# SCIENTIFIC REPORTS

### OPEN

Received: 21 April 2015 Accepted: 29 July 2015 Published: 14 September 2015

## Lyophilization protects [FeFe]hydrogenases against O<sub>2</sub>-induced H-cluster degradation

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Nature has developed an impressive repertoire of metal-based enzymes that perform complex chemical reactions under moderate conditions. Catalysts that produce molecular hydrogen (H<sub>2</sub>) are particularly promising for renewable energy applications. Unfortunately, natural and chemical H<sub>2</sub>- catalysts are often irreversibly degraded by molecular oxygen (O<sub>2</sub>). Here we present a straightforward procedure based on freeze-drying (lyophilization), that turns [FeFe]-hydrogenases, which are excellent H<sub>2</sub>-producers, but typically extremely O<sub>2</sub>-sensitive in solution, into enzymes that are fully resistant against O<sub>2</sub>. Complete dryness protects and conserves both, the [FeFe]-hydrogenase proteins and their inorganic active-site cofactor (H-cluster), when exposed to 100% O<sub>2</sub> for days. The full H<sub>2</sub>-formation capacity is restored after solvation of the lyophilized enzymes. However, even minimal moisturizing re-establishes O<sub>2</sub>-sensitivity. The dry [FeFe]-hydrogenase material is superior also for advanced spectroscopic investigations on the H-cluster reaction mechanism. Our method provides a convenient way for long-term storage and impacts on potential biotechnological hydrogen production applications of hydrogenase enzymes.

The global market for enzymatically catalyzed reactions and their products is continuously growing. The use of enzymes often is preferable over conventional chemical processes because of their stereo-selective chemistry and superior turnover rates under moderate temperatures and pressures. With the large-scale cultivation of bacteria and yeast as homologous and heterologous expression hosts, valuable products can be generated in large and therefore economically attractive quantities. Besides the industrial production of metabolites like vitamins, amino acids, vaccines or antibiotics a number of highly active and specific enzymes are produced and purified for the *in vitro* generation of further high-value products and applied in food-, pharmaceutical-, and detergent-industry<sup>1</sup>. Also the production of second-generation biofuels by enzymatic polymer-degradation using cellulases and of first-generation biofuels like ethanol and methane from biomass fermentation are commercially well established procedures<sup>2</sup>.

A promising fuel resource that can also be produced biologically is molecular hydrogen (H<sub>2</sub>). Two H<sub>2</sub>-producing biocatalysts are found in nature, which inspire modern biomimetic catalyst research. Nitrogenases turn molecular nitrogen into ammonia, a process which yields H<sub>2</sub> only as a byproduct<sup>3</sup>. On the other hand, hydrogenase enzymes directly catalyze the reduction of protons to form H<sub>2</sub> as the primary product<sup>4,5</sup>. With reported turnover rates of up to 10,000 molecules of H<sub>2</sub> per second, [FeFe]-hydrogenases are the most efficient known biocatalysts for H<sub>2</sub> generation and therefore promising targets for potential applications<sup>6–8</sup>.

The catalytically active cofactor (H-cluster) of [FeFe]-hydrogenases is a [6Fe6S] complex (further on denoted [6FeH]), consisting of a standard cubane [4Fe4S] cluster ([4FeH]) and a unique diiron complex

<sup>1</sup>Ruhr-Universität Bochum, Fakultät für Biologie und Biotechnologie, Lehrstuhl für Biochemie der Pflanzen, AG Photobiotechnologie, 44801 Bochum, Germany. <sup>2</sup>Freie Universität Berlin, Institut für Experimentalphysik, 14195 Berlin, Germany. <sup>3</sup>Ruhr-Universität Bochum, Fakultät für Chemie und Biochemie, Anorganische Chemie I–Bioanorganische Chemie, 44801 Bochum, Germany. Correspondence and requests for materials should be addressed to T.H. (email: thomas.happe@rub.de) ([2FeH]) which represents the actual active site of reversible  $H_2$ -production<sup>8-10</sup>. The two iron atoms in [2FeH], in distal (Fe<sub>d</sub>) or proximal (Fe<sub>p</sub>) positions to [4FeH], bind unusual carbon monoxide (CO) and cyanide (CN<sup>-</sup>) ligands, which tune the electrochemical and redox properties of the H-cluster and guide state transitions during the catalytic cycle<sup>11,12</sup>. An azadithiolate (adt =  $S_2(CH_2)_2NH$ ) ligand connects the two iron ions in [2FeH] and its secondary amine bridgehead shuttles protons between a conserved H<sup>+</sup>-transfer pathway in the protein and the site of catalytic turnover<sup>13-15</sup>. The sub-clusters in [6FeH] are covalently and electrochemically coupled through one of the four cysteine ligands that bind the [4FeH] complex to the protein scaffold<sup>16</sup>.

A very high hydrogenase activity and thus  $H_2$ -production yield can be achieved *in vitro* using purified [FeFe]-hydrogenase protein when the natural electron mediators are substituted by redox-active chemicals or the enzyme is immobilized on an electrode<sup>7,17,18</sup>. Under such conditions, the rate-limiting metabolic pathways of electron supply as present in the living cell are omitted and competitive redox processes are absent<sup>6,18,19</sup>. The major obstacle for introducing these promising  $H_2$ -producing biocatalysts into biotechnological applications, however, is their sensitivity towards molecular oxygen (O<sub>2</sub>). While all hydrogenases are inhibited by O<sub>2</sub> to a certain extent, [FeFe]-hydrogenases exhibit the highest level of O<sub>2</sub>-sensitivity due to irreversible degradation of the H-cluster within seconds to minutes during exposure even to traces of dioxygen<sup>20-22</sup>.

In this study, we present a method which prevents the O<sub>2</sub>-induced inactivation of two [FeFe]-hydrogenases representing different prototypic enzyme structures. HYDA1 from the green alga *Chlamydomonas reinhardtii* represents the minimal M1-type, binding only the H-cluster, whereas CPI from the bacterium *Clostridium pasteurianum* represents the most complex monomeric M3-type, containing in addition to the H-cluster three [4Fe4S] and one [2Fe2S] cluster<sup>23</sup>. We developed and applied a technique for protein lyophilization (freeze-drying by water sublimation at low pressure and cryogenic temperatures<sup>24</sup>), which removes bulk and protein-bound water from protein preparations, respectively, for the preservation of [FeFe]-hydrogenases. Our biochemical analyses show that these extremely O<sub>2</sub>-sensitive enzymes after lyophilization are completely protected against O<sub>2</sub>. Structural characterization using X-ray absorption spectroscopy (XAS)<sup>25-28</sup> on the lyophilized proteins revealed the integrity of the H-cluster. The procedure of freeze-drying thereby is fully reversible, yet also restoring O<sub>2</sub>-sensitivity in solution. We suggest that the H-cluster degradation requires both, O<sub>2</sub> attack and either specific protein bound H<sub>2</sub>O molecules or at least a water-dissolved enzyme state. These findings are an important milestone for biotechnological use of [FeFe]-hydrogenases as sustainable catalysts for H<sub>2</sub> fuel generation and provide a new impulse in the challenging task to overcome the enzyme's Achilles heel.

#### Results

Improved purification yields of HYDA1 [FeFe]-hydrogenase. Significant improvements in purification yield and specific activity have been achieved for [FeFe]-hydrogenase HYDA1 from C. reinhardtii since the first isolation about 30 years ago (Supplementary Tab. S1)<sup>29</sup>. Here, besides HYDA1 a second [FeFe]-hydrogenase type, CPI from C. pasteurianum, was produced by overexpression in Escherichia coli, which contained the active-site cofactor in two different maturation states. The inactive proteins denoted [4FeH] bind only the [4Fe4S] subsite of the H-cluster that is assembled by the iron-sulfur cluster housekeeping (ISC) machinery in E. coli, whereas the fully active holoenzyme denoted [6FeH] is equipped with both, the four-iron and the diiron ([2FeH]) sub-complexes of the H-cluster. In the living host, synthesis and assembly of the [2FeH]-subsite requires specific maturases, namely HYDE, HYDF and HYDG<sup>30,31</sup>. By replacing the previously used Strep-tag II (STII) with a hexahistidine-tag (HIS<sub>6</sub>) fused to the HYDA1 protein, we improved the yield of HYDA1[6FeH] expressed in E. coli containing the HYDE,F,G background by a factor of 1.5 and the yield of HYDA1[4FeH] expressed in E. coli without HYDE,F,G by a factor of 3 (Supplementary Fig. S1; Supplementary Tab. S1) in comparison to previously reported purification procedures<sup>32</sup>. An efficient method for *in vitro* H-cluster maturation has been recently reported, in which the [4FeH] protein is activated with a chemically synthesized analogue (here denoted 2Fe<sub>adt</sub>) of the diiron site, thereby forming the [6FeH] enzyme with full activity<sup>14</sup>. The combination of the HIS<sub>6</sub>tagged HYDA1[4FeH] construct and in vitro maturation with 2Fe<sub>adt</sub> thus facilitated the production of up to 90 mg of fully active HYDA1[6FeH] with a typical specific activity of about 950  $\mu$ mol H<sub>2</sub>  $\cdot$  mg<sup>-1</sup>  $\cdot$ min<sup>-1</sup> per liter of *E. coli* culture (Supplementary Tab. S1)<sup>14</sup>. This amount of enzyme is capable to produce about 2 liters of H<sub>2</sub> per minute under maximum turnover conditions.

**Lyophilization of two different [FeFe]-hydrogenases.** The process of lyophilization, also called freeze-drying, is commonly used to preserve valuable and labile molecules for application<sup>33,34</sup>. HYDA1 and CPI proteins in the [4FeH] and [6FeH] maturation states were first deep-frozen in liquid nitrogen and then dehydrated under vacuum through water sublimation, resulting in lyophilized samples. Thereafter, the H<sub>2</sub>-evolution activity was assayed after re-solvation of aliquots of the dry proteins in buffer solutions. The lyophilized [6FeH]-containing holoenzymes, *in vitro* maturated with the  $2Fe_{adt}$  complex prior to lyophilization, were assayed directly after rehydration. Freeze-dried hydrogenases, which only contained the [4FeH] were *in vitro* activated during rehydration with a 10-fold molar excess of  $2Fe_{adt}$  to yield active [6FeH]-containing enzymes after lyophilization. In addition, the enzyme activities of non-lyophilized (control) [6FeH]- and  $2Fe_{adt}$ -activated [4FeH]-samples and lyophilized protein samples were compared (Fig. 1).



**Figure 1.** Specific H<sub>2</sub>-evolution activities of HYDA1 and CPI before and after lyophilization. Specific H<sub>2</sub>-production rates of the [FeFe]-hydrogenases HYDA1 (green) and CPI (brown) were determined using the sodium-dithionite-driven methyl-viologen-reduction assay (1 $\mu$ g HYDA1 or 0.5 $\mu$ g CPI). The enzymes were both assayed before or after lyophilization and subsequent re-suspension; in addition, [4FeH]-proteins were maturated with 2Fe<sub>adt</sub> prior to activity determination. Mean activities from at least 3 independent measurements are shown, error bars give the standard deviations.



Figure 2.  $O_2$ -sensitivity of lyophilized and dissolved [FeFe]-hydrogenases. Sample annotations are as in Fig. 1 and H<sub>2</sub>-activities were determined accordingly. Hydrogenase proteins were (**a**) exposed after lyophilization to 100%  $O_2$  for up to 24 hours (green symbols, HYDA1; brown symbols, CPI) or to ambient air for 24 h (triangles) at room temperature or (**b**) solutions of non-lyophilized proteins (6.33 nM) were incubated with the indicated  $O_2$ -levels for 15 min at 37 °C. Values are the mean of at least 3 independent measurements, error bars give standard deviations.

The HYDA1 and CPI preparations showed a 1:3 ratio of their specific activities, which is in agreement with previous results<sup>14</sup>. No significant differences were found between the  $H_2$ -evolution rates of previously purified  $2Fe_{adt}$ -activated [6FeH]-enzymes and [4FeH]-enzymes that were *in vitro* activated immediately prior to the assay. The specific  $H_2$ -activities after lyophilization were practically identical to the activities determined for the non-lyophilized control samples for both HYDA1 and CPI, demonstrating the complete reversibility and activity recovery after rehydration of the lyophilized [FeFe]-hydrogenases. These results reveal that both, chlorophycean M1-type HYDA1 and bacterial M3-type CPI [FeFe]-hydrogenases can be lyophilized without any significant loss of catalytic activity (Fig. 1). Accordingly, the feasibility and reversibility of lyophilization is independent from the level of structural complexity of the [FeFe]-hydrogenase. Moreover, there was no need for additives like lyo- or cryo-protectants to shelter the proteins, because full activity recovery was achieved for enzymes lyophilized in a common storage buffer (Tris-HCl, pH 8).

 $O_2$ -tolerance of lyophilized [FeFe]-hydrogenases. The  $O_2$ -sensitivity levels of control and lyophilized HYDA1 and CPI samples were compared. The lyophilized [4FeH] and [6FeH] proteins were exposed to dry pure  $O_2$  gas for up to one day and H<sub>2</sub>-evolution activities were determined after subsequent enzyme dissolution (and maturation with 2Fe<sub>adt</sub> of the [4FeH] proteins) (Fig. 2a). In addition,





lyophilized proteins were exposed to ambient air for 24h. Solution samples of the non-lyophilized hydrogenases were incubated with increasing  $O_2$ -concentrations for only 15 min for comparison (Fig. 2b).

The results show that the lyophilized hydrogenases retained their full catalytic competence even after 24 h exposure to 100%  $O_2$ . This holds no matter whether [4FeH] or [6FeH] proteins were exposed to  $O_2$ . The enzymes in solution, however, almost entirely lost H<sub>2</sub>-production activity already after 15 min under 10%  $O_2$  (CPI) or 2.5%  $O_2$  (HYDA1). The degrees of inactivation of the solution samples were similar for [6FeH] and [4FeH] proteins, meaning that not only the complete H-cluster, but also the [4FeH] cluster alone apparently is damaged by  $O_2$ . Interestingly, all lyophilized hydrogenase proteins were almost completely inactivated after incubation in ambient air for 24h (Fig. 2a). A plausible explanation for this behavior was that the humidity (water vapor) in the air caused partial rehydration of the proteins<sup>35,36</sup>, which restored the  $O_2$ -sensitivity, as examined further below.

**Restoration of O<sub>2</sub>-sensitivity by humidity.** We analyzed the impact of atmospheric humidity on the O<sub>2</sub>-induced inactivation rate of the lyophilized [FeFe]-hydrogenases. Increasing amounts of H<sub>2</sub>O-saturated O<sub>2</sub> gas were injected into sealed reaction vials containing lyophilized hydrogenase in an atmosphere of O<sub>2</sub> and the samples were incubated at room temperature for 30 min prior to H<sub>2</sub>-evolution activity determination (Fig. 3). The data show that the catalytic activity of all lyophilized proteins in the presence of O<sub>2</sub> was decreasing for increasing humidity levels. For CPI no difference between the activities of [4FeH] and [6FeH] proteins was observed. For HYDA1, however, the O<sub>2</sub>-sensitivity seems to be slightly higher for increasing humidity in the [4FeH] protein compared to the [6FeH] protein. This finding corroborated the observation described above that already the protein containing only the [4FeH] cluster is a target for O<sub>2</sub> attack.

H-cluster integrity studied by X-ray absorption spectroscopy. XAS was employed to characterize the oxidation state and molecular structure (iron-ligand bond lengths and Fe-Fe distances) of the H-cluster sub-complexes in the lyophilized HYDA1 proteins (Fig. 4). The redox level and metric parameters are indicative of the H-cluster integrity<sup>24</sup>. We compared the following preparations: (1) HYDA1[4FeH] protein overexpressed in *E. coli* in the absence of the three maturases HYDE,F,G was prepared anaerobically and studied in the anoxic state ([4FeH]  $-O_2$ ) and after exposure to dry pure  $O_2$ gas for ~15 min ([4FeH]  $+O_2$ ). (2) HYDA1[6FeH] protein, which contains the complete H-cluster was studied in the anoxic state ([6FeH]  $-O_2$ ), after exposure to dry pure  $O_2$  gas for ~15 min ([6FeH]  $+O_2$ ), and after exposure to air at ambient humidity for ~5 days ([6FeH]  $+O_2 + H_2O$ ). XAS experiments on lyophilized HYDA1 proved to be highly advantageous, because the X-ray fluorescence signal intensity was increased about 10-fold compared to solution samples with similar protein contents and the scattering background was drastically diminished due to the absence of water, which largely improved the XAS data quality at shortened measuring times.

The XANES and EXAFS spectra of lyophilized HYDA1[4FeH] even after extended  $O_2$  exposure were very similar to the spectra of HYDA1[4FeH] in anoxic solution<sup>28</sup>, indicating a similar iron coordination environment and thus an intact [4Fe4S] subsite of the H-cluster under both conditions (Fig. 4a). The EXAFS analysis revealed the typical four Fe-S bonds (~2.3 Å) and three Fe-Fe distances (~2.7 Å) per Fe ion in the cubane cluster, both in the absence and after exposure to  $O_2$  (Tab. 1). The structure of the [4Fe4S] cluster thus was not significantly affected by lyophilization and  $O_2$ -exposure. The XAS spectra of lyophilized anoxic HYDA1[6FeH] differed pronouncedly from the spectra of HYDA1[4FeH] (Fig. 4b), indicating quantitative binding of the [2FeH] unit to the cubane cluster in the 2Fe<sub>adt</sub>-maturated HYDA1[6FeH]<sup>13,14</sup>. The K-edge energy of the lyophilized HYDA1[6FeH] was similar within ±0.15 eV to



**Figure 4. Fe-XAS analysis of lyophilized HYDA1 proteins. (a)** XANES spectra of the indicated [FeFe]hydrogenase samples. (b) Fourier-transforms (FTs of experimental data) of EXAFS spectra in the inset (black lines, experimental data; colored lines, simulations with parameters in Table 1). The color code is the same and spectra were vertically displaced for comparison in panels (a and b).

that of heterologously expressed HYDA1[6FeH] from *Clostridium acetobutylicum* in solution, suggesting that not more than one Fe ion in the lyophilized protein differed by one oxidation step from the reduced state of the H-cluster  $(H_{red})^{27,28}$ . The EXAFS was well simulated using similar iron-ligand bond lengths and Fe-Fe distances as for enzyme from *C. acetobutylicum* in solution<sup>10,20,21</sup>, showing that a complete and intact H-cluster was present in the lyophilized HYDA1[6FeH] (Tab. 1).

After ~15 min O<sub>2</sub>-exposure of lyophilized HYDA1[6FeH], the XANES revealed only small changes and the EXAFS still was compatible with an intact H-cluster (Tab. 1). However, slight shortening of the Fe-Fe distances attributable to the [4Fe4S] cluster and shortening of the Fe-C(=O/N) bonds and elongation of the Fe-Fe distance in [2FeH], as well as an increased K-edge energy and amplitude compared to the anoxic protein may be explained by one-electron oxidation of the H-cluster and possible binding of an oxygen species to [2FeH]<sup>16,21</sup>. Accordingly, O<sub>2</sub>-induced H-cluster degradation as observed in solution was prevented in the lyophilized HYDA1[6FeH] sample<sup>21</sup>. After long-term exposure to air at ambient humidity, however, the drastically altered XANES reflected symmetrization of the iron coordination in octahedral sites and the EXAFS indicated the loss of most Fe-Fe distances and extensive replacement of iron-sulfur by iron-oxygen bonds (Tab. 1)<sup>20,21</sup>. O<sub>2</sub>-induced H-cluster degradation thus only occurs upon rehydration of the lyophilized protein.

#### Discussion

Increasing efforts have been made to stabilize  $O_2$ -sensitive enzymes for utilization as sustainable catalysts in biotechnological applications. Several highly promising members of the enzyme class of oxidoreductases are especially prone to rapid inactivation under aerobiosis including nitrogenases and hydrogenases<sup>20,21,37</sup>. Oxidoreductases carry redox-active cofactors that facilitate essential electron and proton transfer steps during substrate turnover. In particular, iron-sulfur clusters are ubiquitously employed as electron transfer relays in redox enzymes and often a target of  $O_2$  attack<sup>38</sup>. Removal of harmful oxygen species from protein preparations *in vitro* may be achieved using chemical reductants or enzyme-based  $O_2$ -scavengers. However, these approaches often have certain disadvantages, like undesired pH-changes, and therefore are not well suitable for sustained biotechnological processes<sup>39</sup>. Removal of  $O_2$  from the processing atmosphere usually requires significantly larger technical efforts than removal of water (drying).

Hydrogenases are efficient catalysts for both, the production and oxidation of  $H_2$  and in general rather sensitive to  $O_2$ . Depending on the enzyme family,  $O_2$  induces either reversible or irreversible inactivation processes, both of which have been analyzed at the mechanistic and structural level<sup>4,22</sup>. In recent years a variety of promising strategies have been pursued to understand and increase the varying levels of  $O_2$ -tolerance in the hydrogenases, including research on the mechanisms of the  $O_2$  reactions at the cofactors<sup>20,21,40-42</sup>, enzyme modifications by mutagenesis<sup>43,44</sup>, and development of indirect protection systems. A recently reported *in vitro* approach uses embedding of the enzymes in a polymer-hydrogel matrix to shelter electrode-coupled hydrogenase from  $O_2$ -attack<sup>19</sup>. This system stabilized  $H_2$ -oxidation, but not  $H_2$ -evolution in an atmosphere with up to 7%  $O_2$ , which, however, was still considerable lower than atmospheric  $O_2$  partial pressure (~21%). Electrons for the reduction of detrimental oxygen species at the active site were delivered via viologen moieties in the matrix, mimicking the function of the recently discovered unusual proximal [4Fe3S] cluster in membrane-bound [NiFe]-hydrogenases, which natively show significant  $O_2$ -tolerance<sup>19,41,45-47</sup>.

		N [per Fe ion]/R $[Å]/2\sigma^2 \times 10^3 [Å^2]$					R <sub>F</sub> [%]
HYDA1:		Fe-C/O	Fe-S	Fe-Fe, [2FeH]	Fe-Fe, [4FeH]	Fe <sup></sup> O/N	
[4FeH]	-O <sub>2</sub>	-	4/2.28/7	-	3/2.73/10	-	9.0
	+O <sub>2</sub>	-	4/2.28/7	-	3/2.73/10	-	12.2
[6FeH]	-O <sub>2</sub>	1/1.91/5#	3.5/2.27/10	0.5/2.55/2#	2/2.72/7	1/3.00/5#	12.5
	+O <sub>2</sub>	1/1.87/5# 0.82*/1.86/5#	3.5/2.26/9 3.65*/2.26/10 <sup>&amp;</sup>	0.5/2.58/2# 0.5/2.57/2#	2/2.71/10 1.87*/2.71/7&	1/3.05/5# 1/3.07/5#	14.4 14.7
	$+O_2 + H_2O$	3.07*/2.04/5#	1.73*/2.29/10 <sup>&amp;</sup>	0.05*/2.55/2#	0.39 <sup>*</sup> /2.71/7 <sup>&amp;</sup>	0.34*/3.18/5#	13.8

**Table 1. EXAFS simulation parameters of lyophilized HYDA1 proteins**<sup>a</sup>. <sup>a</sup>N, coordination number; R, interatomic distance;  $2\sigma^2$ , Debye-Waller parameter; R<sub>p</sub> fit error sum calculated for reduced distances of 1–3Å<sup>25</sup>. Data correspond to EXAFS spectra in Fig. 4b. Fit restraints: N-values were fixed to the given numbers in the fits representing the expected coordination numbers for the [4Fe4S] cluster in HYDA1[4FeH] or the approximate N-values for a complete H-cluster in HYDA1[6FeH]<sup>21,28</sup>, \*except for N-values that were allowed to vary in the fits; <sup>&</sup>the same  $2\sigma^2$ -values were used for respective fits of spectra plus or minus O<sub>2</sub>; <sup>#</sup> $2\sigma^2$  values that were fixed in the simulations. For [6FeH] +O<sub>2</sub>, the fit result for unrestricted N-values (\*) emphasizes that the Fe-CO/N and Fe-S bonds and the Fe-Fe interactions in the H-cluster are fully preserved in the presence of O<sub>2</sub>.

The catalytic efficiency for proton reduction to  $H_2$  is at least ten-times larger for [FeFe]-hydrogenases compared to [NiFe]-hydrogenases<sup>48,49</sup>. An approach to generate a non-immobilized [FeFe]-hydrogenase preparation, which can be stored in a stable and catalytically intact state under aerobiosis has not been described yet. We show that lyophilization is a way to produce [FeFe]-hydrogenase material resistant to  $O_2$ . By applying a lyophilization technique under reducing conditions, an  $O_2$ -resistant protein was obtained for representatives of two major [FeFe]-hydrogenase sub-families, namely HYDA1 (M1-type) and CPI (M3-type). No special precautions such as the use of cryo- or lyo-protectants as described for other enzymes<sup>35,50</sup> were necessary for the [FeFe]-hydrogenases to facilitate recovery of full catalytic activity after rehydration. Moreover, the absence of surface-exposed thiol groups, which are a major factor in oxidative damaging of many lyophilized proteins due to formation of inter- or intra-molecular disulfide bonds<sup>51</sup>, may prevent the generation of irreversible aggregates and activity loss in the [FeFe]-hydrogenases.

Our XAS analysis clearly shows that the structure of the [4Fe4S] cluster in HYDA1[4FeH] is not affected by lyophilization nor does the cluster react with  $O_2$  in the dry protein. In contrast, HYDA1[4FeH] loses its competence for *in vitro* maturation with the synthetic [2FeH] analogue 2Fe<sub>adt</sub> to yield active holoenzyme when exposed to  $O_2$  under humid conditions. The almost identical humidity-dependences of the  $O_2$ -induced [6FeH] holoenzyme inactivation and of the loss of the maturation competence of the [4FeH] apoenzyme show that under 100%  $O_2$  even the [4Fe4S] cluster in the absence of the [2FeH] site is reacting with  $O_2$ .

For  $O_2$ -sensitive bacterial [4Fe4S]-cluster ferredoxins it has been shown that  $O_2$ -exposure may either lead to a relatively stable [3Fe4S] product that can be reactivated or causes complete cluster degradation<sup>52</sup>. The susceptibility of a cofactor to oxidative attack may correlate with its surface accessibility<sup>53,54</sup>. The vacant [2FeH] binding-site of inactive [4FeH] protein exposes a relatively large water-accessible surface so that oxygen species likely can reach and damage the [4Fe4S] cluster<sup>31</sup>. This might explain the loss of the maturation competence of the [4FeH] proteins. It remains to be shown whether  $O_2$ -exposure produces a [3Fe4S] species, which can be reactivated, or other detrimental degradation reactions at the cofactor or the protein scaffold. However, the partial surface accessibility of the [4Fe4S] cluster also in HYDA1[6FeH] may suggest that reaction steps are involved in inactivation that are independent of the [2FeH] site.

Our structural characterization revealed that the metric parameters of lyophilized HYDA1[6FeH] derived by *in vitro* maturation of HYDA1[4FeH] from *E. coli* with 2Fe<sub>adt</sub> closely resemble the parameters of HYDA1[6FeH] from *C. acetobutylicum in vivo* maturated via HYDE,F,G<sup>10,20,21,27,28</sup>. This indicated an intact structure of the H-cluster in the *in vitro* maturated holoenzyme, which was fully preserved after lyophilization. The XAS data further show that an intact H-cluster was still present after 15 min exposure of lyophilized HYDA1[6FeH] to pure  $O_2$  gas. However, the observed slight geometry changes in the cofactor are compatible with  $O_2$ -binding and single-electron oxidation of the complex, meaning that  $O_2$  may have partially reacted with the H-cluster in the lyophilized [FeFe]-hydrogenase. Binding of a single  $O_2$  molecule at Fe<sub>d</sub> has been proposed to initiate the oxidative H-cluster degradation reactions<sup>20,21</sup>. Furthermore, this process may crucially depend on specific protonation events of formed reactive oxygen species<sup>55</sup>.

The removal of bulk and protein-bound water from the [FeFe]-hydrogenase by lyophilization may be expected to cause a rupture of the proton transfer pathway leading from the protein surface to the active site. For lyophilized [NiFe]-hydrogenases it has been shown that H/D-isotope exchange activity, which depends on a functional proton transfer pathway, can only occur at dramatically lowered rates<sup>56,57</sup>.

Accordingly, the absence of protonation of reactive oxygen species formed at the H-cluster after the binding of  $O_2$  provides a plausible explanation for the  $O_2$ -resistance of the lyophilized [FeFe]-hydrogenases. Thereby, the H-cluster may be trapped after the first reaction step when  $O_2$  is still bound to Fe<sub>d</sub>. This situation would prevent binding and reaction of further  $O_2$ , which in solution leads to partial or complete H-cluster degradation<sup>21,40</sup>. This mechanism implies that the degradation process may continue as soon as proton transfer is re-established, thereby explaining the inactivation of the lyophilized [FeFe]-hydrogenases at increasing rehydration levels. That lyophilized and  $O_2$ -exposed HYDA1 was fully active in anaerobic solution suggests that the oxygen species bound to the H-cluster was safely removed by dissolution under reducing conditions of the protein, thus preventing irreversible  $O_2$ -inhibition.

In conclusion, our biochemical and spectroscopic experiments on lyophilized [FeFe]-hydrogenases provide clear evidence that water plays an important role in the  $O_2$ -induced inactivation and degradation reactions at the H-cluster. These reactions are effectively prevented in the lyophilized proteins, preserving full activity even after long-term exposure to pure  $O_2$ . Lyophilized and thereby highly up-concentrated and water-free [FeFe]-hydrogenase protein samples are particularly suitable for advanced spectroscopic analyses resulting in superior signal quality. The catalytic competence and structure of the active-site H-cluster in the enzymes is unaffected by lyophilization and rehydration. Lyophilization of [FeFe]-hydrogenases thus is a straightforward method to conserve the versatile chemical properties of these fascinating H<sub>2</sub>-catalysts.

#### Methods

**Organisms and growth conditions.** Escherichia coli strain DH5 $\alpha$  MCR was used for the cloning procedures of the expression constructs described below. Heterologous expression of the [4FeH] (apo-) and holo- protein [6FeH] of *Chlamydomonas reinhardtii* HYDA1 as well as of *Clostridium pasteurianum* HYDA (CPI) was carried out in *E. coli* strain BL21 (DE3)  $\Delta iscR^{58}$  according to the previously published procedure<sup>13,14,32</sup>.

**Cloning of HIS**<sub>6</sub>**-tagged** *C. reinhardtii* **HYDA1** and *C. pasteurianum* **CPI** variants. *E. coli* codon optimized cDNA of *HYDA1* or gDNA of *CPI* both fused at their 3'end to a short DNA sequence that encodes a Strep-Tag II affinity tag and cloned into pET21b<sup>14,32</sup> were used as templates for the respective amplifications with Pfu DNA polymerase. For the PCR based C-terminal fusion of the HIS<sub>6</sub>-tag sequence to HYDA1 the oligonucleotides 5'-CTCGATCCCGCGAAATT-3' (forward), binding upstream of the *HYD* genes and 5'-CGG**GTCGAC**TTA*GTGATGATGGTGGTGATG*GGGCTGATTTTTTCATCTTTT TC-3' (reverse) were used. For the C-terminal fusion of HIS<sub>6</sub> to CPI the same forward primer, but the CPI specific reverse primer 5'-CGG**GTCGAC**TTA*GTGATGATGATGATGATGGTGGTGATGGTGATG*GGCTGATTT TTTATATTTAAAGTGCAGGATTTCATGGGCACGACCTTC-3' was used. Bold letters indicate the restriction site for *Sal*I which together with *Nhe*I was used for cloning, while bold italic letters indicate the HIS<sub>6</sub>-tag coding sequence. The amplified constructs were again cloned into the expression vector pET21b using *Nhe*I and *Sal*I restriction sites and the resulting plasmids verified by DNA-sequencing. The resulting constructs pET21b-HydA1opt-6H and pET21b-CpIopt-6H facilitated a regulated expression via the IPTG inducible lac-promoter.

Heterologous production and purification of recombinant HYDA1 and CPI. E. coli BL21 (DE3)  $\Delta$ iscR expression strains<sup>58</sup> were generated via electrotransformation with constructs pET21b-HydA1opt or pET21b-CpIopt encoding either HIS<sub>6</sub>- or Strep-tag II fusion proteins, commonly without, but also with maturation factors HYDE,F,G,X encoded on a second expression vector pACYCDuet-1, for determining the specific activity of HYDA1[4FeH]+[2FeH] (Supplementary Tab. S1)<sup>32</sup>. Cultivation and enzyme purification via Strep-tag II affinity tag was performed as previously described<sup>14,32</sup>. For purification with Ni-NTA IMAC, E. coli cells, resuspended in 100 mM Tris-HCl (pH 8), plus 2 mM sodium dithionite (NaDT) were lysed by ultrasonication (five times 30s; output 25; Branson Sonifier 250). Cell debris was removed via centrifugation (60 min, 200.000  $\times$  g, 4 °C). For IMAC purification the supernatant was diluted with an aqueous solution containing imidazole and NaDT to final concentrations of 50 mM Tris-HCl (pH 8), 10 mM imidazole, and 2 mM NaDT. The resulting solution was passed through a sterile filter (pore size  $0.2 \mu m$ ). Afterwards the filtrate was loaded on a 4 mL gravity flow Ni-NTA fast-flow column pre-equilibrated with 50 mM Tris-HCl (pH 8), supplemented with 10 mM imidazole and 2 mM NaDT. After washing with 50 mM Tris-HCl (pH 8) containing 20 mM imidazole and 2 mM NaDT the protein was eluted from the column using 100 mM imidazole. Purified proteins were stored at -80 °C until further use after concentrating with Amicon centrifugal filter units (30 kDa cutoff). Protein concentration was determined via spectrophotometry (NanoDrop, Peqlab) at  $\lambda = 280$  nm. Molecular weight and purity of the eluated protein fractions were analyzed via denaturing sodium dodecylsulfate polyacrylamide gel electrophoresis (SDS-PAGE) using Coomassie brilliant blue staining according to standard techniques (Supplementary Fig. S1b). HYDA1[6FeH] and CPI[6FeH] samples were prepared by in vitro maturation of concentrated HYDA1[4FeH] and CPI[4FeH] expression products by addition of the chemically synthesized 2Fe<sub>adt</sub> cofactor with subsequent purification using a desalting column as described previously<sup>14</sup>.

**Lyophilization of [FeFe]-hydrogenases.** The hydrogenases, either [4FeH] or [6FeH] proteins supplemented with 100 mM NaDT, were dispensed in PCR-tubes with a pierced lid. Sample tubes were anaerobically frozen in a round-bottom flask filled with liquid nitrogen and subsequently attached to a freeze-dryer (LyoVac GT 2; Leybold - Heraeus) for 24 h. Protein concentration ranged from 1 to 110 mg mL<sup>-1</sup> for HYDA1 and was 0.5 mg mL<sup>-1</sup> for CPI. After water sublimation the sample-containing flask was sealed airtight under low pressure and the lyophilized samples were reintroduced into the anaerobic tent. To avoid sample humidification, the proteins powder was hermetically sealed and stored at -80 °C until further use. The temporary cryogenic sample storage did not affect the protein properties in any way.

 $O_2$ -exposure of lyophilized [FeFe]-hydrogenases. The lyophilized proteins were continuously gassed with dry pure  $O_2$  gas at room temperature for 1–8h. For 24h  $O_2$ -exposure, samples obtained after  $O_2$ -gassing for 8h were incubated for another 16h in an atmosphere of dry pure  $O_2$  in airtightly sealed suba vials. After the  $O_2$ -treatments, the samples were purged with argon for 10 min before activity measurements were carried out.

**Exposure of lyophilized [FeFe]-hydrogenases to defined moisture levels and O<sub>2</sub>.** Humidified  $O_2$  gas was prepared by filling the headspace of an airtight 20 mL suba vial containing 3 mL of 100 mM potassium phosphate buffer (pH 6.8) with pure  $O_2$  gas followed by a subsequent incubation over night at 80°C in a heating cabinet. For exposure of lyophilized hydrogenases to oxygen gas with increasing levels of humidity, protein aliquots were first flushed for 15 min with dry pure  $O_2$  gas. Afterwards, defined volumes of the water-saturated  $O_2$  gas were injected into airtight sealed sample-containing vials. Protein samples were exposed to the humidified  $O_2$ -atmosphere for 30 min at room temperature. Immediately before determining the specific H<sub>2</sub>-evolution activity the sample vials were anaerobized by flushing with argon for 15 min.

**O<sub>2</sub>-exposure of [FeFe]-hydrogenases in solution.** The two purified [FeFe]-hydrogenases from *C. reinhardtii* and *C. pasteurianum* either with or without the [2FeH] subsite were exposed for 15 min to different concentrations of O<sub>2</sub>. Therefore, solutions of 100 mM potassium phosphate pH (6.8) supplemented with 2 mM NaDT and 6.33 nM of the particular enzyme were flushed first with argon for 3 min, then different volumes of dry pure O<sub>2</sub> gas were injected into the airtight sealed reaction tubes, and the samples were incubated for 15 min at 37 °C in a shaking water bath. For anaerobization, the samples were purged with argon for 5 min and specific H<sub>2</sub>-evolution activities were determined immediately.

*In vitro*  $H_2$ -evolution activity assay. For determining  $H_2$ -evolution activities of [FeFe]-hydrogenase samples, the previously described *in vitro* activity assay was used<sup>59</sup>. To evaluate the maturation competence and thus integrity of hydrogenase apoenzymes ([4FeH]), the proteins were mixed with a 10-fold excess of the synthetic 2Fe<sub>adt</sub> complex immediately prior to activity determination<sup>14</sup>.

**Preparation of samples for XAS on lyophilized HYDA1.** HYDA1[4FeH] expressed in *E. coli* was purified and either used as prepared or after *in vitro* maturation as HYDA1[6FeH]. Samples of HYDA1[4FeH] and HYDA1[6FeH] were concentrated to 2.2 mM protein and NaDT was added to a final concentration of 100 mM.  $16\mu$ L sample volume was injected under anoxic conditions into Kapton-covered acrylic-glass sample holders using a Hamilton syringe<sup>21</sup>. Sample holders loaded with 2 mM of [FeFe]-hydrogenase protein were immediately frozen in a round-bottom flask containing liquid nitrogen and attached to the freeze-dryer for 24h as described above. Afterwards the lyophilized samples were immediately frozen or exposed for 15 min to dry pure O<sub>2</sub> gas and flushed thereafter with argon for 5 min before storage at -80 °C. Storage of the samples at -80 °C was done in a plastic zip bag, enclosed in a 50 mL falcon tube filled with Silica gel 60 to exclude any humidity. In addition, lyophilized HYDA1[6FeH] samples in XAS sample holders were exposed to air at ambient humidity for ~5 days prior to the XAS experiments.

**X-ray absorption spectroscopy.** XAS at the Fe K-edge was carried out at beamline KMC-1 at BESSY (Helmholtz Zentrum für Materialien und Energie Berlin) with the storage ring operated in top-up mode (200 mA). A Si[111] double-crystal monochromator was used for incident energy scanning and a 13-element energy-resolving germanium detector (Canberra) was used for Fe X-ray fluorescence detection (shielded by  $10 \mu$ M Mn foil against scattered X-rays) in a standard XAS set-up<sup>25</sup>. The spot size on the sample was set by slits to about 5 (horizontal) × 0.5 (vertical) mm<sup>2</sup>. Samples were held in a liquid-helium cryostat (Oxford) at 20 K. The absorption spectrum of an Fe foil was measured in parallel and its first inflection point at 7112 eV was used for energy calibration<sup>26,28</sup>. Up to 4 XAS scans of ~30 min duration were carried out per sample (1 scan per spot) and dead-time corrected spectra were averaged, normalized, and EXAFS oscillations were extracted ( $E_0 = 7112 \,\text{eV}$ ) as previously described<sup>25</sup>. Fourier-transform (FT) calculation (using cos windows extending over 10% at both *k*-range ends) and least-squares fit analysis of unfiltered EXAFS spectra<sup>25</sup> were carried out using in-house software<sup>25</sup>. Phase functions were calculated by FEFF7<sup>60</sup> ( $S_0^2 = 0.9$ ;  $E_0$  was refined to 7115 ± 2 eV in the EXAFS fits).

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#### Acknowledgements

T.H. thanks the Cluster of Excellence RESOLV (EXC1069) funded by the Deutsche Forschungsgemeinschaft (DFG) and the EU (Sun2Chem). M.W. gratefully acknowledges support from the Volkswagen Foundation (LigH2t). M.H. thanks the DFG (Ha3265/5-1) and the German Bundesministerium für Bildung und Forschung (grant 05K14KE1 within the Röntgen-Angström Cluster) for funding and the DFG for a Heisenberg Fellowship. We thank H. Dau (FU-Berlin) for access to XAS equipment at BESSY, F. Schäfers and M. Mertin at beamline KMC-1 of BESSY (HZB) for technical support, and P. Schrapers, S. Mebs, and N. Schuth (FU-Berlin) for help in XAS data collection.

#### **Author Contributions**

J.N., M.W. and T.H. conceived and designed experiments. J.N., R.K. and M.H. performed the experiments. J.N., M.W. and T.H. analyzed the data. K.K. provided the 2Fe<sub>adt</sub> complex. R.K. and M.H. performed and analyzed the XAS experiments. J.N., M.W., M.H. and T.H. wrote the manuscript.

#### Additional Information

Supplementary information accompanies this paper at http://www.nature.com/srep

**Competing financial interests:** The authors declare no competing financial interests.

How to cite this article: Noth, J. *et al.* Lyophilization protects [FeFe]-hydrogenases against O<sub>2</sub>-induced H-cluster degradation. *Sci. Rep.* **5**, 13978; doi: 10.1038/srep13978 (2015).

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