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# Electrochemical Kinetic Investigations of the Reactions of [FeFe]-Hydrogenases with Carbon Monoxide and Oxygen: Comparing the Importance of Gas Tunnels and Active-Site Electronic/Redox Effects

Gabrielle Goldet,<sup>†</sup> Caterina Brandmayr,<sup>†</sup> Sven T. Stripp,<sup>‡</sup> Thomas Happe,<sup>‡</sup> Christine Cavazza,<sup>§</sup> Juan C. Fontecilla-Camps,<sup>§</sup> and Fraser A. Armstrong<sup>\*,†</sup>

Inorganic Chemistry Laboratory, Department of Chemistry, University of Oxford, South Parks Road, Oxford OX1 3QR, United Kingdom, Ruhr-Universität, Lehrstuhl fur Biochemie der Pflanzen, AG Photobiotechnologie, 44780 Bochum, Germany, and Laboratoire de Crystallographie et Crystallographie des Protéines, Institut de Biologie Structurale, J.P. Ebel, CEA, CNRS, Université Joseph Fourier, 41, rue J. Horrowitz, 38027 Grenoble Cedex 1, France

Received July 3, 2009; E-mail: fraser.armstrong@chem.ox.ac.uk

**Abstract:** A major obstacle for future biohydrogen production is the oxygen sensitivity of [FeFe]-hydrogenases, the highly active catalysts produced by bacteria and green algae. The reactions of three representative [FeFe]-hydrogenases with O<sub>2</sub> have been studied by protein film electrochemistry under conditions of both H<sub>2</sub> oxidation and H<sub>2</sub> production, using CO as a complementary probe. The hydrogenases are *Dd*HydAB and *Ca*HydA from the bacteria *Desulfovibrio desulfuricans* and *Clostridium acetobutylicum*, and *Cr*HydA1 from the green alga *Chlamydomonas reinhardtii*. Rates of inactivation depend on the redox state of the active site 'H-cluster' and on transport through the protein to reach the pocket in which the H-cluster is housed. In all cases CO reacts much faster than O<sub>2</sub>. In the model proposed, *Ca*HydA shows the most sluggish gas transport and hence little dependence of inactivation rate on H-cluster state, whereas *Dd*HydAB shows a large dependence on H-cluster state and the least effective barrier to gas transport. All three enzymes show a similar rate of reactivation from CO inhibition, which increases upon illumination: the rate-determining step is thus assigned to cleavage of the labile Fe-CO bond, a reaction likely to be intrinsic to the atomic and electronic state of the H-cluster and less sensitive to the surrounding protein.

# Introduction

The increasing need for clean, renewable fuels is stimulating new research on hydrogen  $(H_2)$  production,  $^{1-5}$  and one promising solution is to exploit microorganisms in ' $H_2$  farms'. In biology,  $H_2$  is evolved by metalloenzymes called hydrogenases, in processes ranging from fermentation to photosynthesis. Hydrogenases are highly efficient enzymes—so much so that when attached to an electrode, they are, like platinum, superb electrocatalysts of both  $H_2$  oxidation and  $H_2$  production, at or close to the reversible potential for the  $2H^+/H_2$  couple.  $^{6-8}$ 

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- \* Ruhr-Universitat.
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Of the two main classes, [NiFe]- and [FeFe]-hydrogenases, named according to the metals present in the center at which H<sub>2</sub> is activated, the [FeFe]-hydrogenases are considered to be more active in H<sub>2</sub> production. However, a perceived major disadvantage of [FeFe]-hydrogenases (with respect to [NiFe]hydrogenases) is their higher O<sub>2</sub> sensitivity. 10 The [NiFe]hydrogenases react rapidly with O2 to give inactive, EPRcharacterized, Ni(III) forms that can be reactivated by reduction: Ni-A ('unready') is reactivated very slowly whereas Ni-B ('ready') can be reactivated within seconds, hence there is a rapid repair mechanism for hydrogenases that produce only Ni-B.<sup>6,11</sup> In contrast, the [FeFe]-hydrogenases appear to undergo irreparable damage when exposed to O2 while in their active state (after reduction). $^{12-14}$  The incompatibility of  $O_2$  with [FeFe]-hydrogenases poses a major limitation to progress in 'biohydrogen' production, in particular by modified photosyn-

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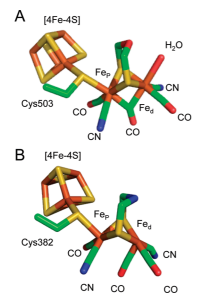
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**Figure 1.** Structures of H-clusters of the [FeFe]-hydrogenase from *Clostridium pasteurianum* (*Cp*I) and *Desulfovibrio desulfuricans* (*Dd*HydAB) constructed using PyMol. A) *Cp*I H-cluster (PDB code: 3C8Y). <sup>23</sup> B) *Dd*HydAB H-cluster. <sup>25</sup> The two structures were modeled with different bridgehead atoms—O for *Cp*I, and N for *Dd*HydAB—but this distinction is not directly relevant for this paper.

thesis.<sup>15</sup> In green algae the O<sub>2</sub>-sensitivity of the [FeFe]-hydrogenase is the bottleneck for producing H<sub>2</sub> from sunlight. Production of H<sub>2</sub> is stimulated during sulfur deprivation, conditions under which only 10% of the photosystem II remains active and the system effectively becomes anaerobic.<sup>16</sup> Green algae that could express an O<sub>2</sub>-tolerant [FeFe]-hydrogenase would therefore provide much increased levels of H<sub>2</sub> production.<sup>15</sup>

The buried active site of [FeFe]-hydrogenases is actually a complex 6Fe unit known as the 'H-cluster' which contains a [4Fe-4S] subcluster (generally referred to as [4Fe-4S]<sub>H</sub>) in addition to the di-iron subcluster (2Fe<sub>H</sub>). <sup>17</sup> The two independent representations of the H-cluster shown in Figure 1 are directly relevant to the catalytically active states known as Hox and Hred that have been extensively characterized. 18,19 General features of the structure are as follows: (a) the [4Fe-4S]<sub>H</sub> subcluster is linked to one of the Fe atoms of the 2Fe<sub>H</sub> subcluster by a bridging cysteine sulfur (the Fe atoms of 2Fe<sub>H</sub> are thus known as 'proximal' (Fe<sub>p</sub>) and 'distal' (Fe<sub>d</sub>) with respect to the [4Fe-4S<sub>H</sub> subcluster);<sup>20</sup> (b) both Fe<sub>d</sub> and Fe<sub>p</sub> are coordinated by CO and CN<sup>-</sup> ligands; <sup>17</sup> (c) an unusual SCH<sub>2</sub>XCH<sub>2</sub>S dithiolate ligand forms a di- $\mu$ -thiolato bridge between Fe $_p$  and Fe $_d$ , and although opinions differ as to whether the bridgehead atom X is an O or N atom, <sup>21–25</sup> recent investigations with <sup>14</sup>N HYSCORE have provided direct evidence that  $X = N.^{21}$ 

In the structure of  $H_{ox}$ , as determined with the CpI enzyme from Clostridium pasteurianum, Fep is coordinated by one CO and one CN<sup>-</sup> ligand and shares a bridging CO with Fe<sub>d</sub>.<sup>20</sup> In turn, Fe<sub>d</sub> is also coordinated by one CO and one CN<sup>-</sup> ligand, and an additional binding site is vacant or occupied by an exchangeable O-ligand, most likely a water molecule (Figure 1A). In the structure of the [FeFe]-hydrogenase from Desulfovibrio desulfuricans, which should be in the H<sub>red</sub> form, the bridging CO is replaced by a terminal CO on Fed (Figure 1B). 26 Recent EPR spectroscopic investigations favor an oxidation state assignment of [4Fe-4S]<sup>2+</sup>-Fe<sub>p</sub>(I)Fe<sub>d</sub>(II) for H<sub>ox</sub>, with some spin density delocalized onto the [4Fe-4S]<sub>H</sub> subcluster, <sup>27</sup> although H-clusters from different enzymes show minor variations in electronic structure. 28 The EPR-silent H<sub>red</sub> state is assigned as [4Fe-4S]<sup>2+</sup>-Fe(I)Fe(I) which, if protonated, is formally equivalent to the hydrido species [4Fe-4S]<sup>2+</sup>-Fe(II)Fe(II)-H<sup>-</sup>.<sup>17</sup> As also determined by EPR spectroscopy, exogenous CO, a competitive inhibitor of H2 oxidation, reacts with H<sub>ox</sub>. <sup>29</sup> Crystallographic and infrared spectroscopic studies of  $H_{ox}$ -CO further show that binding of CO (which is photolabile) occurs at  $Fe_d$ . Inactivation by anaerobic oxidants gives rise to a form known as Hox inact, usually formulated as [4Fe-4S]<sup>2+</sup>-Fe(II)Fe(II), which can be reactivated upon reduction—a process occurring via an intermediate known as H<sub>trans</sub> which has been formulated as [4Fe-4S]<sup>+</sup>-Fe(II)Fe(II), i.e. with the [4Fe-4S]<sub>H</sub> subcluster reduced.<sup>33</sup> The H-cluster is remarkable among non-macrocycle cofactors because the 2Fe<sub>H</sub> subcluster at which H<sub>2</sub> is produced is connected to the protein through just a half-share of a cysteine sulfur: it is very much an organometallic-like compound physically enclosed in protein.

Despite these intense studies by crystallography and spectroscopy, numerous aspects of the reactions of [FeFe]-hydrogenases remain unresolved. These aspects include the activation process (there is evidence from electrochemical titrations that a two-electron process is also involved)<sup>33</sup> and many details of the mechanism of catalysis in either direction, including the exact function of the [4Fe-4S]<sub>H</sub> subcluster. The nature and products of the degradation by  $O_2$  are only now coming to light,<sup>34</sup> and a major issue is whether and how  $H_2$  production could be sustainable at all in the presence of  $O_2$ .

This article describes mechanistic investigations, by protein film electrochemistry, of the O<sub>2</sub> inactivation kinetics of three representative [FeFe]-hydrogenases. These are: the hydrogenase

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from a sulfate-reducing bacterium, Desulfovibrio desulfuricans, abbreviated as *Dd*HydAB which has been crystallographically characterized;<sup>26</sup> the hydrogenase from Clostridium acetobutylicum, abbreviated as CaHydA, which is potentially of importance for H<sub>2</sub> production by anaerobic fermentation and has high sequence similarity with the crystallographically characterized CpI hydrogenase from C. pasteurianum, 22 and the hydrogenase known as CrHydA1 from the green alga Chlamydomonas reinhardtii, which is of interest for photosynthetic H2 production. 35 Both *Dd*HydAB and *Ca*HydA contain a series of Fe-S clusters<sup>17</sup> to relay electrons within the protein for transfer to and from the redox partner (in our case the electrode); in contrast CrHydA1 possesses no Fe-S clusters apart from the [4Fe-4S]<sub>H</sub> subcluster, 36,37 but it is nonetheless electrocatalytically active when adsorbed on an electrode.<sup>34</sup> Although the three enzymes differ in their overall tertiary and quaternary structures, their H-domains that house the H-cluster are very similar.<sup>17</sup>

In protein film electrochemistry, an enzyme is immobilized on the surface of an electrode such that its properties are controlled directly by the electrode potential. 6,38 Catalytic activity in either direction, oxidation or reduction, can be driven and recorded at any particular potential value and the catalytic rate is directly proportional to the current that flows. Various gas mixtures (produced by mass-flow controllers) can be introduced and flushed from the sealed electrochemical cell in which the electrode is rotated rapidly to provide precise hydrodynamic control (supply of reactants and removal of products). A particular advantage of this approach is that turnover activity in either direction is immediately and directly observed from the catalytic current; thus, rates of change of activity, such as those induced by CO or O2, are extracted directly from the variation of current with time, all as a precise function of the electrode potential. In this way, extremely complex reactivities become resolvable; therefore, this technique both complements and instigates structural and spectroscopic

We first establish, for each enzyme, how H<sub>2</sub> oxidation and  $H_2$  production are affected by the concentrations of  $H_2$  and CO; next we examine the kinetics of CO binding and release and correlate these data with equilibrium values; we then examine the kinetics of O<sub>2</sub> inactivation of H<sub>2</sub> oxidation activity; finally, we exploit CO inhibition as a tool to investigate H<sub>2</sub> production in the presence of O<sub>2</sub>. Recent studies have established that CO is able to protect [FeFe]-hydrogenases against inactivation by O<sub>2</sub>, <sup>34,39</sup> an observation suggesting that the sequence of destruction is initiated by  $O_2$  coordinating to the same site at which exogenous CO binds, i.e.  $Fe_d$ . Thereafter, the mechanism remains less clear, but recent EXAFS evidence obtained with CrHydA1 shows that the [4Fe-4S]<sub>H</sub> subcluster is altered more than the 2Fe<sub>H</sub> subcluster.<sup>34</sup> Our experiments show clearly how the destructive power of O<sub>2</sub> varies among the hydrogenases with an interesting dependence on catalytic direction (H<sub>2</sub> oxidation compared to H<sub>2</sub> production)—thus implicating sensitivity to the oxidation level of the active site. The results provide insight for the quest for solutions to the oxygen problem in biohydrogen production by photosynthesis. The functional differences between the [FeFe]-hydrogenases are not immediately evident from the structures that have been obtained for DdHydAB and Cp1.

## Methods

Previously reported protocols were employed to obtain pure samples of *Dd*HydAB, <sup>40</sup> *Cr*HydA1 and *Ca*HydA. <sup>41</sup> In each case levels of CO, O2 and H2 present during cell growth were extremely low. Protein film electrochemistry experiments were carried out in an anaerobic glovebox (M Braun) comprising a N2 atmosphere. The solutions contained 0.05 M phosphate buffer with 0.10 M NaCl as additional supporting electrolyte, and were prepared using standard reagents NaCl, NaH2PO4 and Na2HPO4 (Analytical Reagent grade, Sigma) in purified water (Millipore 18 M $\Omega$  cm). The working electrode was a disk (area 0.03 cm<sup>2</sup>) of pyrolytic graphite oriented so that the 'edge plane' faced the solution. This is called a pyrolytic graphite edge (PGE) electrode and it was used in conjunction with an electrode rotator (EcoChemie) that fitted snugly into a specially designed, gastight, glass electrochemical cell. The cell featured a water jacket for temperature control. Due to the light sensitivity of [FeFe]-hydrogenases the entire surface of the cell was blacked-out with adhesive masking tape. In experiments to detect photolabilization, the tape at the bottom of the cell was removed to allow illumination from a 150 W lamp placed just beneath the cell (see Figure SI.1, Supporting Information). A saturated calomel reference electrode (SCE) was placed in a side arm containing 0.1 M NaCl, connected to the main cell compartment by a Luggin capillary. A Pt wire was used as the counter electrode. Potentials (E) are quoted with respect to the standard hydrogen electrode (SHE) using the correction  $E_{SHE} =$  $E_{\rm SCE}$  + 242 mV at 298 K.<sup>42</sup> Electrochemical experiments were performed using an electrochemical analyzer (Autolab PGSTAT10 or 20) controlled by a computer operating GPES software (Eco-Chemie). Mass flow controllers (Smart-Trak Series 100, Sierra Instruments, U.S.A.) were used to prepare precise gas mixtures (headspace fractions accurate to within 1%) and to impose constant gas flow rates into the electrochemical cell during experiments. Gases used were H2 (Premier grade, Air Products), O2 (Air Products), CO (Research grade, BOC), 1% CO in N<sub>2</sub> (Research grade, BOC), N2 (Oxygen-free, BOC) or mixtures of these gases. Henry's law was applied to estimate the concentration of gas in solution in each experiment. 43 Values taken for use at 10 °C were: 100% CO, 1248 μM; 100% O<sub>2</sub>, 1624 μM.

To prepare each enzyme film, the PGE electrode was first polished for 10 s with an aqueous slurry of  $\alpha$ -alumina (1  $\mu$ m, Buehler) and sonicated for 5 s in purified water, before enzyme solution (1.5  $\mu$ L, containing 0.015–0.15  $\mu$ g, pH 8) was applied and removed after a few minutes. The electrode was then placed in enzyme-free buffered electrolyte so that the only enzyme molecules being addressed were on the electrode and subjected to the same regime of strict potential control. In all experiments the electrode was rotated at a constant high rate (3000–5000 rpm) to ensure efficient supply of substrate and removal of product.

In experiments measuring the effect of  $\mathrm{O}_2$  on hydrogenase-catalyzed  $\mathrm{H}_2$  production, it was necessary to estimate the concentration of  $\mathrm{O}_2$  that the enzyme molecules at the electrode surface actually experience, given that a portion of the total dissolved  $\mathrm{O}_2$  is consumed by the electrode at this potential (-0.4 V). Using a

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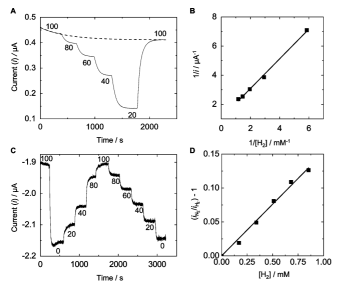
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**Figure 2.** Determination of  $K_{\rm M}^{\rm H_2}$  (A and B) and  $K_{\rm I}^{\rm app}$  (C and D) for  $Ca{\rm HydA}$ . (Panel A) chronoamperometric experiment performed at -0.05 V, at pH 6, 10 °C. The broken line traces the progression of film loss throughout the experiment. (Panel B) Lineweaver—Burk plot from experiment shown in A fitted to a straight line. (Panel C) chronoamperometric experiment performed to determine  $K_{\rm I}^{\rm app}$  at -0.4 V, at pH 6.0, 10 °C. The concentration of  $H_2$  in  $N_2$  (%) in the headspace of the cell is indicated. The rotation rate was varied from 3000 to 4000 and 5000 rpm at each concentration of  $H_2$  to ensure that inhibition was not mass-transport limited. (Panel D) Plot according to the procedure described by Léger et al. <sup>45</sup> from which  $K_{\rm I}^{\rm app}$  is determined, showing the line of best fit.

previously published method, <sup>44</sup> it was estimated that 0.7% O<sub>2</sub> (11  $\mu$ M in solution at 10 °C) survives to be experienced by the enzyme when 1% O<sub>2</sub> is flowed through the headspace of the cell and 4.5% (73  $\mu$ M) survives when 5% O<sub>2</sub> is used.

# Results

Measurements of the H<sub>2</sub> Concentration Dependencies for H<sub>2</sub> Oxidation and Production. We first evaluated the affinity of the enzymes for  $H_2$  both in terms of  $K_M$  for  $H_2$  as the substrate in H<sub>2</sub> oxidation  $(K_{\rm M}^{\rm H_2})$  and the apparent inhibition constant  $K_{\rm I}^{\rm app}$ for H<sub>2</sub> as the product inhibitor of H<sup>+</sup> reduction. Figure 2 shows experiments carried out for CaHydA: analogous experiments were carried out for CrHydA1, but films of DdHydAB were not sufficiently stable to obtain accurate measurements over the period of time required (30-60 min.). Experiments to determine  $K_{\rm M}^{\rm H_2}$  were conducted by varying the ratio of H<sub>2</sub> to N<sub>2</sub> in the headgas and measuring the oxidation current after allowing time for the gas mixture to equilibrate with the cell solution at each concentration of H<sub>2</sub> (Figure 2A). The experiments were performed at an electrode potential of -0.05 V to avoid the anaerobic inactivation that occurs at higher potential (see Figure 3). We and others, have noted that  $K_{\rm M}$  and  $K_{\rm I}$  are potential-dependent quantities, <sup>44,45</sup> and the potential must therefore be specified. It was also important to make measurements under conditions where the current was limited by the catalytic rate of the enzyme rather than by mass transport of substrate to (or product from) the electrode. The experiments were therefore carried out at low temperature (10 °C) to decrease the rate of catalysis. The low temperature also minimized the

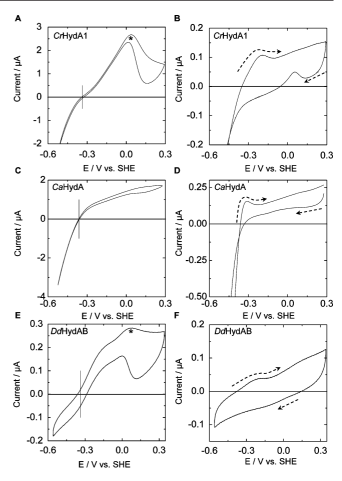


Figure 3. Cyclic voltammograms showing bidirectional electrocatalytic H+ reduction and H2 oxidation by CrHydA1 (A), CaHydA (C) and DdHydAB (E) at pH 6.0 under 1 bar H<sub>2</sub>. Panels B, D, F show the timedependent voltammograms recorded following introduction of CO, introduced prior to the scans shown here, as the potential was being cycled between -0.55 V and +0.3 V, as it is removed from the cell: these reveal different reactions as CO dissociates and rebinds as a function of potential. Experimental conditions for CrHydA1: 10 °C, scan rate 20 mV/s, inhibition was achieved by injection of CO-saturated buffer to give an immediate concentration of  $100 \,\mu\mathrm{M}$  CO in solution 5 min prior to the scan shown in panel B. Experimental conditions for CaHydA: 32 °C, electrode rotation 3000 rpm, scan rate 20 mV/s, inhibition was achieved by flushing 100% CO through the cell 5 min prior to the scan shown in panel D. Experimental conditions for DdHydAB: 10 °C, electrode rotation 3000 rpm, scan rate 10 mV/s, inhibition was achieved by injection of CO-saturated buffer to give an instant concentration of 30  $\mu$ M CO in solution 5 min prior to the scan shown in panel F. The dashed arrows indicate the direction of scanning. The asterisks (\*) indicate the potential above which anaerobic inactivation takes place for CrHydA1 and DdHydAB.

rate of film loss. At each  $H_2$  concentration the electrode rotation rate was stepped between 3000 and 5000 rpm to ensure that the current (and thus the rate of reaction) was independent of rotation rate. Values of  $K_M^{H_2}$  were calculated from the *x*-intercept of the Lineweaver–Burk plot shown in Figure 2B and are summarized in Table 1. Experiments to measure the apparent inhibition constant ( $K_1^{app}$ , defined in eq 1 where  $K_M^{H^+}$  is the  $K_M$ 

**Table 1.** Values of  $K_{\rm M}^{\rm H_2}$  and  $K_{\rm I}^{\rm app}$  for  $Cr{\rm HydA1}$  and  $Ca{\rm HydA}$  Determined at pH 6, 10 °C from the Experiments Shown in Figure

enzyme	$K_{\rm M}^{\rm H}2/{\rm mM}$	$K_{l}^{app}/mM$
<i>Cr</i> HydA1	$0.19 \pm 0.03$	$3.7 \pm 0.6$
<i>Ca</i> HydA	$0.46 \pm 0.06$	$6.2 \pm 1.1$

 $<sup>^{</sup>a}$   $K_{\rm M}{}^{\rm H_{2}}$  and  $K_{\rm I}{}^{\rm app}$  were calculated at -0.05 and -0.4 V, respectively.

<sup>(44)</sup> Goldet, G.; Wait, A. F.; Cracknell, J. A.; Vincent, K. A.; Ludwig, M.; Lenz, O.; Friedrich, B.; Armstrong, F. A. J. Am. Chem. Soc. 2008, 130, 11106–11113.

<sup>(45)</sup> Léger, C.; Dementin, S.; Bertrand, P.; Rousset, M.; Guigliarelli, B. J. Am. Chem. Soc. 2004, 126, 12162–12172.

for binding of the substrate  $H^+$  and  $K_1$  is the real inhibition constant) were conducted at -0.4 V vs SHE using a similar experimental method (Figure 2C). Values of  $K_1^{app}$  were calculated by adapting the method based on that reported by Léger et al.<sup>45</sup> which requires plotting the data according to eq 2 (in which  $i_{N_2}$  is the current recorded under 100%  $N_2$  and  $i_{H_2}$  is the current recorded at each concentration of  $H_2$ ) as shown in Figure 2D. The  $K_1^{app}$  values are included in Table 1.

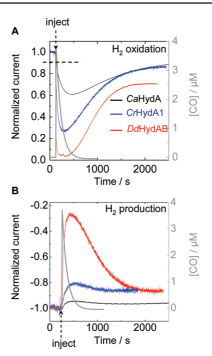
$$K_{\rm I}^{\rm app} = \frac{K_{\rm I}[{\rm H}^+]}{K_{\rm M}^{\rm H^+}} \left(1 + \frac{K_{\rm M}^{\rm H^+}}{[{\rm H}^+]}\right) \tag{1}$$

$$(i_{N_2}/i_{H_2}) - 1 = \frac{[H_2]}{K_1^{app}}$$
 (2)

The  $K_{\rm M}^{\rm H_2}$  values are much higher than those we have measured recently for some O<sub>2</sub>-tolerant [NiFe]-hydrogenases<sup>46</sup> and they suggest that the catalytic current for H<sub>2</sub> oxidation will be sensitive to changes in H<sub>2</sub> levels even close to 1 bar partial pressure (this is particularly so for CaHydA). The results showed that headspace H<sub>2</sub> levels should be maintained constant in quantitative H<sub>2</sub> oxidation experiments. In contrast, the  $K_{\rm I}^{\rm app}$  values are so high that the presence of H<sub>2</sub> is not expected to pose a problem in studies of H<sub>2</sub> production. The order of magnitude difference between  $K_{\rm I}^{\rm app}$  and  $K_{\rm M}^{\rm H_2}$  suggests immediately that the electronic/catalytic state of the H-cluster exerts a strong effect on binding affinity, with H<sub>2</sub> binding more weakly to a more reduced state.

Reactions with CO. Cyclic voltammograms of the electrocatalytic activities of DdHydAB, CrHydA1 and CaHydA at pH 6.0 under a flow of 100% H<sub>2</sub> are shown in Figure 3 (A, C, E—left-hand column). These voltammograms show that all the enzymes are bidirectional, with CaHydA being particularly biased in the direction of H<sub>2</sub> production. The traces in either potential direction cut through the zero current axis at the cell potential (see vertical line) for the 2H<sup>+</sup>/H<sub>2</sub> couple. At high potentials, all three enzymes undergo inactivation to give a species that is most likely  $H_{ox}^{inact}$  although this process is very slow for CaHydA. The voltammograms in the right-hand column (B, D, F) were recorded during the efflux of CO that had been introduced by injecting a saturated solution (giving a concentration of 100  $\mu$ M CO in the cell for the experiment on CrHydA1, 30  $\mu M$  for the experiment on DdHydAB) or flowing 100% CO briefly through the cell (for CaHydA) as the potential was cycled between -0.55 and 0 V prior to the scans shown in panels B, D, and F.

Informative changes in the voltammograms of *Dd*HydAB, CrHydA1 and CaHydA occur upon removal of CO during continuous cycling (B, D, F). In all cases the voltammograms show that once the potential is made sufficiently positive to start H<sub>2</sub> oxidation, there is an initial increase in current which is followed by a decrease (see the dashed arrow indicating the oxidative sweep). The magnitude of this effect is most apparent after a certain time has elapsed, dependent upon the hydrogenase, and finally the voltammograms resume the expected shapes for H<sub>2</sub> oxidation and production at pH 6 under 1 bar H<sub>2</sub>, analogous to those shown in the left-hand column. This cyclic 'inhibitor-on/inhibitor-off' behavior shows that CO binding to the active site is favorable and fast during H<sub>2</sub> oxidation but relatively weak during H<sub>2</sub> production. Thus CO reinhibits strongly as the potential is raised to oxidize H<sub>2</sub>, which addresses a more oxidized form of the enzyme (above -0.3 V). The



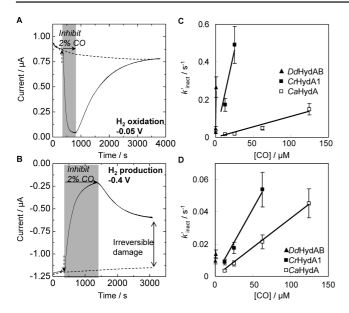
**Figure 4.** Inhibition of *Dd*HydAB (red lines), *Cr*HydA1 (blue lines) and *Ca*HydA (black lines) by injection of 4 μM CO at (A) -0.05 V (H<sub>2</sub> oxidation) and (B) -0.4 V (H<sub>2</sub> production). Experimental conditions: pH 6.0, 10 °C, 1 bar H<sub>2</sub>, electrode rotation 3000 rpm. The final level of the current reached upon recovery of H<sub>2</sub> oxidation activity of *Ca*HydA is marked by the dashed line in A. The exponential decrease in concentration of dissolved CO is shown by the gray trace (details given in Supporting Information, Figure SI.2).

voltammogram for CrHydA1 also shows an oxidation peak soon after commencing the scan in the negative direction. A similar observation was reported in previous experiments on DdHydAB where it arises from the reactivation of some  $H_{ox}^{inact}$  and its rapid inactivation by CO.<sup>47</sup>

Further insight into CO binding and release under different conditions is provided by Figure 4. The CO inhibition profiles for *Dd*HydAB, *Cr*HydA1 and *Ca*HydA were obtained for H<sub>2</sub> oxidation (Panel A) or H<sub>2</sub> production (Panel B) by injecting an aliquot of CO-saturated solution and then recording the catalytic current as the CO is removed by flushing. The same experimental conditions (pH 6.0, 10 °C, 1 bar H<sub>2</sub>, electrode rotation 3000 rpm) were used for all experiments. In each case, injection of CO-saturated buffer gives an immediate initial CO concentration of 4  $\mu$ M which decreases exponentially to zero, 45 as depicted by the gray trace (right y-axis) which represents the dependence of [CO] on time throughout the experiment; < 0.2 μM CO remains in solution after 500 s. (See Supporting Information (Figure SI.2) for how this dependence is determined.) In the case of CaHydA, inhibition continues to increase even though most of the CO has been flushed out of the cell, indicating that the rate of reaction with CO is slow. For H<sub>2</sub> oxidation, the rates and extent of inhibition reached after CO injection decrease in the order DdHydAB > CrHydA1 > CaHydA. The reactivation rates are slow and strikingly similar for all enzymes. For H<sub>2</sub> production (panel B) the rates and extent of inhibition by CO again decrease in the order DdHydAB > CrHydA1 > CaHydA.

<sup>(46)</sup> Ludwig, M.; Cracknell, J. A.; Vincent, K. A.; Armstrong, F. A.; Lenz, O. J. Biol. Chem. 2009, 284, 465–477.

<sup>(47)</sup> Parkin, A.; Cavazza, C.; Fontecilla-Camps, J. C.; Armstrong, F. A. J. Am. Chem. Soc. 2006, 128, 16808–16815.



**Figure 5.** Inhibition by CO and reactivation, observed for  $H_2$  oxidation and  $H_2$  production with DdHydAB, CrHydA1 and CaHydA. Panels A and B show experiments performed on CaHydA at -0.05 and -0.4 V, respectively, in which the headgas is 80%  $H_2$ , 20%  $N_2$  at the start of the experiment. Carbon monoxide (2%, approximately 25 μM) was introduced by injection of CO-saturated buffer (0.5 mL of buffer saturated with 10% CO, 80%  $H_2$ , 10%  $N_2$ ) into the 2 mL of buffer already present in the cell and, simultaneously, flushing the headspace with 80% $H_2$ , 18%  $N_2$ , 2% CO for the period of time marked by the gray boxes. Experimental conditions: pH 6.0, electrode rotation 3000 rpm, potentials as indicated. Panels C and D show the dependencies of rates ( $k'_{inact}$ ) on CO concentration; data are derived from experiments such as those shown in panels A and B, respectively, for DdHydAB, CrHydA1 and CaHydA.

To obtain the kinetics of CO binding for the three [FeFe]hydrogenases it was necessary to maintain a constant CO concentration throughout the time scale of the reaction. This was achieved by injecting an aliquot of solution containing CO and simultaneously changing the gas composition reaching the headspace by replacing, with CO, a certain fraction of the 20% N<sub>2</sub> component of the 80% H<sub>2</sub> mixture. Exemplary results obtained with CaHydA are shown in Figure 5. Panel A shows the time-course for CO inhibition of  $H_2$  oxidation at -0.05 V, and Panel B shows the same time-course for inhibition of H<sup>+</sup> reduction at -0.4 V. Both experiments commenced with the hydrogenase on the electrode being exposed to an atmosphere of 80%  $H_2$ , 20%  $N_2$ . At t = 300 s, an aliquot of CO-containing solution was injected to take the cell concentration to 2% CO, and the cell was flushed with 80% H<sub>2</sub>, 18% N<sub>2</sub>, 2% CO. Once the catalytic current had decreased to a steady level, the CO was removed from the cell by flushing with 80% H<sub>2</sub>, 20% N<sub>2</sub>. (Note that 80% H<sub>2</sub> remained in the headspace of the cell throughout the experiment. This excludes variations in catalytic current resulting simply from changes on H2 concentration, which would otherwise pose a problem, given the high  $K_{\rm M}$  of these enzymes as indicated in Panel 1.) Pseudo first-order rate constants ( $k'_{inact}$ ) were determined for each experiment from the slope of plots of ln(i) vs t, with linearity generally exceeding 75% for H<sub>2</sub> oxidation, although less so for H<sub>2</sub> production. Panels C and D show the corresponding dependencies of  $k'_{inact}$  on CO concentration, obtained for DdHydAB, CrHydA1 and CaHydA by numerous experiments conducted analogously to those shown in panels A and B.

Values for the equilibrium inhibition constant  $K_1^{CO}$ (equil) were determined for CrHydA1 and CaHydA by measuring the  $H_2$  oxidation current stabilized at different concentrations of CO.

The term  $K_1^{\rm CO}({\rm equil})$  depends on  ${\rm H_2}$  concentration and potential, but we kept these variables constant (80%  ${\rm H_2}, -0.05$  V). These titrations were similar in design to those shown in Figure 2C and are described in more detail in Supporting Information (Figure SI.3). Values of  $K_1^{\rm CO}({\rm equil})$  are shown in Table 2. Comparable data for  $Dd{\rm HydAB}$  could not be obtained because the enzyme was fully inhibited at the lowest practical CO concentrations.

Panel B of Figure 5 shows that CO inhibition of  $H_2$  production of these enzymes is only partially reversible. This observation was consistently made with CaHydA and CrHydA1 but the instability of DdHydAB films prevented us from making similar measurements with this enzyme. By comparison, CO inhibition is fully reversible when measuring  $H_2$  oxidation at -0.05 V. In all cases, the background decrease in current that we refer to as film loss (traced by the dashed line) was checked through control experiments carried out without introducing CO. The rates determined for the inactivation of  $H_2$  production by CO are therefore approximate. The potential dependence of the rate of inactivation of  $H_2$  production was not investigated further, although we noted that the inactivation process became noticeably biphasic as the potential was lowered below -0.4 V. We raise this issue later in the Discussion.

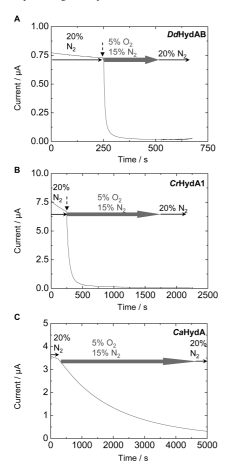
Panels C and D of Figure 5 show that: (i) DdHydAB is always the fastest to react with CO and CaHydA the slowest, and ii) for all three enzymes, inhibition of H<sub>2</sub> oxidation measured at −0.05 V is substantially faster than inhibition of H<sub>2</sub> production at -0.4 V, but the same order DdHydAB > CrHydA1 >CaHydA is observed in both catalytic directions. The secondorder rate constants ( $k_{\text{inact}}$ ) are provided in Table 2 (see later). In all cases, the reactivation rates ( $k_{re-act}$ ) are the same, within reasonable error, for H<sub>2</sub> oxidation and H<sub>2</sub> production. Experiments carried out with CrHydA1 and CaHydA showed that  $k_{\text{re-act}}$  is strongly light sensitive (see Supporting Information, Figure SI.1) in agreement with earlier electrochemical observations made with DdHydAB.<sup>47</sup> In addition,  $k_{re-act}$  increased significantly when the temperature was raised to 25 °C. From the ratio of rate constants for the reactivation and inactivation reactions at 10 °C, we derived the kinetic inhibition constants  $K_{\rm I}^{\rm CO}({\rm kin}) = k_{\rm re-act}/k_{\rm inact}$ . In Table 2,  $K_{\rm I}^{\rm CO}({\rm kin})$  values are compared with the equilibrium values,  $K_1^{CO}$  (equil), obtained by titration for CrHydA1 and CaHydA. The two values for CrHydA1 are in good agreement, although for CaHydA, K<sub>I</sub><sup>CO</sup>(equil) is rather higher than  $K_{\rm I}^{\rm CO}({\rm kin})$ . The ratio of the  $k_{\rm inact}$  values for CO inhibition of H<sub>2</sub> oxidation and H<sub>2</sub> production—the "catalytic direction discrimination"- is about 49 for *Dd*HydAB, 22 for CrHydA1 and 3 for CaHydA: these variations clearly arise from differences in the rate that CO binds because  $k_{\text{re-act}}$  is similar for all three enzymes.

Reactions with  $O_2$ . Figure 6 shows experiments in which each enzyme is subjected to 5%  $O_2$  during  $H_2$  oxidation at -0.05 V vs SHE. As before, the  $H_2$  concentration was kept constant throughout the entire time-course. Each experiment began with 80%  $H_2$ , 20%  $N_2$  flushing through the cell headspace. The gas mixture was then switched to 80%  $H_2$ , 15%  $N_2$ , 5%  $O_2$  for the duration of the reaction (this involved simultaneous injection of an aliquot of gas-equilibrated buffer for DdHydAB and CrHydA1; the kinetics of  $O_2$  inactivation of CaHydA were so slow that the injection was unnecessary). The gas was finally changed back to 80%  $H_2$ , 20%  $N_2$  once most of the activity (>90%) had been eliminated. We noted that the rate of inactivation by  $O_2$  showed a dependence on  $H_2$  concentration, with inactivation occurring more rapidly at lower levels, thus

**Table 2.** Compilation of  $k_{\text{inact}}$  and  $k_{\text{re-act}}$  Values for CO Inhibition of H<sub>2</sub> Production and H<sub>2</sub> Oxidation for *Da*HydAB, *Cr*HydA1 and *Ca*HydA, with the Corresponding Catalytic Direction Discrimination Factors and Values of  $K_1^{\text{CO}}$  Determined by Kinetic and Equilibrium Methods<sup>a</sup>

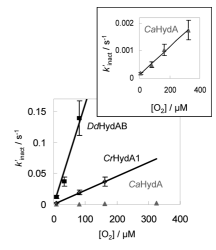
	<i>Dd</i> HydAB	CrHydA1	<i>Ca</i> HydA
$H_2$ oxidation $k_{inact}/s^{-1}\mu M^{-1}$	$3.9 \times 10^{-1} $ $\pm 3 \times 10^{-1}$	$1.9 \times 10^{-2}$ $\pm 1 \times 10^{-2}$	$1.1 \times 10^{-3}$ $\pm 7 \times 10^{-4}$
$H_2$ production $k_{inact}/s^{-1}\mu M^{-1}$	$8.0 \times 10^{-3}$ $\pm 7 \times 10^{-3}$	$8.4 \times 10^{-4} $ $\pm 2 \times 10^{-4}$	$3.6 \times 10^{-4} \\ \pm 1 \times 10^{-4}$
Catalytic direction discrimination $H_2$ oxidation $k_{\text{re-act}}/s^{-1}$	$1.9 \times 10^{-3} \pm 2 \times 10^{-4}$	$\begin{array}{c} 22 \\ 2.3 \times 10^{-3} \\ \pm 1 \times 10^{-3} \end{array}$	$   \begin{array}{c}     3 \\     1.8 \times 10^{-3} \\     \pm 1 \times 10^{-3}   \end{array} $
$H_2$ production $k_{\text{re-act}}/s^{-1}$	$\begin{array}{l} 2.7 \times 10^{-3} \\ \pm 1 \times 10^{-3} \end{array}$	$1.2 \times 10^{-3}$ $\pm 3 \times 10^{-3}$	$2 \times 10^{-3}$ $\pm 2 \times 10^{-4}$
$K_{\rm I}^{\rm CO}$ (equil)/ $\mu M$ for $H_2$ oxidation (at $-0.05~{\rm V}$ )	_	$1.0 \times 10^{-1}$	$2.2 \times 10^{-1}$
$K_{\rm l}^{\rm CO}$ (kin)/ $\mu M$ for H <sub>2</sub> oxidation (at $-0.05$ V)	$4.8 \times 10^{-3}$	$1.2 \times 10^{-1}$	1.6
$K_{\rm I}^{\rm CO}$ (kin)/ $\mu M$ for $H_2$ production (at $-0.40~{ m V}$ )	0.34	1.4	5.6

<sup>&</sup>lt;sup>a</sup> All values pertaining to H₂ production were measured at −0.40 V.



**Figure 6.** Inactivation of H<sub>2</sub> oxidation by *Dd*HydAB, *Cr*HydA1 and *Ca*HydA by 5% O<sub>2</sub>. The headspace mixture is composed of 80% H<sub>2</sub> and the remaining 20% is as indicated. For *Dd*HydAB and *Cr*HydA1 a 0.67 mL aliquot of 20% O<sub>2</sub>/80% H<sub>2</sub>-saturated buffer was injected into the cell which initially contained 2 mL of buffer, to give an instant concentration of 5% O<sub>2</sub> at the points marked by the dashed arrows. For *Ca*HydA, the reaction was sufficiently slow that it could be initiated simply by changing the headgas mixture. Other conditions: pH 6.0, 10 °C, electrode rotation 3000 rpm, -0.05 V.

indicating that O<sub>2</sub> and H<sub>2</sub> are competitive.<sup>34</sup> We also noted that after removing O<sub>2</sub> from the cell, a very small amount of activity was consistently recovered for all three enzymes. Oxygen



**Figure 7.** Dependence of rate constants ( $k'_{\text{inacl}}$ ) for inactivation of enzymatic  $H_2$  oxidation by  $O_2$ , on  $O_2$  concentration, for DdHydAB, CrHydA1 and CaHydA (shown as an expanded scale). Experimental conditions: pH 6.0, 10 °C, electrode rotation 3000 rpm, -0.05 V vs SHE, headspace gas mixture composed of 80%  $H_2$ , 20% mixture of  $N_2$  and  $O_2$ .

undergoes very slow reduction at graphite at -0.05 V, therefore control experiments (Figure SI.4, Supporting Information) were performed to assess the contribution to the current from  $O_2$  reduction. This contribution to the current was then subtracted from the experiments to verify the end point and the activity remaining. In separate anaerobic experiments we determined that small quantities of hydrogen peroxide that would be formed during electrodic  $O_2$  reduction did not cause inactivation, by injecting aliquots of  $H_2O_2$  solution and monitoring the effect on the  $H_2$  oxidation current at -0.05 V (10 °C). This decrease in current was observed to be much slower than that due to equivalent concentrations of  $O_2$  for all three enzymes.

Figure 7 shows how rate constants for  $O_2$ -inactivation of  $H_2$  oxidation activity vary with  $O_2$  concentration for all three [FeFe]-hydrogenases. In each case the rate of inactivation is first-order in  $O_2$  concentration. Table 3 shows the rate constants for inactivation by  $O_2$  alongside the rate constants for inhibition of  $H_2$  oxidation by CO. For each enzyme, the rate of reaction with CO is much faster (80–200-fold) than with  $O_2$  and this ratio, which we refer to as the "gas identity discrimination", is also included in Table 3.

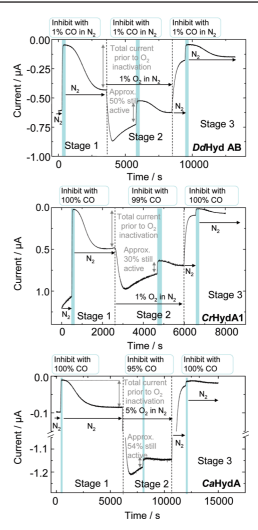
**Table 3.** Comparison of the Second-Order Rate Constants for Inhibition by CO ( $k_{\text{inact}}(\text{CO})$ ) and Inactivation by O<sub>2</sub> ( $k_{\text{inact}}(\text{O}_2)$ ) of DdHydAB, CrHydA1 and CaHydA and Evaluation of the Gas Identity Discrimination—the Ratio  $k_{\text{inact}}(\text{CO})/k_{\text{inact}}(\text{O}_2)$ 

enzyme	$k_{\mathrm{inact}}(\mathrm{CO})/\mathrm{s}^{-1}\mu\mathrm{M}^{-1}$	$k_{\mathrm{inact}}(\mathrm{O_2})/\mathrm{s^{-1}}\mu\mathrm{M}^{-1}$	$k_{\text{inact}}(\text{CO})/k_{\text{inact}}(\text{O}_2)$
DdHydAB	$3.9 \times 10^{-1}$	$1.8 \times 10^{-3} \pm 3 \times 10^{-4}$ $2.2 \times 10^{-4} \pm 1 \times 10^{-4}$ $5.1 \times 10^{-6} \pm 1 \times 10^{-6}$	217
CrHydA1	$1.9 \times 10^{-2}$		86
CaHydA	$1.1 \times 10^{-3}$		216

 $H_2$  Production in the Presence of  $O_2$ . Finally, we carried out experiments to estimate the extent to which the rate of inactivation by  $O_2$  depends on whether the hydrogenase is operating in the direction of  $H_2$  production or  $H_2$  oxidation. We used a procedure described recently,<sup>44</sup> in which the problem of distinguishing the current due to enzymatic  $H^+$  reduction from that due to electrochemical  $O_2$  reduction is resolved by adding an inhibitor. The decrease in current observed when the inhibitor is added provides a direct measure of the component of the current due to enzyme-catalyzed  $H^+$  reduction. The experiments conducted for DdHydAB, CrHydA1 and CaHydA are shown in Figure 8.

The experiment performed on DdHydAB started under an atmosphere of N<sub>2</sub>, and a current corresponding to enzymecatalyzed H<sub>2</sub> production was recorded (stage 1). The headgas was then switched to 1% CO in N2, and a rapid and almost complete loss of current was observed. When the CO was flushed from the cell, the current increased and reached a steady level (albeit not the same level as prior to CO introduction, due to the partial irreversibility of this reaction, see Figure 4). The current that was recovered during CO efflux was adopted as the normalization unit for the next stage. In stage 2, O<sub>2</sub> was introduced: the current initially increased due to direct reduction of O<sub>2</sub> at the graphite electrode but then began to decrease as the enzyme became inactivated. Introduction of 1% CO after 2000 s under O<sub>2</sub> resulted in a rapid loss of current, the magnitude of which reports on the enzyme-catalyzed current component prior to CO inhibition. Removal of CO caused the current to increase again, but a further decrease in current was observed, as expected, when  $O_2$  was removed from the cell after t =8500 s. Finally in stage 3 the inhibition step with CO was repeated to establish the extent of survival of the hydrogenase. Similar sequences of steps were used in the experiments with CrHydA1 and CaHydA, except that 99% and 95% CO, respectively, were used instead of 1% CO in order to compensate for the much slower kinetics and lower CO affinity of these hydrogenases compared to DdHydAB; in addition, 5% O<sub>2</sub> was used to obtain a higher rate of inactivation for CaHydA. To estimate the half-life for inactivation in each case, the decrease in current upon addition of CO after reaction with  $O_2$  for t seconds (during stage 2) was divided by the original increase in current observed when CO was removed during stage 1. The half-life was calculated using  $t_{1/2} = -t \ln 2/\ln(x)$  where x is the fraction of H<sub>2</sub> production current surviving after t seconds. From control experiments such as those previously described, 44 it was estimated that the concentrations of O<sub>2</sub> experienced by the enzyme at -0.4 V under headgas conditions of 1% and 5% were 0.7% and 4.5%, respectively. The corresponding half-lives for H<sub>2</sub> oxidation activity under similar O<sub>2</sub> concentrations were calculated from the rate constants in Table 3. The results and comparisons are shown in Table 4.

All enzymes remained at least 30% active after 2000 s under 1% bulk O<sub>2</sub> (i.e., at least 0.7% O<sub>2</sub> at the electrode<sup>44</sup>) for *Dd*HydAB and *Cr*HydA1 and even under 5% bulk O<sub>2</sub>



**Figure 8.** Chronoamperometric experiments designed to measure the survival of  $H_2$  production activity in the presence of headspace levels of 1%  $O_2$  for DdHydAB and CrHydA1, and 5%  $O_2$  for CaHydA (note change in current scale upon introduction of 5%  $O_2$ ). All procedures were carried out using 1 bar pressure of the gases indicated. Other conditions: pH 6.0, 10 °C, electrode rotation 3000 rpm, -0.4 V vs SHE. The blue vertical lines show introductions of CO to inhibit and reveal enzyme-catalyzed  $H_2$  production. The fraction of enzyme surviving  $O_2$  after time t is given by the ratio of CO-sensitive current measured in Stage 2 relative to that measured after the recovery in Stage 1 (indicated by vertical double-headed arrows in each case).

Table 4. Comparative Half-Lives (s) for O<sub>2</sub>-Inactivation of H<sub>2</sub> Production and H<sub>2</sub> Oxidation of *Dd*HydAB, *Cr*HydA1 and *Ca*HydA<sup>a</sup>

catalytic direction	<i>Dd</i> HydAB	CrHydA1	<i>Ca</i> HydA
H <sub>2</sub> oxidation	30	305	1890
H <sub>2</sub> production	$\sim 2000$	$\sim 1320$	$\sim$ 2250
Catalytic direction discrimination	60 - 70	4-5	approximately 1

<sup>a</sup> The half-lives for the H₂ production reaction are estimated on the basis of the percentage of activity remaining after an extended period (1800–2000 s, different for each enzyme) of catalytic turnover in the presence of O₂. Experimental conditions for H₂ production: DdHydAB, CrHydA1; 1% O₂ in the headgas (i.e. at least 0.7% O₂ surviving at the electrode); CaHydA, 5% O₂ in the headgas (at least 4.5% surviving at the electrode) −0.4 V, pH 6, 10 °C, electrode rotation rate 3000 rpm. Data for H₂ oxidation were calculated using the second-order rate constants (see Table 3) and values of 0.7% O₂ for DdHydAB and CrHydA1; and 4.5% O₂ for CaHydA.

(around 4.5%  $O_2$  at the electrode) for CaHydA. More significantly, there were important differences among the three enzymes when comparing their survival to  $O_2$  exposure

during  $H_2$  production with the data obtained for  $H_2$  oxidation. In the case of DdHydAB, the  $H_2$ -production activity remaining is about sixty-fold greater than expected on the basis of the results described above for  $O_2$  inactivation of  $H_2$  oxidation. The enhancement is also observed for CrHydA1, but to a lesser extent than DdHydAB. On the other hand, no clear difference was observed for CaHydA between the rates of inactivation observed when monitoring  $H_2$  production at -0.4 V or  $H_2$  oxidation at -0.05 V.

## **Discussion**

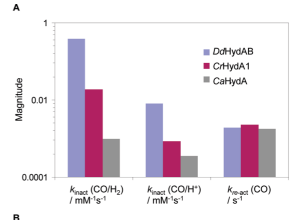
The three hydrogenases we have investigated include two with potential applications in large-scale H<sub>2</sub> production (*Cr*HydA1 for photosynthesis and *Ca*HydA for fermentation) and one of known crystal structure (*Dd*HydAB). He addition, *Ca*HydA is closely related to *Cp*I for which the structure is known. Some important comparisons have been made, exploiting the unique ability of protein film electrochemistry to measure, simultaneously, the rates and extent of changes in catalytic activities under well-defined potentials (driving force). A summary of the quantitative observations and interrelationships is provided in Figure 9.

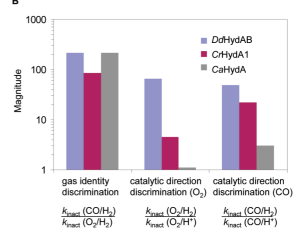
The reactions with CO are highly informative, and we<sup>34</sup> and others<sup>39</sup> have noted that CO protects [FeFe]-hydrogenases against  $O_2$  degradation, suggesting both inhibitors target the same site. In all cases we could use CO as a strong inhibitor of both  $H_2$  oxidation and  $H_2$  production, helped by the fact that binding of  $H_2$  under both conditions is much weaker than CO binding<sup>49</sup> (see Table 1).

Light sensitivity of CO inhibition is a well-established property of [FeFe]-hydrogenases 13,30,33,47,50-53 and originates from the photolability of the Fe-CO bond. A particularly useful result is the similarity in the rates of *dark* reactivation of the CO-inhibited hydrogenases. In all cases, the rate is accelerated by illumination (as reported in an earlier study for *Dd*HydAB<sup>47</sup>), and this suggests strongly that the reaction being observed in all cases is an elementary dissociation of the Fe-CO bond. This result demonstrates an intrinsic property of the H-cluster, maintained regardless of the slightly differing protein environments among the [FeFe]-hydrogenases. Evidently, all that is required for reactivation is to liberate the coordination site and ensure that CO escapes from the pocket before it can recombine. On the coordination of the pocket before it can recombine.

Our  $K_1^{CO}$  data for CaHydA determined during  $H_2$  oxidation lie broadly in the range of values (around 1  $\mu$ M) obtained by Thauer and co-workers<sup>50</sup> for CO binding to the related enzyme from C. pasteurianum, although those experiments also used a higher temperature and we found a consistently higher value

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**Figure 9.** Bar charts representing various comparisons between DdHydAB, CrHydA1, and CaHydA. (A) Comparative rates of CO-inhibition of  $H_2$  oxidation ( $k_{inact}(CO/H_2)$ ,  $H^+$  reduction ( $k_{inact}(CO/H^+)$ ) and rates of recovery from CO-inhibition ( $k_{re-act}(CO)$ ). (B) Discrimination factors characterizing the favorability of binding CO over  $O_2$  ( $k_{inact}(CO/H_2)/k_{inact}(O_2/H_2)$ ), binding  $O_2$  when the enzyme is catalyzing  $H_2$  oxidation compared to  $H_2$  production ( $k_{inact}(O_2/H_2)/k_{inact}(O_2/H^+)$ ) and binding CO when the enzyme is catalyzing  $H_2$  oxidation compared to  $H_2$  production ( $k_{inact}(CO/H_2)/k_{inact}(CO/H^+)$ ) for DdHydAB, CrHydA1 and CaHydA. As the rate of reactivation from CO inhibition is essentially independent of the process being catalyzed, it is simply represented by the term  $k_{re-act}(CO)$ . The  $k_{inact}(O_2/H_2)/k_{inact}(O_2/H^+)$  ratios are approximate because the values of  $k_{inact}(O_2/H^+)$  are estimates. This ratio is approximated to 1 for CaHydA.

(weaker binding) for the kinetic compared to the equilibrium value. The reasons for this are unclear at present, and doubtless the model we now discuss is oversimplified. The rate data for all three enzymes, both qualitative (Figure 3) and quantitative (Figure 4 and Figure 5), reveal a preference for CO binding to the enzymes under H<sub>2</sub> oxidation—the direction enforced by a higher electrode potential that should ensure that H<sub>ox</sub> predominates over H<sub>red</sub> during the catalytic cycle. The "catalytic direction discrimination" for CO decreases in the order DdHydAB >  $CrHydA1 \gg CaHydA$ —the same order as found for the equivalent factor estimated for O<sub>2</sub> inactivation. The order also matches that observed for the rates of inactivation of H<sub>2</sub> oxidation by CO and O2. These interrelationships lead us to propose a model for the reversible binding of CO to [FeFe]hydrogenases which can be extended to account for reactions with  $O_2$ . The model is represented schematically in Figure 10A.

The model considers that the attack by CO (rate constant  $k_{\text{inact}}$ ) involves two stages: the first stage is transport of CO (we treat this generically as the small molecule X) from the external medium ( $X_{\text{ext}}$ ) through the enzyme (for simplicity we show this as a single tunnel without branches or sites at which X could

<sup>(48)</sup> Nicolet, Y.; Piras, C.; Legrand, P.; Hatchikian, C. E.; Fontecilla-Camps, J. C. Struct. Folding Des. 1999, 7, 13–23.

<sup>(49)</sup> This allowed for the same concentration of H<sub>2</sub> to be employed in the experiments at -0.4 and -0.05 V, thus ensuring that the experiments were comparable. For other hydrogenases, strong H<sub>2</sub> inhibition of H<sub>2</sub> production at -0.4 V would have prevented experiments performed at this potential from being performed under 80% H<sub>2</sub> as they were at -0.05 V.

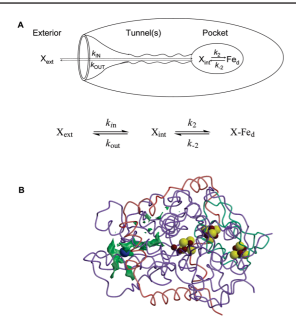


Figure 10. (A) Cartoon depicting the stepwise reaction of an inhibitory gas molecule X (= CO) with the buried H-cluster (Fe<sub>d</sub>) of a [FeFe]-hydrogenase, showing transfer from the external medium ( $X_{ext}$ ) to a position ( $X_{int}$ ) close to Fe<sub>d</sub> ( $k_{in}$ ,  $k_{out}$ ), and inner-sphere coordination/dissociation at Fe<sub>d</sub> ( $k_2$ ,  $k_2$ ). (B) C-α tracing of the [FeFe]-hydrogenase from D. desulfuricans: DdHydAB(large subunit: blue and green, small subunit: red) and the cavity observed in the crystals (probe size: 0.8 Å, program CAVsel, A. Volbeda, unpublished). The H-cluster occupies the center of the molecule. The blue sphere represents the experimentally observed Xe site. Other color codes: red: Fe, yellow: S.

be trapped) to a noncoordinating site close to the H-cluster; the second stage is migration of  $X_{int}$  from the noncoordinating site to a coordination site that we assume to be  $Fe_d$  (the actual innersphere binding reaction). The process of reactivation (overall rate constant  $k_{re-act}$ ) is the reverse of this reaction scheme.

The terms  $k_{\rm in}$  and  $k_{\rm out}$  are rates of transport of X through the protein in either direction, and  $k_2$  and  $k_{-2}$  are the elementary rates of binding and dissociation of ligand X, respectively, to Fe<sub>d</sub> within the region of the active-site pocket. With the simplification that  $k_{-2}[\text{X-Fe}_{\rm d}]$  is negligible until the reaction of inhibitor with enzyme is essentially complete, the steady-state approximation with  $d[X_{\rm int}]/dt = 0$  yields:

rate of inactivation = 
$$\frac{k_{in}k_2[X_{ext}][Fe_d]}{k_{out} + k_2}$$
 (3a)

where the pseudo first-order rate constant (as measured directly in experiments) is:

$$k'_{\text{inact}} = \frac{k_{\text{in}}k_2[X_{\text{ext}}]}{k_{\text{out}} + k_2}$$
 (3b)

An assumption of this model is that  $k_{\rm in}$  and  $k_{\rm out}$  should depend on the nature of the gas molecule and gas filter but not on the catalytic state of the H-cluster—the state predominating for a particular electrode potential, i.e.  $H_{\rm ox}$  or  $H_{\rm red}$ . Based on the evidence that CO binds preferentially (and perhaps exclusively) to  $H_{\rm ox}$ , <sup>29</sup> we expect that  $k_2$  for CO will be large for  $H_{\rm ox}$  and small, even zero, for  $H_{\rm red}$ . Note however that inhibition is still observed during  $H_2$  production because  $H_{\rm ox}$  may always appear briefly in the catalytic cycle, even at the lowest potentials we have used (-600 mV in the cyclic voltammetry experiments).

Recognizing this is a simplistic model, we now consider the following limiting scenarios: (i) if  $k_{\text{out}} \leq k_2$ ,  $k'_{\text{inact}} \sim k_{\text{in}}[X_{\text{ext}}]$  so the rate of inhibitor binding depends only on the external concentration and rate of internal transport of X; in this case little discrimination is expected based on the redox state of the H-cluster. Alternatively, (ii), if  $k_{\text{out}} \gg k_2$ , i.e. if the protein's internal structure does not provide an effective barrier to transport of X, it follows that  $k'_{\text{inact}} = k_2 k_{\text{in}} [X_{\text{ext}}]/k_{\text{out}}$ . In this scenario, the rate of inhibition depends not only on the nature of X but also upon  $k_2$  and therefore should also be faster for conditions favoring  $H_{\text{ox}}$  ( $H_2$  oxidation) compared to  $H_{\text{red}}$  ( $H_2$  production).

Reactivation follows the reverse sequence, and for  $k_{in}[X_{ext}]$  = 0 (because CO is removed from the solution) we obtain

rate of re-activation = 
$$\frac{k_{\text{out}}k_{-2}[X-Fe_{\text{d}}]}{k_{\text{out}}+k_2}$$
 (4a)

and the first-order rate constant (as measured experimentally) is given by

$$k_{\text{re-act}} = \frac{k_{\text{out}}k_{-2}}{k_{\text{out}} + k_2} \tag{4b}$$

Under the limiting condition  $k_{\rm out} \ll k_2, k_{\rm re-act} = k_{\rm out}k_{-2}/k_2$ , whereas if  $k_{\rm out} \gg k_2$ , the rate of reactivation reduces to  $k_{-2}$ , reflecting the likelihood that CO escapes from the enzyme  $(k_{\rm out})$  before it can recoordinate  $(k_2)$ . Our data suggest that the latter situation must generally be the case, with a more intermediate situation (a smaller  $k_{\rm out}$  relative to  $k_2$ ) applying for CaHydA (see below). Overall, the dissociation constant is given by  $K_1^{\rm CO}({\rm kin}) = k_{\rm re-act}/k_{\rm inact} = k_{\rm out}k_{-2}/k_{\rm in}k_2$ , which always depends on the kinetics of making and breaking the Fe-CO bond.

This analysis can be extended to the reaction of [FeFe]-hydrogenases with O<sub>2</sub>, although that reaction is essentially irreversible. Table 3 and Figure 9 show that trends among the hydrogenases as observed for their reactions with CO are mirrored in their reactions with O<sub>2</sub>; for example, *Dd*HydAB shows the highest rates of inhibition in both cases and the greatest discrimination based on catalytic direction.

In mechanistic terms, the minuscule protection that DdHydAB possesses against attack by  $O_2$  is provided only within the active-site pocket in which  $O_2$  is able to discriminate between different catalytic states of the enzyme (a strong  $k_2$  dependence, according to the model). In contrast, the small catalytic direction discrimination observed for CaHydA can be interpreted in terms of it showing a less excessive value of  $k_{\text{out}}$  (a more restrictive tunnel or filter) in relation to  $k_2$ . Values for  $k_{\text{out}}$  should correlate closely with those for  $k_{\text{in}}$ ;thus, it is significant that CaHydA also shows the slowest rates of reaction with CO and  $O_2$  and, with a half-life of several minutes under atmospheric  $O_2$  levels at 10 °C, looks to be a promising model for aerobic biohydrogen production even though it stems from a strict anaerobe.

The evidence (strong light enhancement) that the rate-determining step in reactivation is the elementary scission of the Fe-CO bond in the H<sub>ox</sub>-CO state, and the observation that the rate measured in the dark is quite similar for all enzymes (which share only 40% sequence similarity) shows that the kinetics of reactivation are governed more by the intrinsic properties of the H-cluster than by the nature of the surrounding enzyme. Lubitz and colleagues have proposed that the H-clusters in *Cr*HydA1 and *Dd*HydAB are similar, although they differ slightly in electronic detail.<sup>28</sup> Note that were CO to coordinate

to different states of the H-cluster at -0.4 and -0.05 V, we would expect  $k_{\text{re-act}}$  to depend significantly on potential, but it does not. This supports the view that CO (and by extension,  $O_2$ ) binds to  $H_{ox}$  but not  $H_{red}$ . The other comparison in Figure 9 which is reasonably constant among all three enzymes is the gas identity discrimination (CO vs O<sub>2</sub>). This again may reflect intrinsic behavior of the H-cluster because CO is a superior ligand to  $O_2$  in terms of its  $\pi$ -acceptor capability. Dominant intrinsic effects are not unexpected, given the unusually independent status of the 2Fe<sub>H</sub> subcluster, which was described in the Introduction as an enzyme cofactor resembling an organometallic compound buried in a protein. Clearly our model is an oversimplification, albeit necessary at this stage, and to understand this observation more fully we are undertaking theoretical calculations, including predictions of relative transport rates through the enzyme.

The notion of a filter or a tunnel connecting the H-cluster to the molecular surface is supported by the two available [FeFe]hydrogenase structures. In *Dd*HydAB and *CpI* there is a 'static' tunnel that can be revealed using a cavity-searching program.<sup>55</sup> The tunnel in each enzyme has a central cavity that can bind a Xe atom and a narrower path leading to Fe<sub>d</sub>. <sup>17,26</sup> Figure 10B depicts the experimentally observed tunnel and Xe site in DdHydAB. 56,57 The tunnel connects the molecular surface to the active site, and one possibility is that the Xe atom occupies a cavity in which dissociated CO could reside before rebinding to Fe<sub>d</sub> or escaping to the medium. Molecular dynamics simulations based upon the CpI structure revealed a second tunnel that also connects to the central cavity.<sup>58</sup> What can be concluded from both crystallographic and theoretical studies is that dynamic fluctuations are important for intramolecular gas diffusion in [FeFe]-hydrogenases.

Two further mechanistic points emerge from this study. First, we always observed that CO inhibition of H<sub>2</sub> production is only partially reversible. At present we have no explanation for this, although Adams reported in 1987 that CO binds irreversibly to a catalytic intermediate of CpI.<sup>59</sup> Further investigations including a full study of the potential dependence of CO binding during H<sub>2</sub> production are clearly required to resolve this issue, which may have mechanistic relevance. Second, we always recorded a small proportion of activity returning after O<sub>2</sub> inactivation, an observation in line with those reported by Baffert et al.<sup>39</sup> also in studies on CaHydA. Such a part-reversal is consistent with the mechanism proposed by Stripp et al.<sup>34</sup> in which O<sub>2</sub> must first bind in a reversible manner at the distal Fe of the 2Fe<sub>H</sub> subcluster before causing irreversible damage to the [4Fe-4S]<sub>H</sub> subcluster.

From a biological perspective, the precise, quantitative data that we have been able to extract and compare for the three different hydrogenases should be understandable in terms of the lifestyles of the organisms that express them. This is true, in part. Recent studies on the O<sub>2</sub> detoxification mechanism in C. acetobutylicum have shown that this fermentative bacterium can survive limited exposure to air and can even undergo cell division at surprisingly high concentrations of O<sub>2</sub>.60,61 The relative O2 stability of CaHydA may therefore be a consequence of concerted evolutionary adaptation to an O<sub>2</sub>-rich atmosphere. However, certain species of the Desulfovibrio genus have also been reported to exhibit short-term survival when exposed to O<sub>2</sub>;<sup>62,63</sup> thus, from a microbiological viewpoint the large disparity in O2 sensitivity between CaHydA and DdHydAB is puzzling. The 'intermediate' degree of O<sub>2</sub> sensitivity displayed by CrHydA1 is consistent with the observation that although it is only expressed in *C. reinhardtii* under anaerobiosis<sup>64,65</sup> it is likely to be in contact with at least trace amounts of O<sub>2</sub> that are produced by photosystem II.66

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Supporting Information Available: Photolability of the CObound state; determination of the variation of the concentration of CO with time in the experiments shown in Figure 4; determination of  $K_{\rm I}^{\rm CO}({\rm equil})$ ; control experiments for experiments investigating O<sub>2</sub> inactivation of H<sub>2</sub> oxidation. This material is available free of charge via the Internet at http:// pubs.acs.org.

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