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The Structure of the Active Site H-Cluster of [FeFe] Hydrogenase from the Green Alga Chlamydomonas reinhardtii Studied by X-ray Absorption Spectroscopy[†]

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ABSTRACT: The [FeFe] hydrogenase (CrHydA1) of the green alga Chlamydomonas reinhardtii is the smallest hydrogenase known and can be considered as a "minimal unit" for biological H₂ production. Due to the absence of additional FeS clusters as found in bacterial [FeFe] hydrogenases, it was possible to specifically study its catalytic iron—sulfur cluster (H-cluster) by X-ray absorption spectroscopy (XAS) at the Fe K-edge. The XAS analysis revealed that the CrHydA1 H-cluster consists of a [4Fe4S] cluster and a diiron site, 2Fe_H, which both are similar to their crystallographically characterized bacterial counterparts. Determination of the individual Fe-Fe distances in the [4Fe4S] cluster ($\sim 2.7 \text{ Å}$) and in the 2Fe_H unit ($\sim 2.5 \text{ Å}$) was achieved. Fe-C (=O/N) and Fe-S bond lengths were in good agreement with crystallographic data on bacterial enzymes. The loss of Fe-Fe distances in protein purified under mildly oxidizing conditions indicated partial degradation of the H-cluster. Bond length alterations detected after incubation of CrHydA1 with CO and H₂ were related to structural and oxidation state changes at the catalytic Fe atoms, e.g., to the binding of an exogenous CO at 2Fe_H in CO-inhibited enzyme. Our XAS studies pave the way for the monitoring of atomic level structural changes at the H-cluster during H₂ catalysis.

Hydrogenases are efficient catalysts for the production and cleavage of molecular hydrogen (H₂) (1, 2). Their use in biotechnological applications is expected to significantly contribute to future renewable fuel production (3, 4). The molecular principles of H2 turnover in these enzymes may lead to novel non-platinum catalysts (5). It is thus an important challenge to unrayel the reaction mechanism of H₂ catalysis in hydrogenases.

In biocatalysis the [FeFe] hydrogenases are of particular interest as they show extremely high rates of H₂ production (6, 7). The structures of [FeFe] hydrogenases from two bacterial species, namely, Clostridium pasteurianum and Desulfovibrio desulfuricans, have been resolved by protein crystallography (8, 9). Their active site is a unique iron—sulfur cluster, comprising six Fe atoms commonly known as the H-cluster (6, 10). It can be

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Abbreviations: 2Fe_H, binuclear Fe unit of the H-cluster; adt, azadithiolate; AI, as-isolated state; CpI, Clostridium pasteurianum [FeFe] hydrogenase; DdH, Desulfovibrio desulfuricans [FeFe] hydrogenase; EPR, electron paramagnetic resonance spectroscopy; EXAFS, extended X-ray absorption fine structure; FTIR, Fourier-transform infrared spectroscopy; *CrHydA1*, [FeFe] hydrogenase protein of C. reinhardtii; MS, multiple scattering; XAS, X-ray absorption spectroscopy.

divided into a [4Fe4S] cluster and a binuclear iron unit, 2Fe_H, ¹ cysteine-linked to the cubane. This diiron moiety is probably the active site of H₂ catalysis (8, 11, 12). This situation differs from that in the [NiFe] hydrogenases where a heterobimetallic complex forms the active site (13). Depending on the redox conditions, the two Fe atoms of 2Fe_H bind three to four CO and two CN ligands. One CO may be found in an Fe-Fe bridging position (14-17). An azadithiolate (adt) moiety has been proposed, but the precise chemical identity of this ligand on top of the 2Fe_H site is not yet settled (12, 18). Besides the H-cluster, bacterial [FeFe] hydrogenases harbor up to four additional FeS clusters, serving as electron transfer relays (19). Signal overlay from these clusters complicates investigations on the reaction cycle of the bacterial H-cluster by Fe-specific techniques, i.e., Mössbauer (20) and X-ray absorption spectroscopy (XAS) (18, 21, 22).

[FeFe] hydrogenases exist not only in anaerobic bacteria but also in photosynthetic eukaryotes. For several green algae, the hydA genes and respective proteins have been identified (23–26). These extraordinarily small proteins may be regarded as a "minimal unit" of biological H₂ conversion. Sequence alignments show that the small F-domain carrying the FeS relay clusters in bacteria is missing (24, 25). However, four cysteine residues necessary for H-cluster accommodation in the H-domain are well conserved in all hydrogenase-coding hydA genes (23).

The biophysical characterization of algal [FeFe] hydrogenases is hampered by the fact that these enzymes are extremely O₂ sensitive and synthesized in vivo only in small amounts (23, 28). To overcome low protein yields, we have established a general

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system for heterologous expression and synthesis of [FeFe] hydrogenases in *Clostridium acetobutylicum*. The [FeFe] hydrogenase from *Chlamydomonas reinhardtii*, *Cr*HydA1, is efficiently assembled due to the maturation apparatus of the host organism, yielding 2 mg of protein/L of cell culture (28). Hence, a sufficient amount of *Cr*HydA1 [FeFe] hydrogenase is available for indepth spectroscopic explorations.

CrHydA1 is an ideal candidate to study the assembly, structure, and catalytic mechanism of [FeFe] hydrogenases. First, EPR studies on CrHydA1 and two further algal [FeFe] hydrogenases revealed similar signals of their CO-inhibited state of the active site as found in bacterial enzymes. However, deviations in the EPR g-tensors suggested distinct differences in the electronic structure of the H-cluster (27). Direct information on the geometric structure of the CrHydA1 H-cluster is required to compare it to the situation in bacteria.

In the present study X-ray absorption spectroscopy (XAS) (29) at the Fe K-edge was performed to obtain atomic level structural and electronic information on the H-cluster of CrHydA1. We show that the structural parameters imply an overall organization of the algal H-cluster similar to that in bacterial enzymes. Structural alterations induced by the inhibitor CO and the substrate H_2 are addressed.

MATERIALS AND METHODS

Protein Sample Preparation. C. reinhardtii [FeFe] hydrogenase CrHydA1 was heterologously synthesized and isolated as previously described (30). C. acetobutylicum ATCC 824 recombinant strains were grown in CGM media containing up to 60 g/L glucose in a 2.5 L MiniFors bioreactor (Infors, Augsburg, Germany) (28, 31). Both cell growth and protein purification were carried out under strictly anoxic conditions. Isolated protein was concentrated up to 1 mM (about 48 mg/mL) on Vivaspin 6 and Vivaspin 500 columns (Sartorius Stedim Biotech, Göttingen, Germany) and stored in 10% glycerol and 2 mM sodium dithionite (NaDT) for stabilization.

The specific H_2 -evolving activity was determined by an *in vitro* assay as described before (23). The gas mixture was injected into a gas chromatograph (GC-2010; Shimadzu, Kyoto, Japan) equipped with a PLOT fused silica coating Molsieve column (5 Å, 10 m by 0.32 mm) from Varian (Palo Alto, CA). The specific activity of the hydrogenase was calculated from the detected amount of produced H_2 .

For gas treatments, protein samples ($\sim 1 \text{ mM}$) were filled into open 200 µL PCR tubes and placed in 8 mL Suba tubes, which were rubber sealed under the N_2/H_2 atmosphere of the anaerobic tent. To humidify the working gases (CO, H₂), degassed water was bubbled with each of the two gases for 30 min, respectively. The headspace of the hermetically sealed Suba tubes was flushed with gas for 15-30 min. The final protein concentration of the samples used at the DESY Hamburg was adjusted right after the gas treatments; hence, more diluted protein was flushed. During the treatments, protein was kept on ice and protected against light. For the "as-isolated" samples (AI), no gas treatment was applied. Here, AI^{red} denotes an (optimal) enzyme isolation procedure in the presence of 2 mM NaDT, whereas AI^{ox} samples are preparations without any reducing agents (NaCl was used instead of NaDT to adjust the ionic strength of the buffers). Samples were filled into Kapton-covered acrylic-glass sample holders for XAS and frozen in liquid nitrogen. EPR measurements on the XAS samples (not shown) revealed no indications for significant unspecific iron in the as-isolated reduced *Cr*HydA1 samples as the "rhombic iron" signal at a *g*-value of ~4 was practically absent. The absence of water-dissolved Fe ions in as-isolated reduced *Cr*HydA1 samples was also likely according to the K-edge spectra, which showed no evidence for Fe–O bonds (see Results).

X-ray Absorption Spectroscopy (XAS). Kα fluorescencedetected XAS spectra at the Fe K-edge were collected at 20 K using energy-resolving Ge detectors and helium cryostats as previously described (22, 32) at beamline D2 of the EMBL outstation (at HASYLAB, DESY, Hamburg, Germany) and at beamline KMC-1 of BESSY (Berlin, Germany). Harmonic rejection was achieved by detuning of the Si(111) double-crystal monochromators to 50% of their peak intensities. The energy resolution of the used monochromators was ~3 eV at DESY and optimized by appropriate setting of aperture slits and slightly lower, \sim 4 eV, at BESSY due to the beamline specifics (33). The slightly different energy resolution of the XAS spectra does not affect the conclusions drawn from the data analysis. Spectra were collected maximally for a scan range of 6950–8450 eV, i.e., up to $k = 19.5 \text{ Å}^{-1}$, within $\sim 1 \text{ h}$ each. Dead time-corrected XAS spectra were averaged after energy calibration of each scan using the peak at 7112 eV in the first derivative of the absorption spectrum of an Fe foil as an energy standard (estimated accuracy ± 0.1 eV) (18). EXAFS oscillations were normalized and extracted as described in refs 29 and 34. The energy scale of EXAFS spectra was converted to the wave vector scale (k-scale) using an E_0 value of 7112 eV. Unfiltered k^3 -weighted spectra were used for least-squares curve fitting employing a curved-wave multiplescattering (MS) approach with the program EXCURV (35). In our hands, the EXCURV program is able to reproduce the expected coordination numbers of Fe-ligand interactions in Fe model compounds with known structure from fitting of respective EXAFS oscillations ranging up to k-values of 20 Å^{-1} (18). An amplitude reduction factor, S_0^2 , of 0.9 was used in the EXAFS fits. EXAFS fits initially were based on a model of the H-cluster derived from the crystal structures (8, 9). The CO and CN ligands were classified as units, with almost linear arrangement of the Fe-C=O/N atoms (bond angles close to 180°). For the units, correlation was enabled. A maximal path length in MS calculations of 10 A and extending over a maximal scattering order of 4 was employed. Debye–Waller factors $(2\sigma^2)$ were either fixed at physically reasonable values (see Table 2) or varied in the simulations. Fourier transforms (FTs) were calculated from k^3 weighted EXAFS data using the program SimX (29) and employing cos² windows ranging over 10% at both ends of the k-range. From experimental K-edge spectra the preedge peak region was extracted by subtraction of a polynomial spline through the main edge rise using the program XANDA (36). K-edge energies were determined by the "integral method" (29), using integration limits of 15% and 90% of normalized intensity.

RESULTS

As-Isolated Reduced State of CrHydA1. Purification of the protein under strictly anaerobic and reducing conditions (30) to obtain the as-isolated state of CrHydA1 (AI^{red}) yielded the most active protein (Table 1). Iron XAS spectra of AI^{red} were collected up to a k-value of 19.5 Å $^{-1}$, i.e., to \sim 1450 eV above the Fe K-edge at \sim 7120 eV (Figure 1). This very long k-range for XAS on proteins allows for resolution even of closely spaced interatomic distances (34, 37).

Table 1: Specific H₂-Evolving Activities of CrHydA1 Preparations

| sample | specific H_2 evolving activity (μ mol of H_2 mg^{-1} min^{-1}) \pm 20 |
|------------------------|---|
| CrHydA1 ^{red} | 650 |
| CrHydA1 ^{ox} | 590 |
| H ₂ treated | 620 |
| CO treated | 40 |

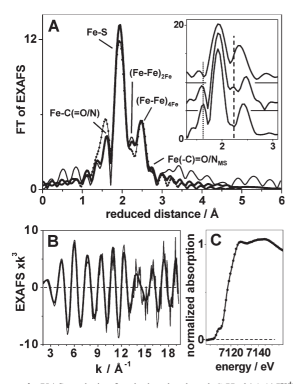


FIGURE 1: XAS analysis of as-isolated reduced CrHydA1 (AI^{red}) at the Fe K-edge. (A) Fourier transform of the EXAFS spectrum in (B). The FT was calculated from k-values ranging over $4.7-19.5 \, \text{Å}^{-1}$. Inset: FTs calculated from k-ranges starting at 7.4, 4.9, and 1.6 Å^{-1} (top to bottom); vertical lines mark contributions from Fe-C(=O/N) vectors (dots) and from the Fe-Fe vector in 2Fe_H (dashes). (A) and (B): thin lines, experimental data; thick lines, simulations (Table 2C); the dotted line in (A) is a simulation with parameters in Table 2A. (C) Fe K-edge spectra of two AI^{red} preparations measured at DESY (line) and BESSY (dots).

The K-edge spectrum of AI^{red} (Figure 1C) was perfectly reproducible in independent protein samples, indicating the high quality of the CrHydA1 preparation. Its small primary maximum at \sim 7130 eV revealed the predominant coordination of Fe by sulfur ligands (22, 38) as expected due to the presence of the respective conserved cysteines in CrHydA1. Indications for oxygen ligation of Fe, as expected for oxidatively modified (or damaged) FeS species (22), were absent.

The Fourier transform (FT) of the AI^{red} EXAFS spectrum shows three major peaks (Figure 1A), at reduced distances between \sim 1.5 and \sim 2.5 Å (the reduced distance is the true interatomic distance minus \sim 0.4 Å due to a phase shift). This immediately suggests at least three Fe-backscatterer shells, i.e., of Fe-C(=O/N), Fe-S, and Fe-Fe vectors. A shoulder on the high-distance side of the FT peak due to Fe-Fe interactions likely accounted for the expected Fe(C)=O/N interactions, the contributions of which to the EXAFS spectrum usually are enhanced by multiple scattering (MS) effects due to the almost linear Fe-C=O/N arrangement (see below). The FT peaks due to Fe-C(=O/N) and Fe(-C)=O/N interactions may be

expected to be similarly large (18). Here, the magnitude of the peak due to Fe(-C)=O/N apparently was diminished due to partial cancelation of respective EXAFS oscillations by interference with EXAFS contributions from the Fe-Fe interactions. A fourth peak at ~2.2 Å reduced distance became more prominent when the FT was calculated starting at higher k-values (Figure 1A, inset) where metal-metal interactions dominate the EXAFS oscillations (shown in Figure 1B). This result was suggestive of a Fe-Fe distance of 2.5–2.6 Å in the H-cluster. The Fe-Fe distance of the 2Fe_H unit in crystal structures of bacterial [FeFe] hydrogenases is about 2.5 Å, whereas the Fe-Fe distances in the [4Fe4S] cluster are ~2.7 Å (8, 9). Thus, the EXAFS data apparently allow to discriminate between the Fe-Fe distances in the [4Fe4S] and the 2Fe_H subgroups of the C. reinhardtii H-cluster.

Precise Fe-ligand distances in AI^{red} were determined by simulations (curve fitting) of the EXAFS spectrum. We used a multiple scattering (MS) approach (35, 39) to account for the expected (relatively small) MS contributions to the EXAFS from the linear CO and CN ligands at 2Fe_H. It should be noted that very similar fit results with respect to bond lengths and coordination numbers were obtained if MS contributions very neglected in the simulation procedure (not documented). First, coordination numbers were employed in the simulations (Table 2) which where based on the crystal structures of the H-cluster in bacterial [FeFe] hydrogenases (8, 9, 14). Possible weak Fe-Fe interactions between the two parts of the H-cluster were neglected in the simulations as all distances between Fe ions of the [4Fe4S] cluster and of 2Fe_H are ≥4 A in crystal structures of the bacterial enzymes and such interactions did not contribute significantly to the EXAFS spectra. The fit results in Table 2A and the corresponding fit curve (Figure 1A, dotted line) revealed that the main features of the EXAFS spectrum already were reasonably well reproduced by using the crystal structure as a template. All Fe-ligand distances, in particular the two Fe-Fe and Fe-S distances (Table 2A), are similar to those found in the crystal structures of CpI and DdH (8, 9). Sequence similarity of the catalytic H-domain of these bacterial hydrogenases to CrHydA1 is reasonable high. Still, structures derived from X-ray crystallography, especially for metal cofactors, have to be considered with care. The good agreement of XAS analysis and crystal structure, however, suggests that the overall organization of the green algae H-cluster is similar to that of bacteria.

A more elaborated fit approach yielded refined structural parameters of the H-cluster. Allowing the coordination numbers (N_i) of the Fe-C(=O/N) and Fe-Fe distances to vary in the fit yielded a lower $N_{\text{Fe-C}(=O/N)}$ value but a larger N-value of the \sim 2.5 Å Fe-Fe distance (Table 2B). An $N_{\text{Fe-C}(=O/N)}$ value of less than unity, i.e., less than one C(=O/N) ligand per each of the six Fe atoms on the average, was expected for the absence of a CO in a bridging position between the two iron atoms of 2Fe_H. In the literature, the reduced form of the H-cluster (H_{red}) has been reported to lack this bridging CO, and the respective CO group is bound to Fe_p (p = proximal) solely. Upon oxidation, the CO bridge between the distal and proximal Fe atoms is formed (14, 16, 17, 41-43). The as-isolated AI^{red} protein of CrHydA1 from our preparations is not likely to contain oxidized or CO-inhibited impurities. Due to our fast and gentle isolation protocol, EPR spectra of AIred did not show the axial EPR signal with g-values of 2.102, 2.040, and 1.998 as expected for CrHydA1^{ox} and 2.052, 2.007, and 2.007 for CrHydA1^{ox}-CO as reported recently (27). Roseboom et al. explained the frequent

emergence of CO-inhibited species by cannibalization of free CO from denatured protein (17). However, these impurities were not detected in AI^{red}. A larger apparent coordination number than the expected one of 0.33 (due to two Fe–Fe distances of ~2.5 Å per six Fe ions) of the Fe–Fe distance attributed to 2Fe_H may suggest contributions from similarly long Fe–S vectors. In the structure of reduced [FeFe] hydrogenase from *C. pasteurianum* the distance of Fe_p at the 2Fe_H site to the cysteine sulfur atom which links it to the [4Fe4S] cluster is ~2.50 Å, and the Fe–Fe distance in 2Fe_H is only slightly longer, ~2.55 Å. Thus, such a long Fe–S distance may contribute to the apparent coordination number of the Fe–Fe interaction. We note that in the crystal structures all further Fe–S bonds are shorter than about 2.4 Å, and such distances would not interfere with the 2.5 Å feature.

The coordination number of the Fe-Fe distance of ~ 2.7 Å $(N_{2.7})$ was very close to 2 (2.02) if it was allowed to vary in the fit (Table 2). This value is similar to the one of 2.0 that is calculated for the presence of only the H-cluster in CrHydA1 (i.e., for 12 Fe-Fe distances of ~ 2.7 Å in the cubane moiety per 6 Fe ions). In the [FeFe] hydrogenases of CpI and DdH, besides the H-cluster, four and two additional FeS clusters are found so that the $N_{2.7}$ values would be expected to be significantly larger, 2.27 and 2.52, respectively. Thus, our XAS data suggest, in concert with previously performed EPR experiments (27, 30) and multiple sequence alignments (23–25), that only the H-cluster is present in CrHydA1 and further FeS clusters are absent.

A significant further improvement of the fit quality (R_F less than 10%) was achieved by including separate shells for the Fe-C(=O) and Fe-C(=N) interactions and further slight parameter adjustments (Table 2C; Figure 1A,B, thick lines). Now, individual distances for Fe-C(=O) of 1.77 Å and Fe-C (=N) of 1.98 Å were obtained. These distances are close to those of the respective ligands in the crystal structures, where the Fe-C (=O) distance usually is shorter than Fe-C(=N) (8, 9).

We note that the above used "top-down" simulation approach, i.e., starting the EXAFS simulations with a model based on the crystal structures of the H-cluster in bacteria and then refining the structural parameters by additional degrees of freedom in the fit, leads to the same simulation results that were obtained by an inverse ("bottom-up") procedure where the structural model was developed by the stepwise inclusion of increasing numbers of coordination shells in the fit (not documented).

An unusual bridging ligand (which perhaps is an azadithiolate (adt)) has been proposed to be present in the 2Fe_{H} part of the bacterial H-cluster and may serve crucial functions in H_2 production (9, 14, 44–47). The respective carbon and nitrogen or oxygen atoms would be within a range of about 3–4 Å to the Fe atoms of 2Fe_{H} . As such distances overlap with those from Fe (-C)=O/N interactions and only weak contributions from these atoms to the EXAFS spectra were expected, the nature of the bridging species in the *C. reinhardtii* H-cluster cannot be deduced from the XAS data.

H-Cluster Integrity in Protein from Two Purification Conditions. The integrity of the active site was compared in asisolated CrHydA1 samples purified under reducing (AI^{red}) and mildly oxidizing (AI^{ox}) conditions (Figure 2). The specific H_2 -evolving activity of AI^{ox} samples, as determined by the *in vitro* assay, was only slightly lower than in AI^{red} (Table 1). The Fe K-edge was at \sim 0.5 eV higher energies in AI^{ox} compared to AI^{red} (Figure 2A), indicating that overall more reduced Fe was present in AI^{red} . Both K-edges were at higher energies than that for Fe(II)

| Table 2: Structural Parameters of the H-Cluster from EXAFS Simulations ^a | | | | | | | |
|---|------------------------|-----|-------------------------|-------------------------------|--------------------------------------|--------------------|--|
| sample | shell | fit | N _i (per Fe) | $R_{\rm i}(\mathring{\rm A})$ | $2\sigma_{i}^{2} (\mathring{A}^{2})$ | R _F (%) | |
| AI ^{red} | C(=O/N) | A | 1.00^{b} | 1.78 | 0.002^{b} | 22.5 | |
| | S | | 2.17^{d} | 2.28 | 0.001^{c} | | |
| | S | | 1.33^{d} | 2.39 | 0.001^{c} | | |
| | Fe (2Fe _H) | | 0.33^{b} | 2.52 | 0.002^{b} | | |
| | Fe (4Fe4S) | | 2.00^{b} | 2.73 | 0.009 | | |
| | C(=O/N) | В | 0.52 | 1.77 | 0.002^{b} | 21.0 | |
| | S | | 2.56 | 2.29 | 0.002^{b} | | |
| | S | | 1.14 | 2.40 | 0.002^{b} | | |
| | Fe (2Fe _H) | | 0.61 | 2.53 | 0.002^{b} | | |
| | Fe (4Fe4S) | | 2.02 | 2.71 | 0.010 | | |
| | C(=O) | C | 0.45^{e} | 1.77 | 0.001^{b} | 8.7 | |
| | C(=N) | | 0.55^{e} | 1.98 | 0.001^{b} | | |
| | S | | 2.04^{d} | 2.28 | 0.001^{b} | | |
| | S | | 1.46^{d} | 2.40 | 0.003^{b} | | |
| | Fe (2Fe _H) | | 0.77 | 2.53 | 0.002^{b} | | |
| | Fe (4Fe4S) | | 2.02 | 2.72 | 0.012 | | |
| AIox | C(=O/N) | D | 0.55 | 1.84 | 0.001^{b} | 7.7 | |
| (AI ^{red}) | | | (0.74) | (1.86) | $(0.001)^b$ | | |
| | S | | 2.96 | 2.28 | 0.010^{b} | | |
| | | | (3.57) | (2.28) | $(0.010)^b$ | | |
| | Fe (2Fe _H) | | 0.58 | 2.52 | 0.001^{b} | | |
| | | | (0.71) | (2.53) | $(0.001)^b$ | | |
| | Fe (4Fe4S) | | 1.19 | 2.70 | 0.010^{b} | | |
| | | | (2.06) | (2.73) | $(0.010)^b$ | | |
| AIred | C(=O/N) | E | 0.68 | 1.85 | 0.001^{b} | 8.3 | |
| [CO] {H ₂ } | | | [0.92] | [1.79] | $[0.001]^b$ | | |
| | | | {0.42} | {1.78} | $\{0.001\}^b$ | | |
| | S | | 3.45^{f} | 2.29 | 0.010^{b} | | |
| | | | $[3.45]^f$ | [2.27] | $[0.010]^b$ | | |
| | | | ${3.45}^f$ | $\{2.25\}$ | $\{0.010\}^b$ | | |
| | Fe (2Fe _H) | | 0.54^{f} | 2.53 | 0.002^{b} | | |
| | | | $[0.54]^f$ | [2.62] | $[0.002]^b$ | | |
| | | | $\{0.54\}^f$ | {2.51} | $\{0.002\}^b$ | | |
| | Fe (4Fe4S) | | 2.02^{f} | 2.73 | 0.012^{b} | | |
| | | | $[2.02]^f$ | [2.71] | $[0.012]^b$ | | |
| | | | $\{2.02^f\}$ | {2.71} | $\{0.012\}^b$ | | |

 aN_i , coordination number; R_i , Fe-ligand distance; $2\sigma_i^2$, Debye-Waller factor. b Fixed values. $^c2\sigma^2$ restricted to values >0. d The sum of $N_{\rm S}$ values was 3.5. e The sum of $N_{\rm C(=O/N)}$ values was 1. fN_i was coupled to yield equal values for the three spectra. $R_{\rm F}(29)$ was calculated over reduced distances of 1.3–2.8 Å. All simulations comprise a further multiple-scattering Fe(-C)=O/N shell (with the same N value as for the Fe-C (=O/N) shell; respective distances were in the range of 2.95–3.02 Å; $2\sigma^2$ was set to 0.01 Å 2).

species due to the presence of Fe(III) ions in the cubane cluster (48, 49). The shift of the edge to lower energies in AIred was compatible with the reduction of at least one Fe(III) ion to the Fe(II) state. Furthermore, the preedge peak, due to formally dipole-forbidden 1s → 3d electronic transitions, was increased in AI^{ox} (Figure 2A, inset). This suggests a less symmetric overall Fe coordination (40), presumably due to the binding of oxygen species to some Fe atoms in AI^{ox} (see below). Lowering of the site symmetry is expected to cause increased admixtures of metal p levels to the 3d electronic orbitals, increasing the probability of electronic transitions into unoccupied 3d levels and thus the intensity of the respective preedge feature. Due to the limited energy resolution of the monochromator at BESSY, possible spectral substructure on the preedge peaks, as related, i.e., to electronic multiplet interactions (40, 50), remained invisible.

The FTs of EXAFS spectra revealed a pronounced decrease of the peak due to Fe-Fe interactions in AI^{ox} (Figure 2B, arrow). EXAFS simulations (using a simplified approach) revealed a

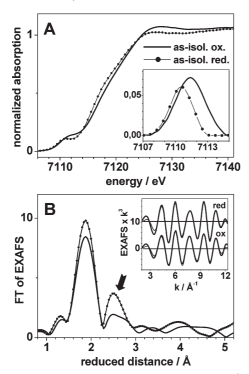


FIGURE 2: XAS comparison of as-isolated reduced (AI^{red}) and oxidized (AI^{ox}) *Cr*HydA1. (A) K-edge spectra (inset, isolated preedge features). (B) FTs of EXAFS spectra (dotted line, AI^{red}; solid line, AI^{ox}). FTs were calculated over a *k*-range of 1.6–12.5 Å⁻¹; the arrow marks the contributions mainly from Fe–Fe interactions. Inset: Fourier isolates over reduced distances of 0.5–3.0 Å (thin lines, experimental data) and simulations according to Table 2D (thick lines).

decrease of the coordination number of \sim 2.7 Å Fe—Fe vectors by a factor of about 2 and a less pronounced decrease of $N_{\rm Fe-S}$ and $N_{\rm Fe-C(=O/N)}$. The N-value of the \sim 2.5 Å Fe—Fe distance in 2Fe_H of AI^{ox}, however, was only slightly smaller than that of AI^{red} (Table 2D). These results may suggest that in AI^{ox} primarily the [4Fe4S] part of the H-cluster was modified or destroyed, whereas in the larger protein fraction the binuclear site at least still contained the \sim 2.5 Å Fe—Fe distance. Whether this Fe—Fe distance belongs to an otherwise unmodified 2Fe_H cluster cannot be concluded from the data.

Effects of Treatments with CO and H₂. The effects of incubation of CrHydA1 with CO and H₂ on the structure of the H-cluster strongly depended on whether concentrated or more dilute protein samples were treated (Figure 3). In diluted samples (Figure 3A), the increased primary maxima of the K-edges (at ~7125 eV) and lowered preedge amplitudes (at ~7111 eV) suggested overall more symmetric coordination of the Fe atoms. Such an effect was expected if Fe ions had become released from the protein to the medium, forming hexaquo—Fe ions. Such deleterious effects in dilute samples were observed with both gases. Analysis of the EXAFS data of H₂- and CO-treated dilute protein (not documented) revealed the partial loss of Fe—Fe interactions, which indicates a degradation of the H-cluster.

 H_2 and CO treatments on more concentrated samples caused only small changes of the K-edge spectra (Figure 3A), suggesting that the integrity of the H-cluster was fully preserved. A slight shift of the edge to higher energies (by \sim 0.4 eV) in H_2 -treated CrHydA1 points to the oxidation of one Fe atom at $2Fe_H$, possibly due to reduction of H-species (8, 18). The respective EXAFS spectrum (Figure 3B) showed only minor changes

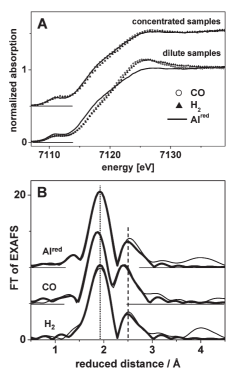


FIGURE 3: Effects of gas treatment on CrHydA1: CO binding and H_2 reduction. (A) K-edges of indicated CrHydA1 samples. (B) FTs of EXAFS spectra (thin lines, calculated as in Figure 2) and simulations (thick lines, Table 2E). H_2 and CO treatments in (B) were carried out on concentrated protein. Vertical dashed and dotted lines mark shifts in the respective FT peaks due to Fe-S and Fe-Fe interactions.

compared to AI^{red} . The main difference in the structure of the H-cluster, according to EXAFS simulations (Table 2E), was a decreased apparent Fe-C(=O/N) distance and a slightly decreased Fe-Fe distance of 2Fe_H in H₂-treated enzyme. Although this small decrease in the Fe-Fe distance may not be significant, it could be caused by the presence of one more oxidized Fe atom (likely Fe_d of 2Fe_H; d = distal) due to the binding and reduction of H-species. Notably, the Fe-Fe distance of the [4Fe4S] cluster was not affected by the H₂ treatment (Table 2E).

Carbon monoxide is an effective inhibitor of [FeFe] hydrogenases. In a crystallographic structure for H_{ox}-CO of CpI, electron density at Fe_d was discussed to be exogenous CO (41). FTIR and EPR analyses support this notion (14, 51). Under certain redox conditions, an endogenous CO is believed to form a "bridge" between the Fe atoms of the 2Fe_{H} moiety (6, 41). A decreased preedge peak in CO-treated CrHydA1 (Figure 3A) was in agreement with an overall more symmetric Fe coordination upon binding of additional CO to 2FeH. The EXAFS data (Figure 3B) revealed an increased coordination number of the Fe-C(=O/N) interactions accompanied by enhanced respective multiple scattering contributions to the spectrum and an overall reduced mean FeC(=O/N) distance (Table 2E). These findings indicate the presence of one to two surplus short Fe-C(=O) interactions. In comparison to AI^{red}, the Fe-Fe distance of 2Fe_H was increased significantly by ~0.1 Å. A similar increase of the Fe-Fe distance has been observed in crystal structures of CO-treated bacterial [FeFe] hydrogenase, where one extra CO was found at Fe $_{\rm d}$ (41). Presumably, in CrHydA1 an extra CO also was bound to Fe_d, causing more symmetric (near-octahedral) coordination of this ion.

DISCUSSION

XAS on biological metal centers (BioXAS) is a powerful tool to derive atomic resolution structural information, to determine the electronic configuration, especially in those states which are not accessible by EPR spectroscopy, and to monitor their dynamics during the catalytic reactions (29, 52-55). Here, we used XAS at the Fe K-edge to specifically investigate the catalytic cofactor of biological H₂ turnover in an [FeFe] hydrogenase protein.

The structural parameters from the present XAS study strongly suggest that the overall atomic organization of the H-cluster in the [FeFe] hydrogenase of the green alga C. reinhardtii is very similar to that found in crystallographically characterized enzymes from bacteria. Recent EPR investigations performed on CrHydA1 support this notion (27, 30). Furthermore, the XAS data strongly suggest that in CrHydA1 no additional FeS clusters are found, as opposed to bacterial hydrogenases (2, 8, 9). EXAFS analysis allowed precise determination of interatomic distances in the H-cluster. The individual Fe-Fe distances in the [4Fe4S] part and 2Fe_H moiety of the H-cluster were measured (precision on the order of $\sim 0.02 \text{ Å}$). Changes of the structural parameters upon treatments of CrHydA1 with the inhibitor CO and the substrate H₂ were detected, allowing to address changes, e.g., in catalytic intermediates. The XAS-derived interatomic distances, because of their higher resolution compared to protein crystallography (56), may help to optimize in silico structures of the H-cluster in DFT calculations (57-59).

There are also limitations of the XAS analysis. By the applied methods, predominantly average coordination environments of the six Fe atoms in the cluster were obtained. Characterization of the individual structure and oxidation state of the Fe atoms, in particular in the 2Fe_H unit, and of the binding sites of substrate and inhibitors is highly desirable. It may be facilitated by future investigations employing site-selective XAS techniques (60). It has been proposed that an azadithiolate is bridging the Fe atoms of the 2Fe_H site and is essential in H₂ turnover (12, 18, 45-47, 61). The respective C, N, and O atoms are almost impossible to detect by XAS because their distances to Fe are relatively large and overlap with the Fe(C)=O/N distances and with contributions of atoms from the protein backbone. Thus, discrimination between different bridging species likely cannot be obtained by XAS.

The integrity of the algal H-cluster is easily perturbed, as observed for CrHydA1 purified under mildly oxidizing conditions and upon incubation of dilute protein with H₂ and CO. Such conditions seem to cause the release of Fe ions from the protein into the medium and, hence, at least partial degradation of the H-cluster. Preliminary evidence was obtained that the [4Fe4S] cluster is the primary target of oxidative modification, whereas the 2Fe_H moiety may be more robust. O₂ sensitivity of FeS clusters is well-known (62, 63). Purification of CrHydA1 under reducing conditions prevents such deleterious effects and stabilizes an intact H-cluster. In bacterial [FeFe] hydrogenases, the H-cluster is deeply buried in the protein. Induced-fit folding models of CrHydA1 (M. Winkler and T. Happe, unpublished results) suggest that its [4Fe4S] cluster is located just beneath the surface whereas the 2Fe_H moiety faces the inside of the protein. This arrangement is likely to allow for easy access of gas molecules and redox partners from the bulk to the [4Fe4S] unit and for its rapid modification by O_2 .

Treatments of CrHydA1 with H2 and CO revealed relatively subtle but discernible structural changes of the H-cluster. Inter-

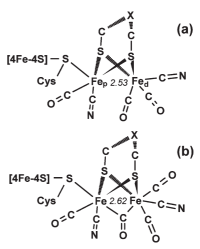


FIGURE 4: Tentative structures of the H-cluster in CrHydA1 for $\mathrm{AI}^{\mathrm{red}}$ and $\mathrm{AI}^{\mathrm{ox}}$ CO. (a) As-isolated reduced and (b) CO-treated protein, in agreement with the XAS results (Fe-Fe distances in Å). The bridging ligand on top of the 2Fe_H cluster is (partly) visible in the crystal structures of bacterial [FeFe] hydrogenases (8-12). Its chemical nature, however, is unclear (X may be N or O). The respective C and N or O atoms were not resolved in this XAS study.

action of the substrate H₂ with 2Fe_H seems to cause slight overall oxidation of the two Fe atoms in the 2Fe_H part, but its Fe-Fe distance was almost unchanged. Accordingly, a change in the Fe-Fe bridging motif upon H₂ binding, compared to reduced asisolated CrHydA1, is unlikely. If H-species become bound to 2Fe_H, they may be located in a terminal position on Fe_d but not in an Fe-Fe bridging position. Further investigations are required to clarify this issue. CO binding to 2Fe_H, on the other hand, caused a significant elongation of the Fe-Fe distance, and additional short Fe-C(=0) distances were found. These results are comparable with crystal data of CO-treated bacterial [FeFe] hydrogenases (41) and previous EPR data on CrHydA1 (27). Presumably, the extra CO on Fe_d inhibits CrHydA1 by causing a structural rearrangement, which blocks a binding site for H-species, in a similar fashion to that proposed for bacterial enzymes (10, 11, 41).

In conclusion, we show that by XAS on algal [FeFe] hydrogenase the site geometry and electronic configuration of the H-cluster can be resolved. It is possible to monitor changes upon substrate and inhibitor binding. For the first time, the bond lengths and Fe-Fe distances of the H-cluster were determined at subangstrom resolution. We summarize our results on the structure of the H-cluster in as-isolated reduced and CO-treated CrHydA1 in the tentative models shown in Figure 4. Further investigations to characterize the active site of C. reinhardtii [FeFe] hydrogenase in its H₂ turnover cycle are in progress in our laboratories.

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