 and subsequently exposed to H$_2$ (absolute spectra and pH adjustment in Fig. S1, ESI†). In each spectrum the signature of H$_{\text{hyd}}$ (blue positive bands) is clearly detectable. A larger fraction of H$_{\text{sred}}$ is formed at pH 8 (100 mM Tris buffer) while a larger fraction of H$_{\text{hyd}}$ is populated at pH 4 (100 mM propionate buffer), in both cases a concomitant loss of H$_{\text{ox}}$ is observed in contrast to earlier results where a depopulation of H$_{\text{ox}}$H was observed at pH 4 due to the presence of NaDT.9,10,16 However, the preference for H$_{\text{hyd}}$ accumulation at low pH is clearly independent of the presence of NaDT. As H$_{\text{hyd}}$ and H$_{\text{sred}}$ are generally considered to reflect two different tautomers, their relative stability as a function of pH value is arguably a consequence of the protonation state of the H-cluster surrounding. The reaction of auto-oxidised HydA1 films with H$_2$ gas proceeds within one-two seconds while auto-oxidation, observed when H$_2$ was replaced by N$_2$, occurs over 10-20 seconds (compare Fig. S2, ESI†). Both kinetics report indirectly on the gas accessibility and hydration of the protein film in the ATR-FTIR setup. Dehydrated [FeFe]-hydrogenase samples were reported to be protected from exposure to gases.

Fig. 2 (bottom) shows that H$_2$ exposure causes H$_{\text{hyd}}$ accumulation in the absence of NaDT preferentially at low pH values (compare Fig. 2 top). This trend had been reported by several groups before, however always involving NaDT.16,17,19,25 Apart from the recent proposed binding of NaDT (or one of its oxidized products) to the H-cluster, protonation has been suggested to stabilize the reduction at the [4Fe4S] cluster for
and in analogy to \( H_{\text{red}}' \) and \( H_{\text{oxH}} \) (blue shifted) we propose,\(^3,31,34\) or \( H_{\text{oxC}} \). Conversely, di- and tricarboxylic acids such as succinic acid (\( pK_{a1} = 4.9 \)) and caproic acid (\( pK_{a1} = 4.2 \)), oxalic acid (\( pK_{a1} = 4.14 \)) or citric acid (\( pK_{a1} = 3.1 \)) did not induce the new species (Fig. S3, ESI\(^\dagger\)). Thus, stabilizations of \( H_{\text{oxC}} \) appear to be determined by the number of the carboxylic acid groups, rather than molecular weight of the acid.

Compared to \( H_{\text{ox}} \) the overall band pattern of \( H_{\text{oxC}} \) is retained and in analogy to \( H_{\text{red}} \) (red shifted) and \( H_{\text{ox}} \) (blue shifted) we propose that the H-cluster geometry and electronic structure is highly similar. Moreover, the requirement for low pH in forming the state implies that the buffer molecule interacts in its protonated, uncharged form (Fig. S4, ESI\(^\dagger\)). A complete characterisation of this new H-cluster state is beyond the scope of the current study, but we note that it could be of high relevance in particular as e.g. acetic acid (acetate) is commonly employed in protein film electrochemistry studies of \([\text{FeFe}]\)-hydrogenases.\(^37,33,45\) Formaldehyde has previously been observed to act as an inhibitor of the H-cluster under reducing conditions.\(^46,47\) In the latter case, binding of formaldehyde on the \([2\text{Fe}]\) subsite via a Fe–C bond was proposed from a combination of ENDOR spectroscopy and DFT calculations,\(^48\) and further supported by studies of model complexes.\(^49\) Whether a similar binding model can be applied also to carboxylic acids remains to be verified. The extent of the new state (\( H_{\text{oxC}} \)) formed in electrochemistry setups, its role in catalysis and its implication on catalytic currents, especially at low pH, remain unclear. In contrast to the \( H_{\text{ox}} \) state, the same \( H_{\text{hyd}} \) state forms regardless of the nature of the buffer, rendering a stabilizing role of the buffer or other additives unlikely (Fig. 2 and Fig. S1, ESI\(^\dagger\)). Our data shows clearly that \( H_{\text{hyd}} \) can be accumulated in the absence of NaDT and without introducing loss-of-function mutations. In closing, this underscores the flat energy landscape of the H-cluster during catalysis, and it is tempting to conclude that the previously characterized \( H_{\text{hyd}} \) state indeed represents one of the final intermediates in hydrogen evolution catalysis of \([\text{FeFe}]\)-hydrogenases.

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Conflicts of interest

There are no conflicts to declare.

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