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A dimanganese complex containing a redox-active elongated disulfide bond electrocatalyses proton reduction.
Electrocatalytic Proton Reduction Catalyzed by A Dimanganese Disulfide Carbonyl Complex Containing a Redox-active Internal Disulfide Bond

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A dimanganese hexacarbonyl complex [(Mn(CO)\(_3\))\(_2\)(\(\mu\)-SC\(_6\)H\(_4\)-o-S-S-C\(_6\)H\(_4\)-o-\(\mu\)-S-)] containing an elongated disulfide bond electrocatalyses proton reduction at moderate overpotentials of 0.55 to 0.65V. Cyclic voltammetric, infrared spectroscopy and computational studies suggest that the redox-active sulfur atoms of the disulfide bond serve as the initial reduction site.

Since crystallographic methods revealed that the Fe\(_2\)S\(_2\) core was essential to the function of [FeFe]hydrogenase,\(^{1}\) hundreds of synthetic models have been prepared to mimic the active site of the hydrogenase.\(^{2,4}\) Among the model complexes, the electrocatalytic proton reduction performance of benzenedithiolate ligands bridged diiron complexes has been evaluated.\(^{5}\) An iron hydride intermediate was proposed to act as a hydride donor and reacted with the acid to generate dihydrogen heterolytically. This mechanism was further supported by other reports where iron hydride complexes were characterized by X-ray crystallographic methods.\(^{6,8}\)

Although many diiron catalysts have been used as hydrogenase mimics, dimanganese model complexes have so far not been reported. Our group has shown that thiols could be transformed into disulfides and H\(_2\) using Mn(CO)\(_5\)Br or CpMn(CO)\(_3\) under uv-visible light.\(^{9,10}\) Since enzymes such as glutathione reductase and cysteine reductase can easily reduce the respective disulfides into thiols using proton sources, combining the features of the manganese complexes and the enzymes into one may convert H\(_2\) into H\(_2\) without sacrificing thiols in the process through the reversible dissociation and formation of the S-S and S-H bonds.

In this report, we show that an easily-prepared dimanganese disulfide hexacarbonyl complex, [(Mn(CO)\(_3\))(\(\mu\)-SC\(_6\)H\(_4\)-o-S-S-C\(_6\)H\(_4\)-o-\(\mu\)-S-)] (complex I, see Scheme 1) can catalyze electrochemical proton reduction using acetic acid as the proton source through the initial reduction of redox-active sulfur atoms of the internal disulfide bond of the complex. The mechanism of the proton reduction process is also investigated by cyclic voltammetry, infrared spectroscopy and density functional theory (DFT) calculations.

Scheme 1. The structure of complex I.

Although the synthesis of I has been reported before, a simpler way of preparing I is used here instead.\(^{11}\) Broadband UV photolysis of decacarbonyldimanganese Mn\(_2\)(CO)\(_{10}\) with two equivalent of 1,2-benzenedithiol in hexane for 2 hours generates brown I in good yield. Suitable crystals of I are grown in dichloromethane/hexane solvent and stored at low temperatures before being subjected to X-ray diffraction (see Supporting Information). As the X-ray crystal structure of I has already been described previously, only the salient features of the structure are mentioned.\(^{11}\) Complex I consists of two six-coordinate Mn(I) atoms where each manganese center is coordinated to three sulfur atoms and three terminal carbonyl groups in a distorted octahedral geometry. The internal disulfide bond length in complex I of 2.2061(7)Å (see Supporting Information), is significantly longer than the free disulfide bond in diphenyl disulfide bond (2.02Å).\(^{12}\) There is no Mn-Mn bond as the distance between the two metal centers is too far at around 3.35 Å. Each of the two Mn centers is formally in the +1 oxidation state. The Mn(CO)\(_3\) moiety adopts a near-C\(_{3v}\) symmetry about the metal center as reflected in the v\(_{\text{CO}}\) infrared spectrum (1944, 1965, 2022, 2041 cm\(^{-1}\) in CH\(_3\)CN solvent).
Fig. 1 Cyclic voltammograms of 1 (1 mM) in 0.1 M Bu4NPF6 in DMF at a scan rate of 100 mV/sec. 3 mm glassy-carbon working electrode, platinum counter electrode at 295 K.

The cyclic voltammetric diagram of 1 in DMF in the absence of acid shows three reduction peaks at -1.11 V, -1.38 V and -2.10 V vs Fc+/Fc (Figure 1). Instead of acetonitrile, DMF is the solvent of choice for cyclic voltammetry studies because of the much greater solubility of 1 in DMF. Drawing analogy to a previous study on an organic disulfide,13 the first peak is assigned to the disulfide bond being reduced by one electron at -1.11V vs Fc+/Fc to the radical anion complex 1- (Scheme 2). At -1.38V vs Fc+/Fc, reduction of 1- by a second electron to a dianion complex 12- occurs. We believe that this complex dissociates to form a monomanganese anion complex 2-, a previously-detected stable ionic complex.11 Complex 2- can be further reduced by an electron to form another monomanganese complex 22- at -2.10V vs Fc+/Fc.14 Only the third step is observed to be reversible. Further evidence for the formation of 2- and its reactions will be discussed later.

Scheme 2. Proposed pathways for the electroreduction of complex 1.

The DFT (B3LYP/6-31+G(d)) calculations also support these proposed electroreduction pathways. The Mulliken charge of each of the two sulfur atoms of the disulfide bond changes from 0.205 to 0.058 to -0.116 in complex 1, 1-, 12- respectively. However, the corresponding Mulliken charge of Mn in each of the three complexes changes very slightly, hence suggesting that the disulfide bond is reduced instead of the manganese. The LUMO calculated for 1 is shown in Figure 2. It shows a mixture of aromatic ring character and antibonding character between the sulfur atoms, which would promote S–S bond cleavage upon electron occupation. The calculated disulfide bond is 2.28 Å, in good agreement with the experimental value. Indeed, the first electron upon reduction goes into the SOMO orbital of 1- which is dominated by S–S antibonding character and causes the S–S distance to lengthen from 2.28 Å to 3.10 Å. The second electron upon reduction further lengthens the disulfide bond in 12- to 4.32 Å after which 12- is expected to dissociate to form 2-. The dissociation of 2- to two 12- species is supported by the large computed exoergicity value of -54.4 kcal per mol of 12-.

Fig. 2 (a) LUMO (lowest unoccupied molecular orbital) of 1 (b) SOMO (singly occupied molecular orbital) of 1- (c) HOMO (highest occupied molecular orbital) of 12-. Color representations: grey is for carbon, white is for hydrogen, red is for oxygen, yellow is for sulfur, purple is for manganese. Bond lengths are shown in Å.
The electroreduction of protons by complex 1 carried out under anaerobic conditions is discussed next. A control experiment is first carried out to determine the potential at which acetic acid in DMF is reduced without the catalyst. However, no peaks were observed even when the potential (vs Fe/C) is scanned to -2.5 V. In contrast, when 5 equivalent amount of CH$_3$COOH is added into the DMF solution containing 1, the peak at -2.10 V increases dramatically. This peak increases in intensity and shifts gradually to -2.20 V as more acid is added. However, the positions of the first two reduction peaks at -1.11 V and -1.38 V remain unaffected. The potential at -2.10 V is thus attributed to the catalytic proton reduction potential of 1, which corresponds to an overpotential of 0.55-0.65 V (CH$_3$COOH standard reduction potential in DMF is at -1.55 V vs Fe/C). The electrocatalytic H$_2$ production was further confirmed by bulk electrolysis of 1 with excess CH$_3$COOH in DMF at -2.10 V vs Fe/C with a Faraday yield consistently above 80% in a series of runs.

Fig. 3 Cyclic voltammograms of 1 (1 mM) with CH$_3$COOH (0–50 mM) in 0.1 M Bu$_4$NPF$_6$ in DMF at a scan rate of 100 mV/sec. 3 mm glassy-carbon working electrode, platinum counter electrode at 295 K.

To further understand the cyclic voltammogram, stoichiometric reactions involving complex 1 are carried out in acetonitrile solvent. Upon addition of NaB$_4$H$_4$, the color of the solution changes from brown to red immediately. The ν$_{CO}$ peaks at 1916 and 2010 cm$^{-1}$ disappear while at the same time the signals of CO brown to red immediately. The ν$_{2041}$ cm$^{-1}$ upon addition of NaBH$_4$ (1 mM) in 0.1 M Bu$_4$NPF$_6$ in DMF runs.

Scheme 3. Proposed mechanism for the proton reduction process.

However acetic acid addition does not reverse the reaction, although this observation is consistent with our cyclic voltammetry data interpretation whereby proton reduction using acetic acid as the source does not take place immediately after 2$^{-}$ is generated at the second reduction peak of -1.38 V.

The mechanism of proton reduction by complex 1 is proposed here (see Scheme 3). As discussed previously, complex 1 can be reduced to eventually generate the dianionic monomanganese complex 2$^{-}$. The reduction is facilitated by taking place at the S atoms of an elongated disulfide bond within 1. Upon addition of acetic acid, 2$^{-}$ is protonated to form either 2H$_{M^+}$ or 2 H$_{M}^{-}$. The proton is coordinated to one of the sulfur atoms in the former complex or to the Mn center in the latter one. Regardless of the initial proton coordination, we propose that hydride transfer from sulfur to manganese or vice-versa takes place in a facile manner. Either isomer is further protonated to form a common neutral intermediate 2H$_{M}$H$_{S}$. The subsequent interaction between the two coordinated H atoms eventually produces dihydrogen via a complex 2HH and leaves behind a 16-electron neutral intermediate 2. Finally, the dimerization of two intermediate 2 regenerates complex 1.
Further support for the proposed mechanism is obtained from DFT calculations. The structures of the complexes are first optimized using B3LYP/6-31+G(d) level of theory. Then single-point energy calculation incorporating a solvation model at both B3LYP/6-311++G(d,p) with SMD/B3LYP/6-31G(d) or M06/6-311++G(d,p) with SMD//M06/6-31G(d) levels of theory is carried out on the optimized structures. Two important features of the mechanism are explored in more detail. The first study shows that the $2\text{H}_5^-$ and $2\text{H}_2\text{M}_5^-$ isomers are energetically similar and can perhaps interconvert at room temperature. The free energy diagram in Figure 5a shows the transition state $\text{TS1}$ of the isomerization between $2\text{H}_5^-$ and $2\text{H}_2\text{M}_5^-$. Intermediate $2\text{H}_5^-$ undergoes intramolecular hydrogen shift from sulfur to manganese through $\text{TS1}$ with a small energy barrier $(6.1(5.9)$ kcal/mol) to form $2\text{H}_2\text{M}_5^-$, which itself is $4.0(3.9)$ kcal/mol more stable than $2\text{H}_5^-$. The distance of the S-H bond lengths from 1.36 Å to 1.60 Å while the Mn-H distance shortens from 1.72 Å in $\text{TS1}$ to 1.60 Å in $2\text{H}_2\text{M}_5^-$. In the second part of the calculation, we show that the formation of dihydrogen from $2\text{H}_2\text{M}_5^-$ is exergonic and also with a small activation barrier. The free energy diagram of $2\text{H}_2\text{M}_5^-$ to $2\text{HH}$ through the transition state $\text{TS2}$ is shown in Figure 5(b). The distance between the two H1 and H2 atoms shortens from 2.60 Å to 1.29 Å in $\text{TS2}$ with a 7.9(9.3) kcal/mol energy barrier with respect to $2\text{H}_2\text{M}_5^-$. The intermediate $2\text{HH}$ is 9.4(9.7) kcal/mol more stable than $2\text{H}_2\text{M}_5^-$. The authors would like to thank a NUS research grant (143-000-553-112).

**Notes and references**