Manganese-based Materials Inspired by Photosynthesis for Water-Splitting

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Abstract: In nature, the water-splitting reaction via photosynthesis driven by sunlight in plants, algae, and cyanobacteria stores the vast solar energy and provides vital oxygen to life on earth. The recent advances in elucidating the structures and functions of natural photosynthesis has provided firm framework and solid foundation in applying the knowledge to transform the carbon-based energy to renewable solar energy into our energy systems. In this review, inspired by photosynthesis robust photo water-splitting systems using manganese-containing materials including Mn-terpy dimer/titanium oxide, Mn-oxo tetramer/Nafion, and Mn-terpy oligomer/tungsten oxide, in solar fuel production are summarized and evaluated. Potential problems and future endeavors are also discussed.

Keywords: photosynthesis; manganese; photosystem II; artificial photosynthesis; semiconductor; Nafion

1. Introduction

In nature, oxygenic photosynthesis harvests solar energy from the sun using a range of photosynthetic antenna systems and funnels the excitation energy to the reaction centers, where the light energy is stored and converted into chemical energy and oxygen as a byproduct using by water splitting reaction in plants, algae, and cyanobacteria [1,2]. Recent progresses to understand the natural photosynthesis at the molecular level have been acquired [3,4] and provides a solid basis for unraveling the mechanisms of water splitting and O-O bond formation, which is one of nature’s most fascinating and important chemical reactions. The knowledge inspires and facilitates the creation of artificial photosynthetic model systems [5–10]. A significant amount of research efforts has been placed toward the development of artificial catalytic systems composed of molecular and
supramolecular architectures and discussed in several relevant reviews [4,11–17]. In particular, the synthesis and characterization of Mn-containing compounds such as CaMn₃O₄, CaMn₂O₄, multiple phases of MnO₂, and Mn₃O₄ are fascinating [18–20]. These synthetic models will provide a deeper understanding of the PS II mechanism and complex cluster assembly and open a new window in the design of better molecular catalytic systems for water splitting. Due to the space limitation, the synthesis and characterization of the Mn-containing compounds to mimic the structure of PS II is beyond the scope of this review and will not be discussed.

In this review, manganese-based catalytic systems including Mn-terpy dimer/titanium oxide, Mn-oxo tetramer/Nafion, Mn-terpy oligomer/tungsten oxide, which are highly active in water oxidation catalysis, will be summarized and evaluated in terms of renewable energy production. Potential problems and future endeavors are also presented.

2. Photosynthetic Water Splitting

Photosystem II (PS II) is the site of photosynthetic water oxidation and a large protein complex containing more than 20 subunits with a molecular mass of about 350 kDa. Much of its protein and pigment components control the light harvesting and transferring of photonic energy, and the energy conversion via oxidation of water is carried out at a cluster of metals, Mn₄CaO₅ center, in the oxygen-evolving complex (OEC). When four electron and four protons are extracted from two molecules of water, one molecule of dioxygen is formed. The mechanism of photosynthetic water oxidation in PS II is believed to occur through four distinct oxidation steps as the S-state cycle by Kok et al. and Joliot et al. [21,22].

The structure of PS II has been solved at resolutions of 2.9–3.8 Å from thermophilic cyanobacteria, Thermosynechococcus elongates and Thermosynechococcus vulcanus [23–27]. However, the resolution is not high enough to reveal the structure of Mn cluster and subtract water molecules. A recent report determines the high-resolution structure of PS II at a resolution of 1.9 Å from Thermosynechococcus vulcanus [28]. The structure reveals the complete geometry arrangement of the Mn₄CaO₅ cluster including its oxo bridges and ligands (Figure 1). Of these five metals and five oxygen atoms, three manganese (Mn1 to Mn3), one calcium and four oxygen atoms for a cubane-like structure. Owing to the difference in bond lengths, the Mn₃CaO₄ cubane is not symmetric. The fourth manganese (Mn4) is located outside the cubane and is linked to Mn1 and Mn3. Every two adjacent manganese atoms are linked by di-μ-oxo bridge. The calcium is linked to all four manganese by oxo-bonds, either by d-μ-oxo or mono-μ-oxo bridges.

Four water molecules (W1 to W4) are also identified to be associated with the Mn₄CaO₅ cluster. W1 and W2 are coordinated to Mn4, and W3 and W4 are linked to calcium. No water molecules were found to associate with Mn1, Mn2, and Mn3. All of the amino-acid residues coordinated to the Mn₄CaO₅ cluster were identified and divided into two layers: (1) direct ligands to the cluster include D1-Glu 189, D1-Asp 170, D1-Glu 333, D1-Asp 342, CP43-Gllu 354, D1-His 332, D1-Ala 334; and (2) second coordination sphere including D1-Asp 61, D1-His 337 and CP43-Arg 357. Each of manganese has six ligands whereas the calcium has seven ligands. These pattern and geometric positions of the metal atoms may have important function for the mechanisms of water splitting and O-O bond formation. The important finding of the new Mn₄CaO₅ structure is apparent longer distances
between O5 and metal atoms suggest the corresponding bonds are weak. Therefore, O5 may have a lower negative charge that the normal oxygen atoms and is likely a hydroxide ion in the S1 state. As the W2 and W3 are within the hydrogen-bond distances to O5 and may be the substrate of water oxidation reaction.

**Figure 1.** Structure of the Mn₄CaO₅ cluster and its ligand environment in photosystem II oxygen-evolving complex at a resolution of 1.9 Å. (from reference [28], produced with permission from Macmillan Publisher).

3. Mn-based Catalyst and Their Mechanisms

3.1. Mechanism of Photosynthetic Water Splitting

The four-oxidation steps with five intermediates known as S-states are required for water oxidation; however, there is limited information at the molecular level. Brudvig et al. proposed a possible “molecular” model based on a structural conversion of a Mn₄O₆ “cubane”-like complex [29]. In this model, a tetrameric manganese site in PS II is sequentially oxidized. Each manganese ion in the tetrameric complex is proposed to be coordinated to the protein via N or O ligand to form an octahedral coordination. For the lower oxidation states (S₀-S₂), the manganese complex is in a cubane structure. Upon reaching the S₃ state, the complex may for a Mn₄O₆ adamantane structure. The Dismukes and Christou groups proposed a “butterfly or double pivot mechanism [30,31]. In this model, oxygen is evolved by O-O bond formation across a face of the cube and subsequent loss of O₂ free two of the manganese atoms and open the cube. This idea was supported by gas phase experiments [31].

Figure 2 showed a popular mechanism known as “2 + 2” model proposed by Babcock et al. This model assumes that the structure of Mn center of PS II is a dimer-of-dimer. Each of the two dimers has water ligated in close proximity. The water molecules are progressively oxidized coupled with proton release until a manganese-oxo group formation on one of dimer and production of hydroxide ion on the other in three steps (S₀→S₁→S₂→S₃ transitions). The O-O bond formation with a quick release of O₂ (S₄→S₀) is followed by the removal of the last proton and generation of a peroxo species between
the two dimers (S₃→S₄). Note: the new structure of PS II does not support the structure of “dimer of dimer”. Although the concept of the proposal is valid, the detailed mechanism of PS II water oxidation is unknown and still under debate.

**Figure 2.** Proposed photosynthetic oxygen evolution mechanistic model (from [62], produced with permission from the American Association for the Advancement of Science).

![Proposed photosynthetic oxygen evolution mechanistic model](image)

The key point for the proposals of water oxidation cycle in photosynthesis is the formation of a high-valent metal oxo group, which is produced by the nucleophilic attack by a near water or hydroxide group. Another area of debate is the nature of high-valent manganese-oxo species. Current evidence cannot rule out the possibility of mechanisms involving manganese-oxo or manganese-oxyl species, we refer to this species as a Mn(V)-oxo intermediate.

3.2. Mn-oxo tetramer/Nafion and Mn-oxo tetramer/TiO₂ System

PS II is a very challenging protein to study by biophysical methods. One of the reasons is that PS II OEC is extremely sensitive and vulnerable to environment and experimental conditions. It is well documented that PS II OEC is the target of UV and blue region light radiation [32–36]. The photodamage of PS II is triggered via the direct absorption of a photon by manganese ions in the Mn₄Ca center, as evidenced by the action spectra of PS II photoinhibition and synthetic manganese-oxo dimer compound [34,35]. The UV photoinhibition at the Mn cluster is likely involved the formation of a high-valent manganese species [34,37]. In addition the Mn₄Ca active site in the single crystal of PS II OEC can be damaged by exposure to X-ray radiation revealed by X-ray absorption spectroscopy. [38]

To elucidate the details of the OEC reaction mechanism, bioinorganic chemists have made new complexes to mimic the proposed biological mechanisms. The study of model complexes greatly improved our understanding of the natural water-oxidation mechanism. The design of functional oxygen evolving compounds is successfully achieved. One such example is Mn-oxo tetramer cubic structure developed by Dismukes and co-workers [31]. The key feature of the compound,
[Mn₄O₄(dpp)₆], is its cubical [Mn₄O₄]⁷⁺ core, surrounded by six bridging bidentate chelates to the manganese ions. The water oxidation reaction proceeds with high quantum efficiency (>50%) in the gas phase, and the O₂ product is exclusively produced from the corner oxo’s of the cube. The cubane core facilitates both the selective reduction of two of the four oxygen bridges to water molecules and their photorearrangement to release an O₂ molecule upon photoexcitation at 350 nm. The highest oxidation state observed in the cubane family contains the [Mn₄O₄]⁷⁺ core, or [Mn₄O₂L₆]⁺.

Due to the insolubility of cubium in water and most organic solvents, Mn-oxo tetramer/Nafion system was developed by depositing the Mn-oxo tetramer into a thin Nafion membrane on a conducting electrode. As shown in Figure 3, the system is a robust catalyst that sustains the photoelectrooxidation of water to O₂ [39]. The perfluorinated polymer Nafion comprises hydrophobic domains and provides excellent aqueous channels for proton transfer in water oxidation reaction. The photooxidation of water is likely due to a catalytically active species originating from the photolysis of Mn-oxo tetramer which is bonded to or activated by the sulfonate groups of the aqueous Nafion membrane. The authors postulate that the Mn-oxo tetramer/Nafion follows the similar photoreaction mechanism of Mn-tetramer in gas phase with O₂ being released upon photodissociation to yield the “reduced butterfly species” [Mn₄O₂L₅]²⁺. This species may be more stable in restricted hydrophobic pockets of Nafion.

**Figure 3.** Mn-tetramer/Nafion water splitting catalytic system. Photoanode are made of a titania layer sensitized with the dye, [Ru(bipy)₂(bipy(COO)₂)], and a Nafion film doped with the Mn-oxo tetramer, [Mn₄O₂L₆]⁺. (From reference [9], produced with permission from American Chemical Society).

Recent *in situ* X-ray absorption spectroscopy and transmission electron microscopy revealed that the synthetic Mn-oxo tetramer cluster undergoes a dissociation to form Mn(II) compound in the Nafion matrix [40]. The Mn(II) species can be reoxidized to generate the catalytic Mn(III/V) oxide and close the water oxidation catalytic cycle. The original Mn-oxo tetramer is likely serves only as a precursor to the catalytically active material.

### 3.3. Mn-terpy dimer and Mn-terpy dimer/TiO₂ System

Another excellent example is the first PS II functional model for oxygen evolving complex so called “Mn-terpy dimer,” ([OH₂(terpy)Mn(O)₂Mn(terpy)OH₂](NO₃)₃·6H₂O) [41]. Figure 4 showed a proposed mechanism for the formation of the reactive intermediate in oxygen evolution catalytically by the Mn-terpy dimer in the present of chemical oxidants, XO, such as hypochlorite or
peroxymonsulfate (Oxone). The XO binds initially to the Mn-terpy dimer reversibly (step 1 → 3). Once formed, compound 3 can react with XO to form the Mn(V)=O intermediate species (step 3 → 4). This is the rate-limiting step in oxygen evolution. The O-O bond-forming step involves attack of solvent or Oxone on the oxo ligand [42].

**Figure 4.** Oxygen evolution mechanism of Mn-terpy dimer in the presence of chemical oxidant, XO. Left pathway: attach of XO on the terminal oxo ligand. Right pathway: attack of solvent water on the terminal oxo ligand. (from reference [42], produced with permission from the American Chemical Society).

By mimicking the photoinduced charge separation of natural photosynthesis, molecular and supramolecular model systems can provide important inspiration for practical solar fuel production. Hammarstrom, Styring, and co-workers prepared molecular assemblies coupling Mn complexes with the dye photosensitizer, [Ru(bpy)₃]²⁺, and observed light-induced manganese oxidation and long-lived charge separation [43]. Brudvig et al. have directly deposited the Mn-terpy dimer on the TiO₂ nanoparticles (NP) and found that Mn-terpy dimer binds to near amorphous TiO₂ NP and dimerized to form Mn(IV) tetramer. The Mn(IV) tetramer attached to TiO₂ oxidizes water to oxygen in the presence of Ce⁴⁺ as the primary one-electron oxidant [44–46]. They also [47–49] synthesize a derivative of Mn-terpy dimer on the surface of TiO₂ NPs in situ by first attached the precursor Mn(II) monomers through an anchoring ligand on the TiO₂ NPs and then assembling the Mn-terpy dimer [50]. The resulting Mn-terpy dimer/TiO₂ is able to reversibly change mixed valent Mn(III/IV) to Mn(IV/IV) state by photoexcitation and interfacial electron injection into the conduction band of TiO₂.

### 3.4. Mn-oligomer/WO₃ System

The weakness of Mn-terpy dimer is that the compound is unstable under elevated temperature [51]. The decomposition of the Mn-terpy dimer in aqueous solution over 60 °C occurred involving a change in Mn valence. Interestingly, the thermal decomposition of Mn-terpy dimer was found to generate a Mn-containing precipitate that retained catalytic oxygen-evolving activity [51]. The solid Mn-containing material binds terpy ligand, judged by elemental analysis and atomic absorption analysis, and is not
simple manganese dioxide. The novel Mn-containing precipitate, assigned as a Mn-terpy oligomer, is thermal stable and may be a unique option for constructing catalytic systems in solar fuel production.

When heated over 60 °C, the Mn-terpy oligomer undergoes thermal decomposition and yields a controlled thin layer on a semiconductor such as tungsten oxide. The resulting Mn-oligomer/WO₃ material is able to efficiently split water to generate oxygen using GC and GC/MS under light illumination [6]. The WO₃ material alone is unstable over a long period of time of water splitting. At pH 7.0, WO₃ without Mn-terpy oligomer decayed about 60% within 1 h, whereas approximately 4% performance decrease was observed when the Mn-terpy was present for up to 2 h. In addition to O₂, the amount of H₂ generated by the photoelectrochemical setup was also measured. The 2:1 ratio of H₂ and O₂ generation rate confirms the complete decomposition of H₂O by the photogenerated charges unambiguously. Another piece of evidence is the ¹⁸O isotope experiments. H₂¹⁸O was used in water splitting experiment. A significant amount of ¹⁶,¹⁸O₂ in the headspace gas sample was detected in addition to a small amount of ¹⁸,¹⁸O₂.

Based on the observed experimental data, a possible mechanism of Mn-terpy oligomer/WO₃ system in water oxidation is proposed in Figure 5. Four light photons are required to oxidize the Mn-terpy species accompanying four proton-coupled steps. At each step, the photon causes charge separation in WO₃. The hole generated in WO₃ receives electron from Mn-terpy complex and oxidizes the Mn ion via Mn valence changes. The Mn(V) intermediate species is generated by the fourth photon-driven electron transfer reaction and splits water to dioxygen and regenerate active Mn-terpy catalyst.

**Figure 5.** Proposed mechanisms of Mn-terpy oligomer/WO₃ catalytic system in water oxidation. Black arrows indicate the proton transfer reaction; blue arrows represent the electron transfer steps. Mn-terpy oligomer oxidizes H₂O oxidation by receiving photogenerated holes from WO₃ and undergoing a catalytic cycle, in which Mn valence change is a proton-coupled electron transfer in a similar way to PS II oxygen-evolving complex.
4. Concluding Remarks

In the recent five years, substantial progress in natural photosynthesis, especially protein structural and mechanistic aspects of PS II oxygen-evolving complex, has inspired the revolutionary development in photoelectrochemical and electrochemical water splitting using manganese-based molecular catalysts as well as Fe-, Co-, Ni-, Ru-, and Ir-containing compounds associated with dye-sensitized semiconductors in solar fuel production [52–58]. Another highly desirable use for solar energy is powering fuel generation by hydrogen production via water splitting using visible light [59,60]. These catalytic systems mimic the natural photosynthesis and take the advantages of multiple materials with unique properties in terms of light harvesting, energy transfer, charge separation, electron transfer, water oxidation, and hydrogen production. It is highly likely that the progress in the field of nanomaterial and photosynthesis will offer novel technology for transforming the solar fuels into our future sustainable energy sources in the next 10–20 years, hopefully as eventual personalized energy systems [61]. Grand challenges remain, including the discovery of inexpensive, robust, and efficient water-oxidation catalysts. It is worth noting that nanoscale materials are especially appealing in energy research because their dimensions coincide with the length scales of processes fundamental to energy conversion and storage. Future endeavors will be placed on improvement in efficiency and durability using visible light as well as the development of novel catalytic materials involving dye-sensitized solar cells and photoelectrochemical cells.

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