Bio-Inspired H₂ Production Catalysts

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Abstract

Hydrogen is considered as one of the best possible energy vectors that can effectively store the otherwise intermittent solar energy for proficient renewable energy usage. Conversion of protons to hydrogen is the heart of this scheme and this reaction requires the presence of a catalyst. Currently, the expensive Platinum (Pt)-based materials are available as one of the best catalysts but its low abundance is a barrier for its worldwide usage. Naturally occurring [FeFe]- and [NiFe]-hydrogenases can also efficiently perform the H₂ production reaction albeit only in a narrow chemical space (25-40 °C, pH 4-8, mostly in absence of O₂). In the past decade, hydrogenase enzyme structures acts as the template for the development of numerous structural and functional model complexes. Macro cyclic cobalt complexes and nickel bis-(diphosphine) complexes containing pendant basic groups have emerged as most efficient catalysts, which are active even in heterogeneous conditions (immobilized on electro active surfaces).

Keywords: Bio-Inspired; Mimics; Catalyst; Hydrogen; Hydrogenase; Amino Acid

Abbreviations: Ang: Arginine; Bz: Benzyl; Gly: Glycine; HER: Hydrogen Evolution Reaction; kDa: KiloDalton (mass unit); OCS: Outer Coordination Sphere; P,N₂: 1,5-diaza-3,7-diphosphacyclooctane; Phe: Phenyl; Py: Pyridine; SHE: Standard Hydrogen Electrode; TFA: Trifluoroacetate

Introduction

Currently, more than 85% of our energy requirement is fulfilled by the carbon based fossil fuels such as coal, oil, and natural gas [1]. The increasing human population and improving life style in modern society have affected a steep increase in global energy consumption, which in turn increased the atmospheric CO₂ emission produced from the fossil fuels. Recently, the atmospheric CO₂ level has surpassed 400ppm and this has impacted adversely on climate and the biosphere [2]. Thus, an alternative energy resource is sought for a solar energy is considered as one of the most promising one due to its abundance. However, the intermittent nature of the solar energy requires that it is stored properly, preferably in the form of chemical bonds, for its effective usage. Hydrogen has emerged as one of the best candidates for that role of energy carrier as it provides clean energy along with high energy density (energy density 140MJkg⁻¹) [3,4]. The following reaction is the heart of this scheme of storing solar energy into H–H bonds.

$$2H^+ + 2e^- \rightarrow H_2$$

The conversion of protons to H₂ requires a catalyst and currently two best available options are: (1) Platinum (Pt) based materials and (2) the natural enzyme hydrogenases. Both these systems can efficiently catalyze the reaction and can be utilized solar energy driven H₂ production [5,6]. However, both of them have their shortcomings that have limited their usage for worldwide solar H₂ production. The Pt-based materials are active in various reaction conditions, even in the harshest ones (strongly acidic, high temperature) but Pt is one of the most expensive metals due to its low earth abundance. On the other hand hydrogenases contain abundant Ni or Fe, but they are active only in a narrow chemical space (25-40 °C, pH 4-8, mostly in absence of O₂) [7,8]. In addition to that, the active site concentration is very low for hydrogenases, where it has only one active site in protein that has size >50kDa [9]. This last factor increases the operational cost for a solar energy driven hydrogen production unit that uses hydrogenases as catalyst. However, the hydrogenase enzymes provide the basic structural blue print that can be utilized to produce synthetic catalysts with analogous activity but in a broader chemical space.

Three types of hydrogenase enzyme are found in nature. They are binuclear [NiFe]-, [FeFe]- and mononuclear [Fe] hydrogenase as per the identity of the metals in the active site of the corresponding enzymes [10-12]. Thus, these enzymes contain only readily available first-row transition metals. In biology, the source of protons are generally water, thus it creates a carbon foot print free H₂ cycle [3]. Hydrogen is relatively weak base but it acidity increases after binding with metal [13], thus facilitates reversible heterolytic cleavage and generation of hydrogen. It is interesting to mention that Pt-based materials follow a homolytic process for H₂ production.

The overall catalytic cycle for H₂ production by hydrogenases consists of the following steps:

1. Movement of protons from the bulk to the metal active site,
2. Reduction of metal bound proton to hydride (the metal is oxidized in this step),
3. Reaction of the hydride with a neighboring proton to produce \( \text{H}_2 \).

4. Release of \( \text{H}_2 \) to the bulk.

5. Reduction of the metal center to re-activate it for the next cycle.

The hydrogenases are one of the most efficient catalysts as all the above mentioned steps are poised at thermodynamically reversible states. This is possible due to the presence of the protein scaffold around the metal center. The protein structure ensures fast movement of protons and electrons via proper positioning of basic groups and iron sulfur clusters, respectively [14]. The protein structure also minimizes the energy requirement for the structural changes of the metal active site during redox changes by inducing entatic states [15]. Thus, the surrounding protein structure plays a crucial role and it is known as the outer coordination sphere (OCS) [16].

In last few decades, bio-mimetic catalysts are mainly developed considering only the metal active site of hydrogenases and a plethora of complexes such as Cobalt dioximes, Cobalt dimine, Cobalt and Nickel diphosphines, Cobalt, Molybdenum and Ruthenium polypyridine, bimetallic dithiolato complexes and macrocyclic Cobalt complexes are produced. Although structurally some of the complexes are analogous to the enzyme active site they rarely matched the enzymatic efficiency for \( \text{H}_2 \) production. Thus, it clearly indicated that the protein based OCS features are also needed to be included in the bio-inspired catalyst design. Recently, Shaw and her co-workers have developed a number of bio-inspired Ni-bis-(diphosphine) complexes containing peripheral amino acids in the structure that imitates the enzyme’s OCS scaffold. Interestingly, these complexes improved the catalytic efficiency compared to their parent complexes without the model OCS features. Artero et al. have demonstrated similar results when they included a synthetic Fe cluster into an apo-protein of hydrogenase to produce a fully active holoenzyme [17]. Some of these new generation bio-inspired catalysts are also immobilized on electro active surfaces and they exhibited fairly improved reactivity compared to hydrogenase enzyme in terms of reaction conditions. Current aspects for electrochemical hydrogen production by the series of these bio-mimetic catalysts are critically reviewed here.

**Method and Parameters for evaluation of the synthetic catalysts**

The catalytic reactivity for the synthesized complexes are evaluated via various techniques, among which electrochemical methods are the most commonly, utilized one. The two most essential parameters that determine the efficiency of a catalyst are: (i) turn over frequency (TOF) that defines the rate of the catalytic cycle and (2) over-potential, which is the excess energy required beyond the thermodynamic potential for a particular catalytic reaction (e.g. thermodynamic potential for \( \text{H}^+/\text{H}_2 \) couple is 0.0V vs SHE at standard conditions). Various catalysts can be compared by those two parameters and catalysts active at a low over-potential with high TOF is considered to be better catalysts. Hydrogenase enzymes and Pt metal are the gold standard for \( \text{H}_2 \) production catalysts and both of them exhibit TOF of ~20,000s\(^{-1}\) and over-potential of <100mV.

Both TOF and over-potential can be measured very efficiently from simple cyclic voltammetry (CV) experiments. In this procedure, at first the electrochemical response of a complex is recorded under non-catalytic conditions and the stoichiometric response in current (\( i_p \)) is measured. Then, an analogous experiment is done under catalytic conditions and the catalytic current (\( i_c \)) is recorded. Then the TOF or \( k_{obs} \) can be measured by using the following equation:

$$\frac{i_c}{i_p} = \frac{n}{0.4463} \left( \frac{RTk_{obs}}{Fv} \right)^{1/2}$$

Here \( R \) is universal gas constant, \( F \) is Faraday’s constant, \( T \) is the temperature in K, \( v \) is the scan rate in Vs\(^{-1}\) and \( n \) is the number of molecules involved in the catalytic reaction.

In certain cases (e.g. acidic aqueous solution), it is difficult to measure the stoichiometric current (\( i_p \)) as it difficult to find a non-catalytic condition under similar reaction set up. In these scenarios, the TOF or \( k_{obs} \) can be measured by using the following equation by using only the catalytic current (\( i_c \)):

$$k_{obs} = \frac{1}{D} \left( \frac{i_c}{nFAC_{cat}} \right)^{2}$$

Here \( D \) is the diffusion coefficient of the complex under catalytic condition, \( F \) is Faraday’s constant; \( A \) is the surface area of the electrode, \( [\text{Cat}] \) concentration of the catalytic complex and \( n \) is the number of molecules involved in the catalytic reaction.

The over-potential is measured as the difference of the potential where the catalytic reaction is occurring and the thermodynamic potential of the concerned catalytic reaction.

Now, the potential of the catalytic reaction can be measured at various positions such as

(i) Foot of the wave, where the catalytic response begins,

(ii) At half maxima of the current, where the catalytic response is 50% of the maximum,

(iii) At the maximum catalytic current. The calculation can be done with any of the three conditions once it is mentioned properly. During the comparison of over-potential among various catalysts the same calculation method should be used.

Coulometric technique is also used for measuring the bulk electrocatalytic properties of complexes. In this chronoamperometric experiment, the current is measured at a particular potential (typically where the catalytic reaction occurs) over time. In this experiment, the overall charge (i.e. no. of electrons) passed can be measured. These experiments are generally performed in a closed system and the end of the experiment the \( \text{H}_2 \) produced (for \( \text{H}^+ \) to \( \text{H}_2 \) conversion catalysis) in the head space can be measured via...
Discussion

Hydrogenase enzymes are an integral part of many microorganisms. Among three distinct types of hydrogenase enzyme, [FeFe]-hydrogenase are the best H₂ production catalyst with turnover frequency reported 20,000s⁻¹ [18]. But these enzymes cannot be used directly in application system due to their bulky size and other large-scale synthesis problems along side with their narrow chemical space issues. However, some of their unique structural aspects can be incorporated into various other bio-inspired mimics. For our formal discussion, we divide them into four categories. These are Clathrochelate complexes, Polypyridine complexes, Complexes with outer coordination scaffold and surface supported catalyst.

Bio-inspired clathrochelate complexes

Fisher & Eisenberg [19] first reported water soluble Co(II) Tetraazomycrocycle (2a, 2b) can produce H₂ at an applied over-potential of -1.26V and -1.36V Vs SHE respectively on acetonitrile and water solvent with 80% Faradic Yield. Kellett & Spiro [20] studied on three water-soluble Co (I) porphyrins, CoTMP (1a), CoTPyP (1b) and CoTMPyP (1c) developing towards efficient hydrogen production catalyst. They exhibit catalytic activity at around -0.76V vs SHE in 0.1M TFA over 90% faradic yields on Hg pool electrode. Later Grätzel et al. [21] Co(Sepulchrate)³⁺ (3) complex shows catalytic activity at lower over-potentials (-0.46V Vs SHE) in weakly acidic medium (pH 4 phosphate buffer), but the faradic yield is much lower to 55%. A series of Boron and Fluorine capped clathrochelated tris glyoximato cobalt complex (4a to 4d) has been synthesized, among them 4a and 4b can reduce HClO₄ to produce H₂ even at very low over-potential 0.19V vs SHE [22] but in case of Fluorine capped clathrochelated complexes (4c and 4d) –Cl group is replaced by phenyl and methyl groups its lability of metal center decrease. As a result of that, systems becomes weak efficient having high over-potential and low faradic yield [23]. When this clathrochelate cobalt oximes has been compared with various annlated glyoxymato cobalt complexes (5) in non-aqueous solvent. Annalated cobalt oximes are more active with decreasing its bulkiness around active metal centre. Annalated cobalt oximes suffer from lack of planarity so there is a room to make it even more active with by achieving planar structure (Figure 1).

Bio-inspired polypyridine catalysts

Molybdenum and Cobalt pentapyridine complex are very efficient in neutral aqueous medium but suffer from large over-potential. Due to neutral as well as strong field ligand nature of pyridine it forms series of complex with Co (II). Complexes 6a to 6c can reduce water at neutral pH. 6a shows 99% faradic efficiency at -1.30V vs SHE. Incorporating electron withdrawing group (6b) at para position decrease the over-potential similarly over-potential increases (6c) when -NMe₂ (electron donating group) group introduced. Similar type of pentapyridine Molybdenum ([PY₅Me₂])
MoO$_2^{2+}$ (7) has more efficiency than that of cobalt complexes but their catalytic activity on electrode other than mercury pool is not reported [24] as well as hydrolyzes on acidic water. A similar type of Molybdenum disulfide complex (8) is stable towards the acidic water and shows good catalytic activity [25]. Another important Pyridine based cobalt catalyst [Co(Py,OMe)]$_2^{2+}$ (9) which can produce hydrogen with 99% Faradic yield in presence of TFA as a proton source on acetonitrile [26]. In 2011, Nocera et al. [27] reported macrocycle porphyrin CoHPX-CO$_2$H (10a) for the effective hydrogen production catalyst and interestingly describe the effect pendant acid group adjacent to working metal center (Hangman contribution) by comparing with its Bromo (10b) variant (Figure 2).

**Bio-Inspired catalyst having OCS**

Mononuclear Ni(II) complex having cyclic diphosphine ligand are on-going interest for green energy hydrogen evaluation chemistry. For the sake of our discussion, we grouped these Ni(II) cyclic diphosphine into two category. One is having pendant amine in outer coordination sphere (OCS) and the other type system containing amino acid scaffold.

![Figure 2: Various polypyridine hydrogen production catalyst.](image)

![Figure 3: Various model system of Nickel(II) cyclic diphosphine catalysts.](image)
Nickel(II) Cyclic diphosphine containing pendant amine: DuBois et al. [28] while working with [Ni(depp)]$_2$ [11] found that it has potential to oxidize hydrogen in the presence of external amine [29]. Then they tried to introduce nitrogen backbone inside it [30]. Catalytic rate was still very low. Pendent amine has to be tuned in such a way that amine and metal center involved in the catalytic cycle jointly. Later similar type of modified catalyst has been synthesized [Ni(P$_2$N$_2$Ph$_2$)(CH$_2$CN)$_2$] (12a), [Ni(P$_2$N$_2$Ph$_2$)X$_2$] (R=H [31-34], -OMe [35,36] and [Ni(P$_2$N$_2$Ph$_2$)(ppy)(OEt)$_2$] [37,38] (P$_2$N$_2$Ph$_2$=1,5-diaza-3,7-diphosphacycloctane). Complex 12a shows efficient catalytic activity for hydrogen production, but others are mainly active for H$_2$ oxidation. [Co(P$_2$N$_2$Ph$_2$)(CH$_2$CN)$_2$] (12b) shows activity at relatively high over-potential and partially dissociate into (P$_2$N$_2$Ph$_2$) during electro catalysis [32]. Also, series of other active [Ni(P$_2$N$_2$X$_2$)$_2$] (X=OMe, CF$_3$, Br, H, Me, CH$_2$P(O)(OEt))$_2$ systems have been synthesized [33]. Tuning the proton relay position and chance of enhancing endo isomer formation [Ni(P$_2$N$_2$Ph$_2$)]$^{2+}$ (13) shows extremely high turnover frequency, almost 1,000,000$^{-1}$ in dry acetonitrile [34] (Figure 3).

Nickel(II) Cyclic diphosphine containing amino acids: Successful attempts were made to incorporate carboxylic ester function into the neighbors of the active site that will continuously relay proton source. [Ni(P$_2$N$_2$Ph$_2$)$_2$]$^{2+}$, R=COOEt, -CO-alanine-methyl ester, -CO-phenyl alanine-methyl ester [35] shows less activity than it parent R=H. The inclusion of single amino acid in the pendant amine groups increase the reactivity and resolves the problem of durability and water solubility [39]. First generation [Ni(P$_2$N$_2$Ph$_2$)$_2$](X=OMe, CF$_3$, Br, H, Me, CH$_2$P(O)(OEt))$_2$ systems have been synthesized [33]. Tuning the proton relay position and chance of enhancing endo isomer formation [Ni(P$_2$N$_2$Ph$_2$)]$^{2+}$ (13) shows extremely high turnover frequency, almost 1,000,000$^{-1}$ in dry acetonitrile [34] (Figure 3).

Surfaced supported catalyst: [Ni(P$_2$N$_2$Ph$_2$)$_2$]$^{2+}$ immobilization was the first successful attempt to direct comparison between [NiFe] and Bio-inspired synthetic enyme on the working electrode. Its TOF increase with decreasing pH and need 146mV more over-potential that of hydrogenase modified Pt electrode at pH 5. At low pH and high temperature imobilozed Ni complex operates at a similar potential at room temperature and neutral pH condition of hydrogenase enzyme [36]. Active [Ni(P$_2$N$_2$Ph$_2$)$_2$]$^{2+}$ catalyst immobilized on the 2-amino-6-naphthoic acid modified single wall carbon nanotubes (CNTWs). Immobilization results in the activity on a broader range of pH and make it most performing hydrogen-oxidizing electrode till reported [44,45]. Recently Dey et al. [46] have synthesized a [FeFe] H$_2$ase mimics having azadithiolato (ATD) bridge and a terminal alkyn. Using "click reaction" this azide covalently attached to the edge plane graphite surface and reduced graphite oxide surfaces. Modified catalyst shows HER reactive for prolong hours from acidic water.

Conclusion

Side chain interaction is crucial for reversibility. Carboxylic groups give two important advances; structural stability by hindering isomerization and proton relays. The high reactivity of natural enzyme will be well understood by adding amino acids or peptide chains via "top-down" method. Still, these are in early stages of development to achieve actual OCS as that of the natural enzyme. Structural features, temperature, solvent, active site, OCS, the chain lengths of OCS, proton source all collectively make a system whether active or not. There is an enormous scope in future for tuning OCS in such a way that it will give cost-effective, highly durable efficient material for alternate energy sources.

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