PERSPECTIVE
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Water oxidation using earth-abundant transition metal catalysts: opportunities and challenges
1. Introduction

Our society’s growing energy demands and the impact of rising CO₂ emissions have resulted in the awareness of the necessity of developing sustainable and carbon-neutral alternatives to fossil fuels.¹–³ Of the possible sustainable energy sources, only solar energy has the potential to meet the global energy demands. In this context, an attractive way of harnessing solar energy is artificial photosynthesis, where solar energy is employed to split H₂O into O₂ and a fuel, H₂ (eqn (1)), thus storing the energy provided by the sun in chemical bonds. Therefore, methods for converting and storing solar energy are being intensively pursued.⁴–⁶

$$2\text{H}_2\text{O} \rightarrow \text{O}_2 + 2\text{H}_2 \quad (1)$$

Owing to its natural abundance, H₂O would be the ideal source of reducing equivalents, i.e. protons and electrons. However, the development of efficient catalysts for the oxidation of H₂O remains a bottleneck in the development of technologies for the conversion of solar energy into fuels.⁷ The natural photosynthesis provides a brilliant strategy for harnessing sunlight and using it to oxidize H₂O to O₂ in order to generate the necessary reducing equivalents for producing biomass by reducing CO₂ to carbohydrates. This process, which takes place in the photosynthetic machinery of green plants and cyanobacteria, is the basis for a major part of our global energy supplies and is thus a crucial reaction in order to sustain life on earth.⁸,⁹

1.1. Photosynthesis—light-harvesting for the oxidation of H₂O

The initial step in the photosynthetic machinery involves light absorption aided by photosynthetic pigments. The energy is subsequently funneled to the central P₆₈₀ reaction center, resulting in charge separation to give the oxidized P₆₈₀ (P₆₈₀⁰). Reduction of P₆₈₀⁰ occurs by abstraction of an electron from the Mn₃Ca cluster located in the oxygen-evolving complex (OEC). After four consecutive electron transfers, the Mn₃Ca cluster is regenerated by formally oxidizing two molecules of H₂O to O₂ (eqn (2)).¹⁰,¹¹

$$2\text{H}_2\text{O} \rightarrow \text{O}_2 + 4\text{H}^+ + 4\text{e}^- \quad (2)$$

X-ray crystallography has revealed that the OEC is composed of a cuboidal Mn₃Ca core with a dangling Mn center (Fig. 1).¹²–¹⁵ The mechanism by which this Mn cluster catalyzes the oxidation of H₂O is currently fairly well understood. However, some questions still remain unsolved and it is therefore still actively studied with the view that further insight into the mechanism is important for the design of artificial systems for solar energy to fuel conversion.¹⁶–²²

1.2. Artificial photosynthesis

In contrast to natural photosynthesis where the protons and electrons produced from H₂O oxidation are used to convert CO₂ to biomass, in artificial systems they are converted into simpler chemical fuels, such as H₂. The light-driven H₂O splitting requires the orchestration of several intricate processes, such as light absorption, charge separation, oxidation of H₂O and reduction of the generated protons.⁴ In artificial photosynthetic systems, catalysts for the oxidation of H₂O are currently the bottleneck. To catalyze such an energy demanding reaction...
as H₂O oxidation, the catalysts should be able to accumulate several oxidizing equivalents, forming the essential O–O bond, and operate close to the thermodynamic potential. The design of artificial water oxidation catalysts (WOCs) thus represents a fundamental challenge in schemes for artificial photosynthesis and has therefore been the major focus of research during the past few decades.

The majority of homogeneous WOCs are based on the second- and third-row transition metals Ru and Ir. Complexes based on these metals have been shown to produce robust and highly active WOCs, and are, in general, thought to generate the vital O–O bond either via the water nucleophilic attack (WNA) mechanism or the interaction of two metal–oxo units (I2M) (see Fig. 2).

If water splitting is to become economically viable on a global scale, the high cost and low abundance associated with transition metals, such as Ru and Ir, may become a problem. The construction of WOCs based on earth-abundant first-row transition metals is therefore highly desirable. Despite considerable efforts, the development of robust WOCs incorporating first-row elements that operate under mild conditions has proven particularly challenging. However, if successful,

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this could have a widespread impact on the construction of low-cost devices for solar energy to fuel conversion. This Perspective summarizes recent advances in the design of WOCs based on earth-abundant first-row transition metals with an emphasis on homogeneous systems based on Mn, Fe, Co and Cu. An examination of the advantages and current limitations of the highlighted catalytic systems is also presented, along with potential future directions of this expanding research area.

2. Methods for evaluating water oxidation catalysts

There exist essentially three different methods for evaluating whether a complex is capable of oxidizing H₂O. The first method is chemical oxidation and involves the use of sacrificial oxidants, allowing for rapid screening of potential WOCs. The most widely used one-electron oxidant is ceric ammonium nitrate (CAN, Ce(NH₄)₂(NO₃)₆, CeIV), which has a redox potential of ∼1.70 V vs. NHE (normal hydrogen electrode). However, the O₂ studies with CeIV (eqn (3)) need to be performed in acidic aqueous media, due to the instability of CeIV at higher pH, and this limits the evaluation to homogeneous catalysts that are resistant to decomposition under acidic conditions. A number of two-electron oxidants have also been used for the evaluation of WOCs, including potassium peroxymonosulfate (Oxone®) and sodium periodate (NaIO₄), which have the formation of WOCs, including potassium peroxymono-

$$2\text{H}_2\text{O} + 4\text{Ce}^{IV} \rightarrow \text{O}_2 + 4\text{H}^+ + 4\text{Ce}^{III} \quad (3)$$

A number of two-electron oxidants have also been used for the evaluation of WOCs, including potassium peroxymonosulfate (Oxone®) and sodium periodate (NaIO₄), which have redox potentials of 1.82 V and 1.60 V vs. NHE, respectively. The main advantage with these oxidants is that they can be used under neutral conditions, thus allowing the examination of acid labile catalysts. The two-electron nature of these oxidants limits their relevance to one-electron conditions, which might occur in photoelectrochemical cells. Oxone® and NaIO₄ are also known oxo-transfer reagents, which precludes that both oxygen atoms in the generated O₂ originate from H₂O; however, this can be easily determined by mass spectrometry using ¹⁸O isotopically labeled H₂O (H₂¹⁸O). It should be noted that recent studies on CeIV suggest that this oxidant might also function as an oxo-transfer reagent.

An alternative to CeIV is the one-electron oxidant [Ru(bpy)₃]³⁺ (I⁺, Fig. 3; bpy = 2,2'-bipyridine), which has a redox potential of 1.26 V vs. NHE. Although this and related complexes have not been widely used as oxidants (eqn (4)), perhaps due to their high cost, the advantages of these oxidants are their activity under neutral conditions and the fact that they can be photochemically generated from the corresponding [Ru(bpy)₃]²⁺-type complexes. The photochemistry of these Ru poly(pyridyl) complexes has been extensively studied and their photophysical properties are well documented.

$$\text{2H}_2\text{O} + 4[Ru(bpy)_3]^{3+} \rightarrow \text{O}_2 + 4\text{H}^+ + 4[Ru(bpy)_3]^{2+} \quad (4)$$

In combination with a sacrificial electron acceptor, [Ru(bpy)₃]²⁺ can be used for light-driven H₂O oxidation. Visible-light absorption of the [Ru(bpy)₃]²⁺ photosensitizer results in the formation of the singlet excited state ([Ru(bpy)₃]²⁺*). This undergoes rapid intersystem crossing (ISC) and is converted into an excited triplet state ([Ru(bpy)₃]³⁺*). The presence of a sacrificial electron acceptor, such as sodium persulfate (Na₂S₂O₈), [Ru(bpy)₃]³⁺* is oxidatively quenched, which generates [Ru(bpy)₃]³⁺, a sulfate ion (SO₄²⁻) and a sulfate radical (SO₄⁻). The sulfate radical is a strong oxidant (E = ∼2.4 V vs. NHE) and has the ability to directly oxidize another equivalent of [Ru(bpy)₃]²⁺ to [Ru(bpy)₃]³⁺. The overall light-induced process is summarized in eqn (5) and Fig. 4.

$$2\nu + 4[Ru(bpy)_3]^{2+} + 2\text{S}_2\text{O}_8^{2-} \rightarrow 4[Ru(bpy)_3]^{3+} + 4\text{SO}_4^{2-} \quad (5)$$

In addition to the aforementioned (photo)chemical techniques for studying WOCs, (photo)electrochemical methods are also available. Here, an important parameter is the so-called overpotential, which is the potential that has to be applied in an electrolysis cell in addition to the thermodynamical potential for a given half-reaction. These electrochemical studies have a major advantage, namely that the studied system resembles the conditions that would be employed in the envisioned formation of solar fuels.
3. Distinguishing between heterogeneous and homogeneous species

Earth-abundant WOCs are especially prone to undergo irreversible structural modifications, leading to transformation of the initially homogeneous complex into heterogeneous metal oxide species. The designed metal complexes merely act as precursors for the in situ formation of metal nanoparticles that might be the true active catalytic species. An essential element for researchers working in the field is thus to determine whether the observed activity is derived from the pristine metal complex or whether nanoparticles, produced during the oxidation process, are responsible for converting H2O to O2. Several distinct physical techniques exist for establishing whether the starting well-defined molecular complex is the true catalytic species or merely a precursor and include X-ray photoelectron spectroscopy (XPS), energy-dispersive X-ray spectroscopy (EDX), transmission electron microscopy (TEM), and dynamic light scattering (DLS). In combination, these techniques provide a powerful toolbox for assessing the real catalytic entity. One also needs to consider that a slight change in the reaction conditions may have a great effect on both the stability of WOCs and the operating mechanism.50,51

4. Artificial Mn complexes as water oxidation catalysts

4.1. Dinuclear Mn complexes containing multidentate ligand scaffolds

Inspired by the Mn4Ca cluster located in the OEC, initial attempts to develop earth-abundant WOCs focused on Mn.52 One of the seminal examples of dinuclear Mn complexes capable of oxidizing H2O is the Mn porphyrin dimers 2–4 (Fig. 5) developed by Naruta and co-workers.53,54 In aqueous MeCN solutions containing nBu4NOH, all three Mn dimers showed a similar irreversible increase in current at potentials >1.4 V vs. NHE. Of the three studied Mn porphyrin dimers, complex 4 showed the highest activity, affording a TON of 9.2. The O2 evolution rate was also found to be proportional to the concentration of the Mn dimers. The analogous monomeric Mn porphyrin complexes did not promote O2 evolution and in the absence of the Mn dimers the onset potential was observed at >2.5 V, highlighting the necessity for the dinuclear Mn scaffold.54 Due to the proximity of the Mn centers, the authors proposed that O–O bond formation occurred through nucleophilic attack of OH− on a dimeric MnV-O species, thus generating a MnIV-OO-MnIV species from which O2 could subsequently be liberated.54 A sulfonated Mn porphyrin complex has also been incorporated into a thin layer of poly(terthiophene) (PTTh) for attachment to indium tin oxide (ITO) electrodes. The resulting electrodes were capable of catalyzing photoelectrochemical H2O oxidation with low overpotentials.55

Perhaps the most well-studied Mn complex capable of mediating H2O oxidation is the dimeric [[tpy(OH2)Mn(μ-O)2Mn(OH2)(tpy)]3+] complex (5, Fig. 6; tpy = 2,2′:6′,2″-terpyridine) reported by Crabtree’s and Brudvig’s groups.35,56,57 The crystal structure of the dinuclear Mn complex 5 revealed that it consisted of a mixed valence dimer with one of the Mn centers being in the +III state and the other in the +IV state. The two exchangeable aqua ligands are essential for the catalytic activity of the Mn complex. The initial evaluation of the oxidation capability of the complex was carried out with NaOCl as a chemical oxidant. Using this oxidant at pH 8.6, the dimeric Mn complex 5 afforded a TON of 4 and a TOF of 0.0033 s−1. The low efficiency of the Mn dimer was ascribed to the ligand decomposition, resulting in the formation of catalytically active MnO4−, as monitored by UV-vis spectroscopy.35 Studies have also suggested that H2O oxidation with Mn complex 5 can be driven using Oxone®58,59 or CeIV (ref. 60 and 61) as a
chemical oxidant. Deposition of the dimeric Mn complex 5 onto different materials, such as kaolin,\textsuperscript{62} mica,\textsuperscript{63-65} montmorillonite\textsuperscript{66} or TiO\textsubscript{2},\textsuperscript{67,68} has also been carried out. Additionally, attempts to extend the lifetime of Mn complex 5 by its incorporation into metal–organic frameworks (MOFs) have been reported.\textsuperscript{69,70} A variety of analogs of Mn complex 5 have been synthesized and evaluated as WOCs.\textsuperscript{71-75}

Based on UV-vis spectroscopy and electron paramagnetic resonance (EPR) experiments, the authors proposed a simplified mechanism, depicted in Scheme 1, for O\textsubscript{2} formation from the reaction of Mn complex 5 with NaClO. The proposed mechanism involves an initial oxidation of dimeric Mn complex 5 to produce a Mn\textsuperscript{IV,IV}\textsubscript{2} species (6), which can react with the oxidant to generate Mn\textsuperscript{V=O} (7). This species is the key intermediate and is responsible for promoting O–O bond formation via nucleophilic attack of OH\textsuperscript{−}. The formed Mn–peroxo intermediate is subsequently oxidized, which results in the release of O\textsubscript{2} and ultimately regenerates the dimeric Mn\textsuperscript{V,IV} complex 6.\textsuperscript{35,76} It can be concluded that the mechanism by which Mn complex 5 mediates the oxidation of H\textsubscript{2}O is intricate and is highly dependent on the reaction conditions.\textsuperscript{77,78} This is further supported by findings concerning the possible involvement of a tetrameric Mn species\textsuperscript{79-81} while another report suggests that heterogeneous Mn–oxides are generated.\textsuperscript{82}

Based on previous findings,\textsuperscript{83} McKenzie and co-workers evaluated the dimeric Mn complex [Mn\textsuperscript{II,II}\textsubscript{2}(mcbpen)\textsubscript{2}(OH\textsubscript{2})\textsubscript{2}]\textsuperscript{2+} (9; mcbpen = N-methyl-N′-carboxymethyl-N,N′-bis(2-pyridylmethyl)ethane-1,2-diamine) as a WOC (Fig. 7). The dimeric Mn complex 9 was able to evolve O\textsubscript{2} when driven using t\textsubscript{BuOOH or Ce\textsuperscript{IV} as a chemical oxidant.\textsuperscript{84} Using t\textsubscript{BuOOH as an oxidant afforded high turnovers of up to 15 000 mol of O\textsubscript{2} per
mol of Mn. However, when using isotopically labeled H$_2$O (H$_2^{18}$O), the evolved O$_2$ was shown to be exclusively $^{18}$O$_2$. This highlights that one of the oxygen atoms in the generated O$_2$ originates from the $^t$BuOOH oxidant, demonstrating the ability of this two-electron oxidant to act as an oxygen atom transfer reagent. In the case when Ce$^{IV}$ is used as an oxidant, the only available source of oxygen is the nitrate counterion of the oxidant, thus suggesting the active involvement of the nitrate counterion in the catalytic cycle.

The mcbpen ligand scaffold with its carboxylate and pyridine motifs attached to the ethylenediamine backbone was suggested to provide an ideal environment for coordination to Mn in various redox states. It was initially proposed that the catalytic cycle for O$_2$ generation involved the coupling of two μ-oxo units. However, a subsequent computational study suggested that the key species consisted of a Mn$^{III}(μ$-O)Mn$^{IV}$ – O’ species (12), a mixed valent oxyl radical species, which is in equilibrium with the isomeric diamond-core Mn$^{IV}(μ$-O)$_2$Mn$^{IV}$ (13). This mixed valent Mn$^{III}(μ$-O)Mn$^{IV}$–O’ species (12) is responsible for mediating O–O bond formation through nucleophilic attack of H$_2$O, resulting in a Mn$^{III}$–OOH species (14). The other Mn center, which has thus far acted as a spectator, functions as the additional oxidizing equivalent, liberating O$_2$ to regenerate the monomeric Mn$^{II}$ complex 10 without the need for an additional oxidation event (Scheme 2).

Several Mn complexes have been covalently linked to [Ru(bpy)$_3$]$^{2+}$-type chromophores for mimicking the electron transfer events occurring in Photosystem II. Light absorption of the Ru$^{II}$ photosensitizer core triggers electron transfer from the photoexcited state (Ru$^{II*}$) to a sacrificial electron transfer. The oxidized photosensitizer subsequently abstracts an electron from the attached Mn complex. Of the examined Ru–Mn assemblies, assembly 16 (Fig. 8) displayed promising properties as it was able to undergo three consecutive electron transfers, which converted the initial Mn$^{II,II}_2$ core to Mn$^{III,IV}_2$. Although Ru–Mn$_2$ assembly could accumulate three oxidizing equivalents, it was not able to evolve O$_2$ under the studied reaction conditions.

In light of the attractive features of the dinuclear Mn core in assembly 16, the properties and O$_2$ evolving activities of several different dinuclear Mn complexes (17–20) housing ligand scaffolds based on benzylic amines have been studied (Fig. 9). It could be concluded that the one-electron oxidants Ce$^{IV}$ and [Ru(bpy)$_3$]$^{3+}$ were inefficient oxidants in H$_2$O oxidation. However, with two-electron oxidants, such as H$_2$O$_2$, $^t$BuOOH, Oxone® or NaClO, i.e. oxygen-transfer oxidants, a more efficient O$_2$ evolution was observed.

The fact that imidazole, carboxylate and phenol moieties are essential features of the OEC inspired the group of Åkermark to design the dinuclear Mn complex 21 (Fig. 10, left)
housing a bioinspired ligand scaffold with imidazole, carboxylate and phenolic groups.\(^9\) In the solid state, the crystal structure revealed that Mn complex 21 adopts a tetrameric structure (22, Fig. 10, right), resembling the tetranuclear Mn\(\text{IV,III}\) core in the OEC. An additional advantage of the incorporation of the negatively charged groups was the lowering of the redox potentials, an effect previously observed for other metal complexes,\(^{100,101}\) allowing for H\(_2\)O oxidation to be driven by [Ru(bpy)\(_3\)]\(^3+\). Experiments were initially carried out with a 480-fold excess of the chemical oxidant [Ru(bpy)\(_3\)]\(^3+\) under neutral conditions. Upon addition of an aqueous solution containing Mn complex 21, O\(_2\) evolution was immediately observed, affording a TON of \(~25\) and a TOF of \(~0.027\) s\(^{-1}\). Light-driven H\(_2\)O oxidation was also carried out using [Ru(bpy)\(_3\)]\(^2+\)-type photosensitizers and Na\(_2\)S\(_2\)O\(_8\) as a sacrificial electron acceptor. When employing [Ru(bpy)\(_2\)(deeb)]\(^2+\) as a photosensitizer (deeb = diethyl 2,2′-bipyridine-4,4′-dicarboxylate), a TON of 4 was reached. Experiments with isotopically labeled H\(_2\)O (H\(_2\)\(_{18}\)O) confirmed that both of the oxygen atoms in the produced O\(_2\) originated from the solvent H\(_2\)O.\(^9\)

Subsequent work involved the preparation of a Ru–Mn\(_2\) assembly (23, Fig. 11) where the dinuclear Mn\(_{\text{IV,III}}\) complex 21 was covalently linked to a [Ru(bpy)\(_3\)]\(^2+\)-type photosensitizer.\(^{102}\) Although Ru–Mn\(_2\) assembly 23 was able to mediate chemical

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Fig. 8 Ru–Mn\(_2\) assembly 16 capable of accumulating three oxidizing equivalents. A = acceptor.

Fig. 9 Structures of Mn complexes 17–20.
oxidation of H$_2$O, the photophysical properties of the assembly did not allow for photochemical H$_2$O oxidation in the presence of a sacrificial electron acceptor (Na$_2$S$_2$O$_8$). It was revealed that the excited state of the dyad 23 was too short-lived and could not be modelled with a mono-exponential decay function. Computational studies suggested the presence of an intricate excited-state manifold, along with possible large effects of different protonation states, which may have contributed to the observed intricate decay pattern.

A series of dinuclear derivatives of Mn complex 21 have also been synthesized.$^{103}$ The prepared ligand scaffolds contained a variety of different substituents and the electronic and catalytic features of the corresponding Mn complexes were systematically examined. Of the synthesized Mn complexes, complex 24 possessing the distal carboxylate group (Fig. 12, left) was shown to produce O$_2$ more efficiently than the other Mn complexes. This effect was proposed to originate from pre-orientation of the incoming H$_2$O molecule where the introduced carboxylate moiety directs the incoming H$_2$O through hydrogen bonding (Fig. 12, right). This probably facilitates proton-coupled electron transfer (PCET) and assists the nucleophilic attack on a high-valent Mn–oxyl/Mn–oxo species. Incorporation of non-innocent distal groups into ligand backbones could be a general method for enhancing the catalytic activity of WOCs.

A computational study has been performed on the mechanism by which the dinuclear Mn$_{II,III}$ complex 21 mediates O–O bond formation.$^{104}$ The study suggested an OEC-like mechanism where tetrameric Mn species are involved throughout the catalytic cycle (Fig. 13). The catalytic cycle was suggested to begin with the formal Mn$_{III,III,III,IV}$ species, which is better described as having all four Mn centers in the +III state with a ligand radical. Four PCET events transform the initial formal Mn$_{III,III,III,IV}$ species into a Mn$_{III,IV,IV,IV}$–L–L$^\ast$ species. Intramolecular proton transfer subsequently generates a Mn$_{IV}$-bound oxyl radical that is responsible for creating the O–O bond with the Mn1-Mn2 bridging oxo group (Fig. 13). The proposed mechanistic pathway contains several features that are important clues to understand how the OEC catalyzes H$_2$O oxidation.$^{19}$ Examples of design principles that were obtained from the study are: (1) a Mn-bound terminal aqua molecule is essential for the formation of the crucial Mn$_{IV}$–O$^\ast$ species, (2) nucleophilic attack on the Mn$_{IV}$–O$^\ast$ species is less favorable compared to a route featuring the coupling between this species and a bridging oxo moiety, and (3) the presence of a redox-active ligand backbone decreases the redox potentials for the individual redox events and alleviates the metal centers from being too heavily oxidized.$^{104}$
4.2. Mononuclear Mn complexes: O–O bond formation at single metal centers

A vital feature for metal complexes to mediate H₂O oxidation is the necessity for these catalysts to generate high-valent metal-oxyl/oxo intermediates, which subsequently trigger O–O bond formation. Several formal high-valent Mn–oxo complexes have been prepared using various chemical oxidants, such as hydrogen peroxide (H₂O₂),¹⁰⁵ iodosylbenzene (PhIO),¹⁰⁶ or m-chloroperbenzoic acid (m-CPBA),¹⁰⁷ or photochemically generated [Ru(bpy)₃]⁺.¹⁰⁸

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Fig. 12 (Left) Molecular structure of dinuclear MnII,III₂ complex 24 and (right) calculated structure of the complex in its formal MnIV,V₂ state, showing the hydrogen-bonding interaction between the distal carboxylate moiety and the Mn bound hydroxide(s) (adapted with permission from ref. 103. Published by the PCCP Owner Societies).

Fig. 13 Comparison of the catalytic cycles and transition-states for O–O bond formation for the tetranuclear Mn complex derived from Mn complex 21 and the tetranuclear Mn₄Ca cluster in the OEC. Adapted with permission from ref. 104. Copyright 2015 American Chemical Society.
Although a variety of high-valent Mn–oxo complexes have been reported, only a few mononuclear Mn complexes have been prepared to mediate \( H_2O \) oxidation. An early example of a mononuclear Mn complex capable of evolving \( O_2 \) is the Mn Schiff base complex [Mn(salpd)(OH)_2]⁺ (25, Fig. 14; H_salpd = \( \text{N,N'}\)-bis(salicylidene)propane-1,3-diamine). This Mn\(^{III}\) Schiff base complex was reported to liberate \( O_2 \) and reduce \( p \)-benzoquinone to hydroquinone when irradiated with visible-light. The catalytic activity was shown to be dependent on the quinone concentration, the wavelength of light, temperature and \( \text{pH} \), affording TONs of 0.02–0.06 depending on the reaction conditions. Although the exact mechanism has not been elucidated, the rate of \( O_2 \) evolution followed first-order kinetics with respect to Mn complex 25 and \([\text{Mn(salpd)}]_2\text{O}\) was identified as a deposit during irradiation. Related Mn Schiff base complexes have also been studied for \( O_2 \) evolution by several research groups. An interesting study was conducted by the group of Åkermark and Sun where the mononuclear Mn\(^{IV}\) corrole xanthene complex 26 (Fig. 15) was shown to promote electrochemical \( H_2O \) oxidation. In aqueous solutions in the presence of \( nBu_4NOH \), \( O_2 \) was detected at quite low oxidation potentials. Electrochemical measurements of Mn corrole complex 26 further revealed that high-valent redox states were accessed at relatively low potentials. A computational study suggested that O–O bond formation occurred via nucleophilic attack of \( OH^- \) or \( H_2O \) on a formal Mn\(^{IV}=\text{O} \) intermediate.

In attempts to obtain experimental support for a mechanistic pathway involving nucleophilic addition of \( OH^- \) or \( H_2O \), a subsequent study focused on Mn\(^{III}\) corrole 27 (Fig. 16). The Mn corrole complex 27 was chosen as it was believed to produce a sufficiently stable Mn\(^{V} \) species (28). Treating Mn complex 27 with \( \text{BuOOH} \) furnished the corresponding Mn\(^{V} \) species, as shown by UV–vis spectroscopy. Additional support for the generation of the Mn\(^{V}=\text{O} \) species 28 was obtained from high-resolution mass spectrometry (HRMS). Upon addition of 2 equivalents of \( nBu_4OH \), rapid \( O_2 \) evolution ensued and a peak ascribed to a Mn\(^{IV} \) species (30) appeared, which is most likely generated from oxidation of the Mn\(^{III} \)-hydroperoxo species 29. This peak gradually decreased with the emergence of a peak corresponding to the Mn\(^{III}=\text{OH} \) complex 31. The use of isotopically labeled \( H_2O \) (\( H_2^{18}O \)) supported that the evolved \( O_2 \) is produced by addition of \( ^{18}O \) to the unlabeled Mn\(^{V}=\text{O} \) species. This study demonstrates that nucleophilic attack of hydroxide on a high-valent Mn species is a viable mechanistic scenario for triggering \( O=O \) bond formation in the conversion of \( H_2O \) to \( O_2 \) (Scheme 3).

The group of Smith found that small changes in the ligand backbone of Mn\(^{II}\) pyridinophane complexes (Fig. 17) had a dramatic influence on the catalytic properties of these complexes. Mn complexes 32 and 33, possessing small substituents, were shown to disproportionate \( H_2O \) in aqueous solutions. However, for Mn complex 34 with the sterically encumbered \( tBu \) group, catalase activity is turned off and the complex instead mediates electrocatalytic \( H_2O \) oxidation. When carrying out the electrocatalytic experiments at \( pH \) 12.2 with a potential of 1.23 \( V \) vs. \( \text{NHE} \), TONs of 16–24 were obtained with Faradaic efficiencies of 74–81%. A number of experiments were also undertaken to exclude the involvement of heterogeneous nanoparticles: (1) no spectral changes were observed for the electrolysis solutions, (2) DLS measurements provided no evidence for nanoparticles, (3) the catalytic current does not increase over successive scans, which would be expected if a catalytic deposit was produced, and (4) EDX measurements showed no evidence of the formation of Mn-containing deposits on the electrode. Collectively, these results point to a homogeneous pathway for the oxidation of \( H_2O \). Mechanistic studies revealed that the catalytic current varies
linearly with the concentration of Mn complex 34, indicating that O–O bond formation occurs at a single metal center.\(^{121}\) The results highlight that a ligand-controlled switch in catalytic reactivity has implications for the design of novel Mn-based WOCs.

Brudvig and co-workers recently investigated the \(\text{H}_2\text{O}\) oxidation ability of the three mononuclear Mn complexes depicted in Fig. 18; \([\text{Mn(PaPy})_3(\text{NO}_3)]^{+}\) (35; \(\text{HPaPy}_3^{123} = \text{N},\text{N}-\text{bis}(2\text{-pyridylmethyl})\text{-amine-N-ethyl-2-pyridine-2-carboxamide})\), \([\text{Mn(N4Py)}(\text{OTf})]^+\) (36; \(\text{N4Py}^{124,125} = \text{N},\text{N}-\text{bis}(2\text{-pyridyl-methyl})\text{-N-bis(2-pyridyl)methylamine}\)) and \([\text{Mn(PY5)}(\text{OH}_2)]^{2+}\) (37; \(\text{PY5}^{126,127} = 2,6\text{-bis(methoxydi(pyridin-2-yl)methyl)pyridine})^{128}\)

To facilitate access to the key high-valent Mn–oxo species required to initiate O–O bond formation, an electron-rich ligand scaffold is essential. The PaPy\(^3\) ligand was chosen as it has been shown that deprotonated carboxamido units are strong electron donors and have previously been employed for isolation of high-valent Mn\(^{V=O}\) compounds.\(^{129–131}\) It was envisioned that the inclusion of the carboxamido unit at the trans position to the oxo site could favor a reactive species by weakening the Mn–oxo bond. Catalytic experiments revealed that Mn complex \([\text{Mn(PaPy})_3(\text{NO}_3)]^{+}\) (35) was able to catalyze \(\text{O}_2\) formation using either Oxone\(^\circ\) or \(\text{H}_2\text{O}_2\) as a two-electron oxidant and kinetic studies showed that the reaction was first-order in the catalyst. However, Mn complex \([\text{Mn(N4Py)}(\text{OTf})]^+\) (36) was shown to evolve \(\text{O}_2\) only in the presence of Oxone\(^\circ\) while Mn complex \([\text{Mn(PY5)}(\text{OH}_2)]^{2+}\) (37) was found to be inactive.\(^{128}\)

The use of the anionic carboxamido ligand in Mn complex 35 is most apparent in the comparison of the \(\text{O}_2\) evolution rate, where complex 35 had a significantly higher rate than that of Mn complex 36. The different features of the anionic PaPy\(^3\) ligand compared to the neutral ligand backbones are further highlighted by the fact that complexation with Mn produces the Mn\(^{III}\) complex 35 in air. However, the corresponding Mn\(^{III}\) complex \([\text{Mn(PY5)}(\text{OH}_2)]^{3+}\) (37\(^+\)) requires oxidation with
e.g. PhIO. The MnIII complex \( \text{37}^+ \) is stable in air but decomposes rapidly in aqueous solutions to heterogeneous Mn oxides. The stability difference between the two MnIII complexes \( \text{35}^+ \) and \( \text{37}^+ \) demonstrates the power of using anionic ligand backbones for stabilizing high-valent Mn species, a critical feature for the activation of \( \text{H}_2\text{O} \).

### 4.3. Mn cubanes—bioinspired mimics of the Mn\(_4\)Ca core in Photosystem II

Molecular mimics of the Mn\(_4\)Ca cluster core in the OEC have attracted significant attention from researchers. Such artificial Mn clusters are highly desired and could facilitate the understanding of the chemical features of the OEC, and could ultimately be used for designing artificial molecular Mn-based WOCs. Numerous multinuclear Mn clusters have been developed as artificial OEC mimics, for example, the \([\text{Mn}\text{3CaO}_4]\) sub-units (Fig. 19, bottom), and the \([\text{Mn}\text{IVCa}_2\text{O}_4\text{OTf}_2\text{Bu}_2\text{CO}_2\text{H}_4]\) cubane.

The heterometallic \([\text{Mn}^\text{IIICaNa}]\) cluster \( \text{39} \) based on a Schiff base ligand (Fig. 20) was recently reported by Reedijk and coworkers. This Mn\(_3\)Ca cluster was found to evolve O\(_2\) using NaOCl, Oxone® or \( \text{tBuOOH} \) as a chemical oxidant. However, the molecular nature of the developed cluster was not thoroughly investigated in order to rule out the formation of heterogeneous metal oxides as the active catalytic entity.

Perhaps the most well-studied tetranuclear Mn mimic is the \([\text{Mn}\text{IV}_4\text{dpp}]\) cubane \( \text{40} \) (dpp = diphenylphosphinate), which contains a cubical \([\text{Mn}\text{O}_4]^{7+}\) core surrounded by six facially bridging bidentate diphenylphosphinate groups coordinated to the Mn centers (Fig. 21). Mn cubanes, such as the \([\text{Mn}_3\text{O}_4]\) cubane \( \text{40} \), self-assemble from mononuclear precursors or from dimeric complexes housing a \([\text{Mn}\text{O}_2]^{3+}\) core in the presence of phosphinate ligands. The three oxo units and the three anionic phosphinate ligands provide an electron-rich environment for the Mn centers. The electrochemical properties of the \([\text{Mn}_4\text{O}_4]\) cubane \( \text{40} \) have also been investigated. Electrochemical oxidation of the \([\text{Mn}\text{O}_4]^{6+}\) core in cubane \( \text{40} \) occurs at a potential of \( \sim 0.94 \) V vs. NHE and generates the one-electron oxidized Mn\(_{\text{III}}\)Mn\(_{\text{IV}}\) core. The redox potentials are ligand-dependent, indicating that they can be tuned using supporting phosphinate ligands with stronger electron donation.

The \([\text{Mn}\text{IV}_4\text{dpp}]\) cubane \( \text{40} \) has also been reported to liberate stoichiometric amounts of O\(_2\) upon UV-light irradiation. The necessity for one of the phosphinate ligands to dissociate before O\(_2\) evolution can occur was demonstrated by isotopic labeling experiments in combination with mass spectrometry, with O–O bond formation occurring via direct coupling of two bridging oxo groups. Furthermore, Mn cubane \( \text{40} \) has been suspended in Nafion, a proton-conducting polymer matrix, for sustained O\(_2\) evolution. The generated Nafion polymer films were deposited on a variety of electrodes and reached TONs of \( >1000 \) and rates of 0.075 \( \text{O}_2\) molecules.
per s per Mn$_4$ cluster at an overpotential of 0.38 V at pH 6 when illuminated with light of wavelength 275–750 nm.\textsuperscript{155–159} A photoelectrochemical cell has also been designed where the Nafion film embedded with cubane 40\textsuperscript{40} was coupled to a [Ru(bpy)$_3$]$^{2+}$-type sensitized TiO$_2$ layer (Fig. 22). The produced photoelectrochemical cell was able to catalyze the photooxidation of H$_2$O using visible-light as the sole energy source, reaching TONs of 13 O$_2$ molecules per Mn cluster and TOFs of $\sim$0.013 O$_2$ molecules per s per Mn cluster.\textsuperscript{160} However, a recent report suggests that bulk electrolysis of the Nafion embedded cubane causes it to dissociate into Mn$^{II}$ compounds. These compounds are subsequently reoxidized to produce nanoparticles composed of Mn$^{III/IV}$ oxides; thus the original Mn cubane cluster merely serves as a precursor to the catalytically active nanoparticle material.\textsuperscript{161}

Ca$^{2+}$ is essential in the natural OEC for triggering O–O bond formation and O$_2$ release and tuning the electrochemical properties of the Mn$_4$Ca cluster.\textsuperscript{162–164} Agapie and co-workers therefore recently synthesized a [Mn$_4$CaO$_4$]$^{6+}$ cubane 41 that models the Mn$_4$Ca core of the OEC. Structural and electrochemical comparison between the [Mn$_4$CaO$_4$]$^{6+}$ cubane 41 (Fig. 23, left) and a related Mn$_4$O$_4$ cubane 42 (Fig. 23, right) demonstrated that the redox-inactive metal facilitates access to higher redox states and the assembly of the cluster.\textsuperscript{165} Subsequent work focused on the synthesis of a series of [Mn$_3$M($\mu_3$-O)]($\mu_2$-O)] clusters\textsuperscript{166} (where M = Na$^+$, Ca$^{2+}$, Sr$^{2+}$, Zn$^{2+}$, or Y$^{3+}$) and [Mn$_3$M’O$_4$] clusters\textsuperscript{167} (where M’ = Sr$^{2+}$, Zn$^{2+}$, Sc$^{3+}$, Y$^{3+}$; see Fig. 24 for the structure of the Mn$_3$ScO$_4$ cubane 43). Here, the redox-inactive metal was shown to modulate the redox potentials of the heterobimetallic Mn clusters. These studies have provided a methodology with unprecedented structural control for the rational synthesis of Mn-based mimics of the Mn$_4$Ca core of the OEC.\textsuperscript{168–171} The [Mn$_3$M’O$_4$] cubanes have recently been dropcast on ITO or glassy carbon disk electrodes and used as precursors to heterogeneous electrocatalysts for H$_2$O oxidation.\textsuperscript{172} Multinuclear clusters of the type [Mn$_{12}$O$_{12}$] have also been demonstrated to be viable H$_2$O oxidation electrocatalysts.\textsuperscript{173}

Mechanistic understanding of the biological Mn$_4$Ca cluster in the OEC at a molecular level is highly desirable for the rational design of viable WOCs. The collective work on the Mn-based model clusters has provided valuable insight into
5. Homogeneous Fe-based complexes for water oxidation

Fe is an essential element in a variety of enzymes and perhaps the most important transition metal from a biological point of view. Fe catalysis has emerged as a prominent research field and in recent years there has been a tremendous development in this area. Fe is known to have a rich redox chemistry and to adopt a variety of redox states. In addition to having access to this broad spectrum of redox states, Fe-complexes have the ability to engage in both one- and two-electron processes—essential features for 

H₂O oxidation. The large number of spin states accessible to Fe should also make it amenable to tuning, by varying the ligand environment. Although Fe is associated with these attractive properties, relatively few examples exist of Fe-catalyzed 

H₂O oxidation.

5.1. Mononuclear Fe-based catalysts

Seminal work on Fe-catalyzed 

H₂O oxidation has already been carried out in the 1980s. However, the first examples of homogeneous Fe-based WOCs were reported by Collins and co-workers in 2010. In order to stabilize the Fe center, the authors employed tetraamido macrocyclic ligands (TAMLs). These ligands have previously been shown to efficiently activate 

O₂ and peroxides to produce 

FeIV–μ-oxo species. In aqueous solutions, the 

FeV–μ-oxo species have been shown to participate in a pH-dependent equilibrium with a monomeric 

FeIV–oxo complex. Furthermore, the initial FeV–TAML complexes react with peracids to produce the corresponding 

FeV–oxo complexes. The diverse and high reactivity associated with the Fe–TAML complexes thus rendered these catalysts attractive targets for use in 

H₂O oxidation.

Fe–TAML complexes 44–48 were therefore investigated toward 

H₂O oxidation. In the presence of 

CeIV as a chemical oxidant, 

fast 

O₂ evolution was observed for complexes 45–48. However, no 

O₂ production was detected for the least acid stable 

Fe–TAML complex 44. The rate of 

O₂ evolution was shown to be highly dependent on the electron-withdrawing ability of the macrocyclic ligand. Fe–TAML complex 48 gave the highest TOF, 1.3 s⁻¹, and was shown to display a first-order dependence on the catalyst concentration, implying that 

O–O bond formation occurs at a single metal center. 

H₂O oxidation could also be driven using 

NaIO₄ as a chemical oxidant. Mechanistic studies using 

NaIO₄ as an oxidant also concluded that 

an 

FeIV–μ-oxo–FeIV species is involved in the oxidation process. Although this seminal work highlighted that Fe-based WOCs could be designed, the catalysts were prone to undergo fast deactivation/decomposition, highlighting that more robust WOCs need to be targeted. In an attempt to improve the longevity of the Fe–TAML complexes, they were subsequently immobilized on electrodes for electrocatalytic 

H₂O oxidation.

Dhar and co-workers subsequently employed the modified 

Fe–TAML complex 49 to carry out light-driven 

H₂O oxidation using 

[Ru(bpy)₃]²⁺ as a photosensitizer and 

Na₃S₂O₈ as a sacrificial electron acceptor. The photocatalytic system was shown to produce a TON of 220 and a TOF of 0.67 s⁻¹. An 

FeV–oxo intermediate was proposed to be generated in this system as supported by EPR, UV-vis and HRMS studies. Based on these results and related studies, the authors proposed a mechanism, outlined in Scheme 4, where the generated 

FeV–oxo species 52 undergoes nucleophilic attack by a 

H₂O molecule to form a 

FeIII-hydroperoxo 

(FeIII–OOH) species (53). Subsequent oxidation of this 

FeIII–OOH species leads to...
O₂ liberation and regenerates the starting aqua complex 49, FeIII–OH₂.¹⁸³

An alternative mechanism for the oxidation of H₂O by the Fe–TAML complexes was presented by Liao and Siegbahn.¹⁸⁹ This mechanism was proposed to involve the formation of a formal FeVI–oxo species, which upon further inspection was better described as having an FeV center with a ligand cation radical. Three distinct ligand modification pathways were also realized: (1) H₂O or (2) nitrate (the anion originating from the CeIV oxidant) attack on the ligand framework, and (3) amide oxidation. The pathway involving H₂O attack on the ligand was associated with a low barrier and results in the opening of the benzene ring. The observed reactivity pattern for the Fe–TAML complexes 44–48 could also be rationalized as it was found that the barrier for O–O bond formation decreased with electron-withdrawing substituents. However, the introduction of electron-withdrawing units increases the redox potentials and adds an additional energetic penalty. These effects need to be carefully balanced in future designs of ligands.

Another class of Fe-based WOCs was described by Lloret-Fillol and co-workers.¹⁹⁰ The authors evaluated the catalytic activity of a series of Fe complexes based on tetra- and pentadentate ligand motifs (Fig. 27). From the study, it could be shown that Fe complexes 54–58 having two open coordination sites in a cis configuration were competent WOCs. The structural requirement was further confirmed by complexes 59 and 60, with a trans configuration or possessing only a single coordination site, which were found to be inactive. Of the evaluated catalysts, Fe complex 55α afforded a TON of >1000 and a TOF of 0.062 s⁻¹ using NaIO₄ as a chemical oxidant. Mechanistic investigations employing the dimeric Fe₃III,III₂(μ-OH)(μ-O) complex of catalyst 55α showed that the dimeric complex was associated with a lower reaction rate and a different kinetic behavior, discarding such dimeric structures as essential active intermediates in the studied Fe-based system. Addition of 6 equivalents of CeIV oxidant resulted in the appearance of new
bands in the UV-vis, a species characteristic of an Fe$^{IV}$–oxo intermediate. The persistence of this Fe$^{IV}$–oxo indicated that this species is not responsible for mediating O–O bond formation, and that species of higher valency are needed for O–O bond forming events.

Several mechanistic studies on Fe complexes of the type $54$–$58$ have also been carried out.$^{191–195}$ A follow-up study by Lloret-Fillol and co-workers involved the electronic tuning of the Fe-based catalyst $54$ in which a family of substituted Fe complexes ($61$–$65$, Fig. 28) were synthesized. Using Hammett parameters, the rate of O$_2$ evolution was found to correlate with the electronic nature of the introduced substituents, with electron-withdrawing substituents favoring O$_2$ production. The non-innocent effect of the Ce$^{IV}$ cation was also observed in which a Fe$^{IV}$(OH)(O$–$Ce$^{IV}$) adduct was proposed to be involved prior to the O–O bond forming step.$^{191}$

A recent study identified Fe$^{IV}$(O$–$Ce$^{IV}$) as the key reaction intermediate in H$_2$O oxidation using Fe complex $55$.$^{193}$ Here, Ce$^{IV}$ was found not to act merely as an inner-sphere oxidant but was revealed to generate an Fe$^{IV}$($\mu$-O)Ce$^{IV}$ species, crucial for the catalytic reaction to proceed. The oxidation of H$_2$O with Fe-based complexes such as $55$ not only requires the presence of two open coordination sites in a cis configuration but also a framework that allows the generation of Fe$^{IV}$($\mu$-O)Ce$^{IV}$. The isomeric Fe complexes $55_a$ and $55_b$ also displayed different reactivity, arising either from steric encumbrance or electronic effects. While the complex $55_b$ did not evolve O$_2$, the $\alpha$-isomer was found to produce O$_2$ according to the pathways highlighted in Scheme 5. These findings could thus provide valuable information for future evaluations of earth-abundant WOCs.

The relative ease by which Fe complexes housing tetra- and pentadentate nitrogen-based ligands can be accessed has attracted attention from several research groups.$^{196–198}$ The phosphonate modified Fe complex $71$ (Fig. 29; dpaq = 2-[bis(pyridine-2-ylmethyl)]amino-N-quinolin-8-yl-acetamido)$^{200}$ which has previously been reported to mediate alkane hydroxylation with H$_2$O$_2$ as an oxidant.$^{201}$ Meyer and co-workers found that the
[Fe(dpaq)(OH2)]2+ complex 72 was able to catalyze electrochemical H2O oxidation in propylene carbonate–water mixtures. Electrochemical examination revealed that the Fe catalyst exhibited a quasi-reversible one-electron wave at 0.38 V vs. NHE corresponding to the [FeIII–OH2]2+/[FeII–OH2]+ redox couple. At higher potentials (1.58 V), an irreversible two-electron wave appeared, which was assigned to the oxidation of [FeIII(OH2)]2+ to [FeV(O)]2+. Following this wave, an increase in current was observed. The peak current was shown to display a first-order dependence on the catalyst concentration, suggesting a rate-limiting reaction between [FeV(O)]2+ and H2O. This is consistent with a single-site mechanism for H2O oxidation as depicted in Scheme 6.200

A series of Fe-complexes containing pendant bases have also been synthesized by Yang and co-workers (76–82, Fig. 31).202 The authors envisioned that incorporation of bases into the second coordination sphere would facilitate PCET events and lead to enhanced reaction kinetics. However, Fe complexes 78–80 were shown to be inactive catalysts as they were not able to reach the critical FeIV=O state. For Fe complexes 81 and 82, an undesirable change in coordination took place, in contrast to what was observed for the analogous complexes lacking the introduced heteroatoms. Of the developed Fe complexes, only the pyridazine containing complexes 76 and 77 displayed H2O oxidation activity similar to the analogous complexes lacking the ancillary proton relays. Although the introduction of proton relays did not improve the catalytic activity of the Fe complexes, the concept of designing ligand motifs containing strategically functionalized moieties able to

Scheme 5  Reactivity differences between the isomeric Fe complexes 55α and 55β.
participate in PCET is an attractive feature for accessing improved Fe-based WOCs.

It needs to be stressed that the molecular nature of Fe-based complexes is highly dependent on the reaction conditions. Additionally, the catalytic activity of Fe-based complexes and the formation of metal nanoparticles are extremely susceptible to electronic and structural features. This intricate behavior is elegantly illustrated by recent examinations.
conducted by the groups of Lau and Fukuzumi. In these studies, the Fe-based complexes were shown to act as homogeneous catalysts under more acidic conditions while more basic conditions facilitated the formation of Fe-oxide nanoparticles.

The rational construction of ligand scaffolds, which allow high-valent Fe species to be generated and stabilized, still remains an essential challenge that needs to be addressed if more robust Fe-based catalysts are to be developed. As further insight into the mechanistic details of Fe-catalyzed \( \text{H}_2\text{O} \) oxidation emerges, new ways to design more efficient Fe-based systems will most certainly be devised.

5.2. Dinuclear Fe-based catalysts

Although a variety of mononuclear Fe-based complexes have been shown to catalyze \( \text{H}_2\text{O} \) oxidation, examples of dinuclear Fe complexes capable of mediating this transformation are currently limited. Initial observations on dinuclear Fe species being involved in the catalytic process were reported by Najafpour and co-workers. The authors selected the tris(2-pyridylmethyl)amine (tpa) ligand and synthesized the dinuclear Fe complex \([(\text{tpa})(\text{Cl})\text{Fe}(\mu-\text{O})\text{Fe}(\text{Cl})(\text{tpa})]^{2+} (83)\) which contains a \( \mu \)-oxo bridge. When the dinuclear Fe complex 83 was examined as a WOC using CeIV as a chemical oxidant, the rate for \( \text{O}_2 \) evolution was found to be significantly higher than for its mononuclear analog 58 (see Fig. 27). The rate was found to be first order in both CeIV and catalyst 83, suggesting that O-O bond formation occurs within the dinuclear framework.

Subsequent studies on dinuclear Fe-based WOCs by the groups of Sakai and Ma also centered on using the tpa ligand. The group of Sakai synthesized the dinuclear Fe complex \([(\text{tpa})\text{Fe}(\mu-\text{O})(\text{µ-} \text{SO}_4)\text{Fe}(\text{tpa})]^{2+} (84)\) and established that it mediated \( \text{H}_2\text{O} \) oxidation using NaIO\(_4\) as a chemical oxidant. DLS analysis suggested that the dinuclear Fe complex served as a homogeneous WOC and that Fe-oxide nanoparticles were not formed under the investigated reaction conditions. Kinetic experiments showed a first-order dependence on the catalyst concentration. Unexpectedly, the rate of \( \text{O}_2 \) evolution exhibited a dramatic increase when the pH was decreased. Such an effect is uncommon since the rate-determining step is usually nucleophilic attack of \( \text{H}_2\text{O} \) or coupling of two oxyl units. It was speculated that the observed rate enhancement could be due to reduced anion at low pH. Ma and co-workers subsequently investigated the catalytic activity of the Fe complex \([(\text{tpa})(\text{Cl})\text{Fe}(\mu-\text{O})\text{Fe}(\text{Cl})(\text{tpa})]^{2+} (85)\). In aqueous acetate buffer solutions, using Oxone\(_\text{®}\) as a chemical oxidant, Fe complex 85 exhibited high \( \text{O}_2 \) evolution activity with TONs reaching 2380 and TOFs of 2.2 s\(^{-1}\). HRMS suggested that upon dissolution in acetate buffer solutions, the acetate bridged dinuclear Fe species \([(\text{tpa})\text{Fe}(\mu-\text{O})(\mu-\text{OAc})\text{Fe}(\text{tpa})]^{2+}\) was formed. This was proposed to be essential because the two free coordination sites are constrained in a \textit{cis} fashion, allowing for efficient \( \text{H}_2\text{O} \) oxidation.

Thummel and co-workers recently revealed that the dinuclear Fe\(_{\text{III,III}}\) complex \([(\text{ppq})(\text{OH}_2)\text{Fe}(\mu-\text{O})\text{Fe}(\text{Cl})(\text{ppq})]^{2+} (86)\), Fig. 33; \text{ppq} = 2-(pyrid-2′-yl)-8-(1″,10″-phenanthrolin-2″-yl)-quinoline) was able to mediate \( \text{H}_2\text{O} \) oxidation. The tetradentate ppq ligand was previously employed in the synthesis of a Co-based complex for \( \text{H}_2 \) evolution. Treating FeCl\(_3\) with the ppq ligand afforded the \( \mu \)-oxo bridged dinuclear Fe\(_{\text{III,III}}\) complex 86 in 28% yield. Electrochemical analysis of Fe complex 86 revealed a reversible wave at ~0.21 V vs. NHE and another quasi-reversible wave at ~0.69 V. The first event could be assigned to a two-electron process and was tentatively attributed to the simultaneous oxidation of the two Fe\(_{\text{III}}\) centers, Fe\(_{\text{II,II}}\) → Fe\(_{\text{IV,IV}}\). This produced [Fe\(_{\text{II}}\)\text{Fe}_{\text{IV}}\text{O}\(_2\)]\(^{2+}\) which could undergo disproportionation with loss of two protons to afford a [Fe\(_{\text{III}}\)\text{Fe}_{\text{IV}}\text{O}]=\text{O}_2\) species. \( \text{H}_2\text{O} \) oxidation was evaluated using Ce\(_{\text{IV}}\) as a chemical oxidant. Upon addition of the oxidant to a solution of complex 86, rapid \( \text{O}_2 \) evolution was observed. As a comparison, the mononuclear Fe complex 87 (Fig. 33) was synthesized and examined as a WOC. Although the mononuclear complex 87 was able to oxidize \( \text{H}_2\text{O} \), the dinuclear Fe complex 86 was found to be more active, affording a TON of ~1000 and a TOF of 2.2 s\(^{-1}\). An essential question was whether the dinuclear Fe complex 86 retains its dimeric structure or whether it dissociates to yield a monomeric species in solution. UV-vis analysis and the apparent first-order dependence on the catalyst concentration suggested that the Fe complex remains intact during the catalytic oxidation. The involvement of

![Fig. 32](image-url) Structures of the dinuclear Fe complexes \([(\text{tpa})(\text{OH}_2)\text{Fe}(\mu-\text{O})\text{Fe}(\text{OH}_2)(\text{tpa})]^{2+} (83)\), \([(\text{tpa})\text{Fe}(\mu-\text{O})(\mu-\text{SO}_4)\text{Fe}(\text{tpa})]^{2+} (84)\) and \([(\text{tpa})(\text{Cl})\text{Fe}(\mu-\text{O})\text{Fe}(\text{Cl})(\text{tpa})]^{2+} (85)\).
nanoparticles in the studied catalytic system could successfully be discarded based upon DLS measurements. The collective work on dinuclear Fe-based catalysts highlights that dinuclear species have the potential of generating more active catalytic systems compared to their mononuclear counterparts. The continued endeavors in this field will likely result in a rapid expansion of viable, and more robust, dinuclear Fe-based catalysts. This is further highlighted by the fact that a recently developed homogeneous pentanuclear Fe complex is able to facilitate \( \text{O}_2 \) bond formation with an impressive efficiency.

6. Co-based systems for water oxidation

Inorganic Co salts have been studied as potential WOCs since the late 1960s.175,211–213 Although these early attempts were rather unsuccessful due to precipitation of heterogeneous species, they suggested the involvement of Co\(^{IV} \) species during the catalytic process.214 Spurred by the findings of Nocera and co-workers that CoO\(_x\) species (CoP, catalyst, Fig. 34),215–229 generated \textit{in situ} under neutral conditions in phosphate buffer, exhibited \( \text{O}_2 \) evolution activity, several research groups have attempted to prepare homogeneous molecularly defined Co-based catalysts for the oxidation of \( \text{H}_2\text{O} \).

6.1. Co polyoxometalates (POMs)—structurally defined all-inorganic catalysts

In 2010, Hill and co-workers reported that Co-based polyoxometalates (POMs) were able to evolve \( \text{O}_2 \) when driven by pregenerated or photochemically generated \([\text{Ru(bpy)}_3]^{3+}\).230 The all-inorganic Co-POM catalysts can be considered as the homogeneous alternative to metal oxides with the advantage of being well-defined and consisting of only inorganic elements. Prior to the report dealing with the Co-POMs, the groups of Hill and Bonchio had simultaneously shown that their Ru-analogs were efficient catalysts for the oxidation of \( \text{H}_2\text{O} \).235,236 With the goal of developing an earth-abundant POM, Hill and co-workers synthesized the tetranuclear Co-POM \([\text{Co}_4(\text{OH}_2)_2(\mu-\text{Cl})(\text{ppq})]^{13+}\) (Fig. 35) and showed that it was capable of driving \( \text{H}_2\text{O} \) oxidation with \([\text{Ru(bpy)}_3]^{3+}\). Subsequent studies revealed that Co-POM \( 88 \) also catalyzed photoinduced \( \text{H}_2\text{O} \) oxidation using \([\text{Ru(bpy)}_3]^{2+}\) as a photosensitizer and persulfate (\( \text{S}_2\text{O}_8^{2-} \)) as a sacrificial electron acceptor, generating a TON of 224 and a quantum yield of 30%.231 Co-POM \( 88 \)'s \( \text{H}_2\text{O} \) oxidation activity and other essential features have also been studied by other research groups.237–240 A recent report by the Hill group has shown that the structurally analogous Co-POM \([\text{Co}_4(\text{H}_2\text{O})_2(\text{VW}_9\text{O}_{34})]^{10-}\) is an extremely efficient WOC, reaching TOFs >1000 \( \text{s}^{-1} \) with quantum yields approaching 68%.241 Additional Co-POM derivatives have also been synthesized by the Wang group and shown to mediate photocatalytic \( \text{H}_2\text{O} \) oxidation.242
excess 2,2

oxide film on the electrode could be avoided by addition of H2O oxidation was performed. However, formation of a Co whereas slow release of Co was observed when electrochemical geneous operating mechanism under chemical oxidation NaClO as a chemical oxidant. Experiments supported a homo-

cal catalysis.246

involvement of heterogeneous metal oxide species during

and pH, which needs to be considered when determining the

further highlight that the homogeneity of WOCs is highly

or whether decomposition resulted in Co oxides which were the real catalytic entities. From these extensive experiments it was suggested that Co-POM 88 did not decompose and operated via a homogeneous mechanism.230 Although Hill and co-workers concluded that Co-POM 88 was stable under the studied reaction conditions, subsequent reports challenged these conclusions, showing that POM 88 decomposed to Co oxides as the dominant cata-

lyst.239,243 Additional studies revealed that the slight modifi-
cations of the reaction conditions influenced the operating mechanism and indicated that the conclusions from the original publication on Co-POM 88 were indeed correct.244,245 They further highlight that the homogeneity of WOCs is highly dependent on the reaction conditions, such as the buffer and pH, which needs to be considered when determining the involvement of heterogeneous metal oxide species during catalysis.246

The group of Galán-Mascarós subsequently reported on a high-nuclearity Co-POM, [Co4(OH2)2(OH3)3(HPO4)2(PW9O43)2]16− (89, Fig. 36).247 The nonanuclear Co core is stabilized by three hydroxo and two hydrogen phosphate bridges. The nonnucelar Co-POM 89 was found to catalyze H2O oxidation using NaClO as a chemical oxidant. Experiments supported a homo-

geneous operating mechanism under chemical oxidation whereas slow release of Co was observed when electrochemical H2O oxidation was performed. However, formation of a Co oxide film on the electrode could be avoided by addition of excess 2,2'-bipyridine, which functions as a chelating agent for the liberated CoII or CoIII. A TON of ~400 and a TOF of ~0.1 were measured, with maintained O2 evolution activity for several days without any sign of fatigue or decomposition.247

The same group also synthesized an insoluble salt (Cs salt) of Co-POM 89 for integration into amorphous carbon paste elec-

Additional examples of Co-based POMs that have been reported to mediate visible-light driven H2O oxidation include the mixed dinuclear Co-POM [Co8CoIII(OH)3W11O39]7− (90),249 the octanuclear Co-POM [(Au3SiW10O42)2Co4(μ-OH)(μ3-OH2)2(CO3)4]16− (91,250 [CoII(bpy)3](H2bpy)][Co3(bpy)2(PMo6V2Mo8O30)][(Co3bpy)-

(CO4VMo10O30)] (92),251 [Co2Bi2−(μ-B-CoW9O43)2]14− (93),252 [Co2Mo10O38H4]4+ (94)253 and [(Co(OH2)3]2[CoBi2W7O46(OH)3]10− (95).254 From the plethora of studies conducted on Co-based POMs, it is clear that these catalysts can operate as robust and efficient molecular catalytic entities for the oxidation of H2O.

6.2. Co cubanes as mimics of the catalytic core in the oxygen-evolving complex

Inspired by the structural rearrangement of the Mn4Ca core in the OEC, researchers have devoted considerable efforts to synthesize complexes with similar cubane core structures. Early examples of such complexes include the Co cubanes Co4O4(pyr)4(OAc)4 ([Co2Mo10O38H4]6− (96),255,256 and [Co2O4(bpy)6(OAc)4]2+ (97) (Fig. 37).257,258 The Co4O4(pyr)4(OAc)4 (96) cubane can be easily prepared from Co(NO3)2, NaOAc and pyridine and contains a Co4O4 core, which makes this cubane and related systems attractive as potential WOCs. A seminal report by Dismukes and co-workers showed that cubane 96 indeed was able to promote light-driven H2O oxidation using a three-component system consisting of [Ru(bpy)3]2+ as a photosensitizer and Na2S2O8 as a sacrificial electron acceptor. This photocatalytic system generated a TON of 40 and a TOF of 0.02 s−1.259

Subsequent studies by Bonchio and co-workers involved the examination of isostructural analogs of Co cubane 96.260,261 The authors targeted Co cubanes 98–103 (Fig. 38) in order to study catalyst tuning for boosting the photocatalytic efficiency in the three-component system previously used by Dismukes
and co-workers. Hammett linear free energy plots were employed and revealed a correlation between the photo-induced electron transfer constants and the electron-donating ability of the ligand. However, under the explored conditions all of the investigated cubanes reached similar TONs (∼140), with the difference being the rate by which O₂ was produced.

Ofoli and co-workers recently immobilized Co cubane 96 on ITO and showed that the cubane catalyst retained its catalytic activity similarly to the homogeneous catalytic entity. Sun and co-workers have also coupled Co cubane 96 to [Ru(bpy)₃]²⁺-type photosensitizers to generate two supramolecular assemblies, assembly 104 depicted in Fig. 39 and the cyclic assembly 105 consisting of two Co cubane and two [Ru(bpy)₃]²⁺ units (not shown). The carboxylate motif was chosen since it has been shown that Co cubane 96 can be immobilized onto carboxylate functionalized silica via carboxylate exchange.

Electrochemical measurements revealed that assembly 104 displayed two redox peaks at 1.01 V and 1.28 V vs. NHE. The cyclic assembly 105 also exhibited two redox peaks; however, these were positively shifted and appeared at 1.10 V and 1.34 V vs. NHE. In comparison, a solution containing a mixture of the separate components, Co cubane 96 and [Ru(bpy)₃]²⁺ (1), displayed two waves at 0.91 V and 1.26 V vs. NHE. The two assemblies were subsequently evaluated toward light-driven H₂O oxidation using Na₂S₂O₈ as a sacrificial electron acceptor. When exposed to light, the two assemblies 104 and 105 rapidly evolved O₂. Here, the cyclic assembly 105 proved to be a more efficient catalyst than assembly 104, producing a TOF of 0.023 s⁻¹ compared to 0.0067 s⁻¹ for the linear assembly 104. Although the reason for the striking difference in activity for the cyclic assembly 105 is still not clear, the authors speculated that assembly 105 was more robust than the linear counterpart. This hypothesis originated from the fact that cyclic assembly 105 exhibited fewer changes in the UV-vis spectrum during the photocatalytic experiments compared to assembly 104.
A mechanistic investigation conducted by Tilley and co-workers on the $\text{Co}_4\text{O}_4(\text{pyr})_4(\text{OAc})_4$ cubane (96) suggested that the stoichiometric reaction of the one-electron oxidized cubane (96$^+$, [Co$^{\text{III}}$Co$^{\text{IV}}$]) with OH$^-$ produces O$_2$ with quantitative regeneration of cubane 96. The experimental results also suggested that the Co$^{\text{III}}$Co$^{\text{IV}}$ species (96$^+$) undergoes disproportionation to generate an intermediate with an even higher oxidation state, a formal Co$^{\text{III}}$Co$^{\text{V}}$ or Co$^{\text{II}}$Co$^{\text{IV}}$ species. The proposed mechanism for the reduction of 96$^+$ by OH$^-$ is depicted in Scheme 7. The mechanism by which Co cubanes mediate H$_2$O oxidation has also been investigated by density functional theory (DFT) calculations. Several reaction pathways were considered; however, the lowest energy pathway for Co cubane 96 was found to proceed through a formal Co$^{\text{V}}$ species, which is perhaps better described as a Co$^{\text{IV}}$ center coupled to an oxygen radical. The mechanism with the lowest energy involved water nucleophilic attack on the formed oxygen radical. In a related study, the pathway was observed to involve germinal coupling of a formal Co$^{\text{V}}$-oxo unit with bridging oxo sites. In this study the examined models were revealed to be sensitive to positions of the ligands and the hydrogen bonding environment, resulting in distinct isomers with different energies. The studies described above highlight the necessity of designing Co cubanes, and other Co-based WOCs, that are capable of accessing high-valent redox states. A multimetallic system is well-suited to stabilize such high redox states but is not a prerequisite for designing active Co-based WOCs since the ancillary ligands can also be an essential feature for tuning the redox potentials of the metal complexes.

Co cubane $[\text{Co}_5(\text{hmp})_4(\mu-\text{OAc})_2(\mu_2-\text{OAc})_2(\text{H}_2\text{O})_2]$ (112, Fig. 40; hmp = 2-(hydroxymethyl)pyridine) was synthesized by...
The catalytic activity of Co cubane 112 was evaluated in light-driven H₂O oxidation using [Ru(bpy)₃]³⁺ as a photosensitizer and Na₂S₂O₈ as a sacrificial electron acceptor. From these experiments it was established that the O₂ evolution performance was strongly dependent on the pH. The TOF was shown to change notably over the studied pH range, from 1.8 s⁻¹ at pH 7 to 7 s⁻¹ at pH 9. Several different techniques were used to determine the extent of nanoparticle formation and suggested that the Co cubane core of 112 remains intact. The high catalytic activity of Co cubane 112 was attributed to its flexible architecture consisting of monodentate acetate and aqua ligands in combination with a robust core. A recent computational study suggests that O–O bond formation occurs either through a single-site pathway involving H₂O attack on a Co₃O₅ core or through an oxo–oxo coupling pathway.

Subsequent work on biomimetic Co cubanes consisted of the synthesis and examination of a series of isostuctural Co cubanes (113–116) with the general formula [Co³⁺Ln(hmp)₄(OAc)₅H₂O] (Fig. 41, where Ln = Ho, Er, Tm, Yb). These bioinspired Co cubanes show several design features, such as ligand flexibility and redox-inert Ln³⁺ metal ions for electrochemical tuning. The catalytic performance of the lanthanide cubane series (113–116) was studied using [Ru(bpy)₃]³⁺ as a photosensitizer and Na₂S₂O₈ as a sacrificial electron acceptor in order to enable a direct comparison with the previously developed Co cubane 112. O₂ evolution peaked at pH 8–9, which was also observed for the [Co³⁺Ln(hmp)₄(μ₂-OAc)₅(H₂O)] cubane (112). Table 1 summarizes the catalytic activity of the biomimetic lanthanide cubanes [Co³⁺Ln(hmp)₄(OAc)₅H₂O]. The stability of the lanthanide containing Co cubanes was subsequently assessed employing a variety of different techniques, including UV-vis aging tests, DLS, and extended X-ray absorption fine structure (EXAFS) and X-ray absorption near-edge structure (XANES) solution phase tests, which supported the structural integrity of the [Co³⁺Ln(hmp)₄(OAc)₅H₂O] cubane core under the catalytic conditions.

Although the stability of the various Co cubane WOCs was assessed and pointed to a homogeneous H₂O oxidation pathway operating with the cubane core being intact, a report from the Nocera group suggested that the catalytic activity emanates from a Co³⁺ impurity. The Co³⁺ impurity was proposed to act as a source for the generation of heterogeneous Co species that are the real catalysts. These heterogeneous species are believed to be formed if the potential is sufficiently high to oxidize the Co³⁺ to Co⁴⁺ in the presence of a proton accepting electrolyte, such as phosphate and carbonate. A recent report also questions the homogeneity of the Co-based cubane complexes, and highlights the dependence on the initial structure and the catalytic methodology being used.

From the conflicting results regarding the homogeneity of Co cubanes it is clear that care must be exercised when examining whether a specific WOC operates through a homogeneous mechanism or whether heterogeneous species are produced in situ, and the studied molecular complex merely acts as a precursor. However, deriving a definite conclusion for the examined catalytic system is not always straightforward and can be highly sensitive to the methodologies and experimental conditions being employed.

### Table 1 Comparison of photocatalytic H₂O oxidation activities for Co cubanes 113–116 over the pH range 7–9

| pH   | Catalyst concentration (μM) | TON | TOF (s⁻¹) |
|------|----------------------------|-----|-----------|
|      | [Co³⁺Ho(hmp)₄(OAc)₅H₂O] (113) |     |           |
| 7⁻⁰  | 12                         | 7   | 0.87      |
| 8⁻⁰  | 10                         | 163 | 5.84      |
| 9⁻⁰  | 12                         | 135 | 9.55      |
|      | [Co³⁺Er(hmp)₄(OAc)₅H₂O] (114) |     |           |
| 7⁻⁰  | 12                         | 2   | 2.06      |
| 8⁻⁰  | 10                         | 211 | 5.65      |
| 9⁻⁰  | 12                         | 108 | 5.02      |
|      | [Co³⁺Tm(hmp)₄(OAc)₅H₂O] (115) |     |           |
| 7⁻⁰  | 12                         | 24  | 1.55      |
| 8⁻⁰  | 10                         | 92  | 5.34      |
| 9⁻⁰  | 12                         | 64  | 3.48      |
|      | [Co³⁺Yb(hmp)₄(OAc)₅H₂O] (116) |     |           |
| 7⁻⁰  | 12                         | 1.1 | 1.08      |
| 8⁻⁰  | 10                         | 160 | 6.54      |
| 9⁻⁰  | 12                         | 120 | 5.65      |

* Photochemical experiments were carried out with a catalyst (12 μM), [Ru(bpy)₃]Cl₂ (1 mM) and Na₂S₂O₈ (5 mM) in aqueous phosphate buffer solutions (pH 7, 40 mM Na₂HPO₄/NaH₂PO₄ buffer).
* Experiments were performed with a catalyst (10 μM), [Ru(bpy)₃]Cl₂ (1 mM) and Na₂S₂O₈ (5 mM) in aqueous borate buffer solutions (pH 8, adjusted with HCl addition to 50 mM borate buffer). Experiments were performed with a catalyst (12 μM), [Ru(bpy)₃]Cl₂ (1 mM) and Na₂S₂O₈ (5 mM) in aqueous borate buffer solutions (pH 9, adjusted with HCl addition to 50 mM borate buffer).

### 6.3 Mononuclear Co catalysts

The early observation that Co⁴⁺ species could be involved in the catalytic oxidation of H₂O using simple inorganic salts...
suggested that one key to realizing molecular Co WOCs might be to design ligands that permit stabilization of high-valent Co intermediates. Two seminal reports by the groups of Berlinguette\textsuperscript{274} and Nocera\textsuperscript{275} showed that this principle was indeed viable. Berlinguette and co-workers utilized the pyridine-based pentacoordinating ligand Py5\textsuperscript{126,127} giving Co-based complex [Co(Py5)(OH\textsubscript{2})]\textsuperscript{2+} (117, Fig. 42) with a single open coordination site. The Py5 ligand was chosen as the authors envisioned it to be a good motif for withstanding the harsh oxidative conditions needed to carry out oxidation of H\textsubscript{2}O. Furthermore, the Py5 ligand lacks β-hydrogens, which prevents that the corresponding metal complex undergoes undesirable elimination processes. Electrochemical measurements showed that Co complex 117 displayed two redox events. The second event results in a significant rise in the current, suggesting that a catalytic reaction takes place. The catalytic current was demonstrated to increase with decreasing scan rate (ν), attesting that a chemical process, such as O–O bond formation, is the rate-determining step. The authors attributed this event to nucleophilic attack of H\textsubscript{2}O on a high-valent formal Co\textsuperscript{IV}–oxo/hydroxo species, generated after the initial two redox steps.\textsuperscript{274} A subsequent study conducted on the [Co(Py5)(OH\textsubscript{2})]\textsuperscript{2+} complex 117 supported that the catalytic current was molecular in origin and that the H\textsubscript{2}O oxidation did not emanate exclusively from \textit{in situ} formed heterogeneous Co species.\textsuperscript{276} A detailed computational mechanistic study on Co complex 117 suggested that the resting state of the catalyst contained a Co\textsuperscript{IV}–oxyl species. The O–O bond formation was suggested to proceed \textit{via} nucleophilic attack of OH\textsuperscript{−} on this formal Co\textsuperscript{IV}–oxyl species.\textsuperscript{277} The structurally related Co\textsuperscript{II} complex [Co(Py5OH)(Cl)]\textsuperscript{+} (118, Fig. 42) has also recently been reported to mediate H\textsubscript{2}O oxidation.\textsuperscript{278}

Based on previous observations that Co\textsuperscript{II} hangman porphyrin complexes were able to activate O\textsubscript{2},\textsuperscript{279} Nocera and co-workers decided to investigate whether the related Co\textsuperscript{II} corrole complexes 119 and 120 (Fig. 43) could promote the opposite reaction—oxidation of H\textsubscript{2}O.\textsuperscript{275} These Co xanthene hangman corrole complexes have two vacant coordination sites and contain a proximal base. Measurements revealed that they could indeed drive H\textsubscript{2}O oxidation electrochemically, with the β-octafluoro Co complex 120 being the more active of the two.

**Fig. 42** Structures of Co-based complexes [Co(Py5)(OH\textsubscript{2})]\textsuperscript{2+} (117) and [Co(Py5OH)(Cl)]\textsuperscript{+} (118).

**Fig. 43** (Left) Co corrole complexes 119 and 120, and (right) organization of H\textsubscript{2}O within the Co hangman cleft.
complexes. The importance of the hangman cleft is believed to be its ability to arrange one H$_2$O molecule in the primary coordination sphere of the Co center and another molecule in the secondary coordination sphere via hydrogen bonding to the xanthene hangman motif (see Fig. 43). Subsequent quantum chemical modeling supported the feature of the carboxylate unit in functioning as a general base to activate the attack of the incoming H$_2$O molecule on the metal–oxo species. Additional key features in the catalytic cycle that were uncovered included the non-innocent role of the corrole backbone and that fluorination of the ligand backbone modulates the electrophilicity of the metal–oxo moiety and alleviates the decomposition of the produced corrole radical cations.

Another Co corrole complex that has been reported to catalyze electrocatalytic H$_2$O oxidation is [Co(tpfc)(pyr)$_2$] (121, Fig. 44; where tpfc = 5,10,15-tris(pentafluorophenyl)corrole and pyr = pyridine). Stability tests suggested that Co corrole 121 did operate through a homogeneous mechanism. The catalytic rate of H$_2$O oxidation for Co complex 121 was enhanced by increasing the phosphate concentration, indicative of the importance of proton transfer for O–O bond formation, which was also calculated to constitute the rate-determining step.

Co porphyrins have also been employed as WOCs and are depicted in Fig. 45. Sakai and co-workers recently reported that Co porphyrins 122–124 could function as active WOCs in photochemical oxidation of H$_2$O. Of the studied catalysts, Co porphyrin 124 was found to be the most efficient one (see Table 2). DLS measurements were also carried out to provide information whether nanoparticles are generated during the course of the catalysis. After the reactions, the solutions did not show any dispersion due to nanoparticle formation upon irradiation, which is in contrast to [Co(bpy)$_3$]$^{2+}$ that was shown to efficiently produce nanoparticles when irradiated. The second order catalyst dependence suggests a bimolecular radical coupling event as the rate-determining step. Two pathways for O–O bond formation were therefore proposed (Scheme 8), which featured either radical coupling of two formal Co$^{IV}$–oxyl or two Co$^{III}$–oxyl species, generated from the disproportionation of two Co$^{IV}$ species.

Although nanoparticle formation was not observed for Co porphyrins 122–124, the catalytic activity of these catalysts diminished significantly in subsequent runs after addition of fresh Na$_2$S$_2$O$_8$. The authors attributed this effect to decomposition of the Co porphyrins into catalytically less active or inactive entities. However, these entities must still contain the Co center; otherwise nanoparticles should have been observed. Mass spectrometry analysis of the reaction products revealed that oxidative cleavage of the porphyrin rings occurs at the meso positions, which produces species such as 134 (Fig. 46).

In a previous report, meso-substituted Co porphyrin complexes have been shown to react with singlet oxygen ($^1$O$_2$), which is a powerful oxidant produced during H$_2$O oxidation.

Table 2  Photochemical O$_2$ evolution by Co porphyrins 122–124$^a$

| Catalyst      | TON | TOF  |
|---------------|-----|------|
| Co porphyrin 122 | 88.7| 0.118|
| Co porphyrin 123 | 103.4| 0.138|
| Co porphyrin 124 | 121.8| 0.170|

$^a$ Photochemical experiments were carried out with a catalyst (10 μM), [Ru(bpy)$_3$](NO$_3$)$_2$ (1 mM) and Na$_2$S$_2$O$_8$ (5 mM) in aqueous phosphate buffer solutions (0.1 M, pH 11).
This observation suggests that the studied photosystem can undergo undesired side reactions with $^1$O$_2$, generated by the triplet state of either the Co-porphyrin or $\text{[Ru(bpy)$_3$]}^{2+}$. With this in mind, Sakai and co-workers examined the resistance of Co porphyrins towards $^1$O$_2$. By introducing fluorine substituents at the 2- and 6-positions of the porphyrin aryl units, to afford the fluorinated Co porphyrin 135 (Fig. 47), the attack of $^1$O$_2$ was effectively blocked. Comparing the fluorinated Co porphyrin complex 135 with the previously developed Co porphyrin 124 in light-sensitized H$_2$O oxidation confirmed that the fluorinated catalyst was more resistant towards decomposition, and operates without loss of catalytic activity. In contrast to the earlier Co porphyrin WOCs 122–124, porphyrin 135 exhibited a first order dependence on the catalyst concentration. This disparity implies that the rate-determining step for catalyst 135 is H$_2$O nucleophilic attack on a formal...
high-valent CoV species rather than oxyl coupling between two Co–oxyl units (Scheme 9). This study established the importance of resistance towards $^{1}$O$_2$ during light-driven H$_2$O oxidation and that the rational design of ligands can deliver dramatically improved WOCs.

Groves and Wang have also synthesized a series of cationic Co-based porphyrin complexes (125–127) for H$_2$O oxidation (Fig. 45). These porphyrin complexes were shown to mediate electrochemical H$_2$O oxidation, with Co porphyrin 125, housing the electron-deficient ligand, being the most efficient catalyst. The key species for these WOCs was proposed to be a CoIV–oxo species containing an oxidized radical porphyrin ligand ([[(L')CoIV–oxo]], which can be considered as a formal CoV–oxo species. Support of homogeneous O–O bond formation at a single metal center was also given, thus resembling the mechanistic pathway for the fluorinated Co porphyrin 135 (Scheme 9).

Additional mononuclear Co-based WOCs that have been proposed to promote light-sensitized H$_2$O oxidation through a homogeneous pathway are depicted in Fig. 48.
and include the $[\text{Co}^\text{II}(\text{qpy})(\text{OH}_2)_2]^{2+}$ complex 141 (qpy = 2,2':6,2''-quaterpyridine), the Co$^\text{II}$ salophen complex 142 and related salophen complexes, and the Co$^\text{II}$ complex $[\text{Co}(\text{tpa})\text{Cl}]^+$ 143 (tpa = tris(2-pyridylmethyl)amine). Several Co-based complexes with organic ligand frameworks have been shown to act as precatalysts to heterogeneous Co nanoparticles (Fig. 49). Although the initial well-defined Co complexes are transformed into heterogeneous materials, the carbonaceous residues originating from the ligand frameworks act as modifiers or capping agents of the generated nanoparticles. This indicates that the ligands might affect the structure and efficiency of the nanoparticle catalysts and suggests a ligand dependent route to efficient and robust catalytic materials, opening an appealing avenue for future research.

### 6.4. Dinuclear Co complexes as water oxidation catalysts

A relatively unexplored area is the development of dinuclear Co-based WOCs. Although efforts have been devoted to preparing dinuclear Co-based WOCs, only a few examples of active dinuclear Co catalysts exist. Two of these Co catalysts (148 and 149, Fig. 50) are based on the bridging bispyridylpyrazolate (bpp) ligand. The bpp ligand was chosen since it was considered to provide an environment for the two Co centers which is stable towards hydrolysis and ligand decomposition. Indeed, as confirmed by electrochemical analysis, the two Co-bpp complexes were able to evolve O$_2$ under acidic conditions. Another example of an active dinuclear Co WOC is $[\text{[tpa} \text{Co} - \mu - \text{OH}] \mu - \text{O}_2 \text{Co} - \text{tpa}]^{3+}$ (150), which catalyzes light-induced H$_2$O oxidation using $[\text{Ru(bpy)}_3]^{2+}$ as a photosensitizer and Na$_2$S$_2$O$_8$ as a sacrificial electron acceptor. The dinuclear $[\text{[tpa} \text{Co} - \mu - \text{OH}] \mu - \text{O}_2 \text{Co} - \text{tpa}]^{3+}$ complex 150 was obtained by treating the previously developed mononuclear $[\text{Co(tpa)}\text{Cl}]^+$ complex 143 with LiClO$_4$ and O$_2$, resulting in oxidation and formation of the Co$^{\text{III,III}}$ complex 150. A pH of 8 was found to be optimal for the light-driven O$_2$ evolution experiments, giving a TON of 58 with a TOF of $\sim$1.4 s$^{-1}$. The molecular integrity of the dinuclear Co complex 150 during catalysis was assessed by DLS and suggested that no Co oxide colloids were produced. The proposed catalytic cycle for the oxidation of H$_2$O by $[\text{[tpa} \text{Co} - \mu - \text{OH}] \mu - \text{O}_2 \text{Co} - \text{tpa}]^{3+}$ complex 150 is depicted in Scheme 10 and revolves around the generation of a (O)Co$^{IV}$-Co$^{IV}$(O) intermediate as the key species for intramolecular O–O bond formation. Further mechanistic investigations will likely result in a fundamental understanding of the catalytic process and enable the design of more efficient dinuclear Co-based WOCs.

![Fig. 49](Image)

**Fig. 49** Co-based complexes employed as precatalysts.

![Fig. 50](Image)

**Fig. 50** Structures of dinuclear Co-based WOCs 148–150.
7. Homogeneous Cu-based systems for catalytic water oxidation

Copper is a common catalyst for mediating aerobic oxidations using molecular oxygen as the terminal oxidant.\textsuperscript{298} However, only a handful of Cu-based catalysts are known to be capable of promoting the opposite reaction—H\textsubscript{2}O oxidation. This can perhaps be explained by the metal’s preference to carry out one-electron redox events rather than two-electron processes, which are at the core of H\textsubscript{2}O oxidation.

Early investigations on Cu-catalyzed H\textsubscript{2}O oxidation was conducted by Elizarova and co-workers.\textsuperscript{175} However, the first well-defined homogeneous Cu-based complex was reported by Mayer and co-workers in 2012.\textsuperscript{299} The authors examined the ability of the [Cu(bpy)(OH)\textsubscript{2}] complex \textsuperscript{156} (Fig. 51), a self-assembling Cu complex, to mediate electrochemical H\textsubscript{2}O oxidation. EPR measurements and control studies with Cu oxide implied that the [Cu(bpy)(OH)\textsubscript{2}] complex was homogeneous in nature. However, the stability of this Cu complex was severely hampered as it merely afforded a TON of \(\sim\)30, highlighting that more robust catalysts need to be designed in order to access more efficient Cu-based WOCs.

Lin and co-workers recently studied the catalytic activity of 4,4\textsuperscript{′}- and 6,6\textsuperscript{′}-substituted bipyridine-based Cu complexes.
authors found that Cu complex 157, \([\text{Cu(bpyOH)}(\text{OH})_2]\), housing the 6,6’dihydroxy-2,2’-bipyridine ligand provided an efficient WOC (Fig. 52).\(^\text{300}\) This ligand has previously been employed for other transformations, such as CO\(_2\) reduction\(^\text{301}\) and carboxylation reactions,\(^\text{302}\) where it was proposed to have a non-innocent role during the catalytic transformations. Lin and co-workers therefore utilized the 6,6’dihydroxy-2,2’-bipyridine ligand, envisioning that the ligand could be employed as a redox-active entity to modify the reactivity of the metal center to enhance the catalytic activity.\(^\text{300}\)

For Cu complex 157, O\(_2\) evolution was shown to occur in controlled potential electrolysis (CPE) with an overpotential of \(~640\) mV, giving a TON of \(~400\). The 4,4’dihydroxy-2,2’-bipyridine analogous Cu complex showed a significantly higher overpotential for H\(_2\)O oxidation than Cu complex 156. Both experimental and computational studies suggested the involvement of the ligand framework in electron transfer and proton transfer events, thus enhancing the catalytic activity of the Cu-based WOC.\(^\text{300}\) The chemistry of 6,6’-substituted 2,2’-bipyridine-based Cu complexes in H\(_2\)O oxidation catalysis has also been investigated by the group of Papish.\(^\text{303}\)

The group of Meyer has recently developed a self-assembling Cu tetrapeptide-based complex 158, Fig. 53.\(^\text{304}\) Cu complex 158 contains a triglycylglycine ligand (H\(_4\)tgg), which creates a suitable environment for the coordination of Cu.\(^\text{305,306}\) At pH 11, CPE showed that the catalytic current was maintained for \(~5\) h, resulting in a TON of \(~13\) based on the initial amount of Cu complex 158 in solution. Several lines of evidence supported a homogeneous mechanism: (1) no spectroscopic change of the electrolysis solution was observed, during CPE only minor changes in the peak current or wave shape were observed, (2) an electrode subjected to CPE at 1.32 V vs. NHE with Cu catalyst 158 resulted in no catalytic response when subjected to a fresh catalyst-free electrolyte solution, and (3) no precipitation or film generation was observed by SEM and XPS.\(^\text{304}\)

The catalytic peak current for H\(_2\)O oxidation was found to vary linearly with the concentration of Cu complex 158, suggesting a mechanism involving O–O bond formation at a single-site Cu center. The proposed mechanism for oxidation of H\(_2\)O by Cu complex 158 is shown in Scheme 11 and involves oxidation to produce a Cu\(^{\text{III}}\) species. A subsequent oxidation event furnishes a Cu\(^{\text{IV}}\)-oxo (or a Cu\(^{\text{III}}\)-oxyl) species, which is believed to be responsible for mediating O–O bond formation. The produced hydroperoxide species, Cu\(^{\text{III}}\)-OOH, is subsequently oxidized to Cu\(^{\text{III}}\)-OO’ from which O\(_2\) is liberated, regenerating the Cu\(^{\text{II}}\) catalyst 158 and closing the catalytic cycle.\(^\text{304,307}\) The two related Cu tetrapeptide complexes 159 and 160 (Fig. 53) have also recently been synthesized and studied in H\(_2\)O oxidation.\(^\text{308}\)

Llobet and co-workers recently prepared a family of Cu complexes (161–164) based on tetra-anionic tetradentate amidate ligands.\(^\text{309}\) The prepared Cu\(^{\text{II}}\) complexes (Fig. 54) were found to be four-coordinate with a square planar geometry. Electrochemical measurements for the Cu\(^{\text{III}}\) complexes revealed a significant electrocatalytic current corresponding to H\(_2\)O oxidation together with two redox waves. The first quasi-reversible wave was assigned to a one-electron process, generating a Cu\(^{\text{III}}\) species. A subsequent oxidation of H\(_2\)O by Cu complex 164 involves initial formation of the \([\text{Cu}^{\text{III}}(\text{OH})]\)\(^\text{2+}\) species, which subsequently reacts with OH\(^-\) to produce Cu\(^{\text{III}}\)-peroxo species, \([\text{Cu}^{\text{III}}(\text{HOOH})]\)\(^2+\). This species undergoes a metal-based oxidation to form \([\text{Cu}^{\text{III}}(\text{HOOH})]^-\). A final proton-coupled oxidation gives a \([\text{Cu}^{\text{III}}(\text{HOO})^-]\) species from which O\(_2\) is liberated, thus regenerating the start-
The use of tailored ligands which are able to donate electrons—being non-innocent—during the catalytic oxidation process can be essential for designing novel molecular WOCs where this cooperative effect can alleviate the metal center from being too highly oxidized during the catalytic process.310

The Cu-based WOCs described thus far require alkaline conditions in order to mediate electrochemical oxidation of H2O. However, the dinuclear Cu2II,II2 complex (Fig. 55) based on the 2,7-[bis(2-pyridylmethyl)aminomethyl]-1,8-naphthyridine ligand (bpman) ligand has been reported to operate under neutral conditions.311 Reacting the dinucleating bpman ligand312,313 with 2 equivalents of Cu(CF3SO3)2 produced the CuII,II2 complex in 71% yield. Electrochemical measurements of Cu complex 165 showed an onset potential for catalytic H2O oxidation starting at ∼1.6 V vs. NHE. The catalytic activity of the dinuclear Cu complex 165 was assessed by CPE at 1.87 V vs. NHE to give a TOF of ∼0.6 s−1 with a Faradaic efficiency of ∼98%. Computational studies suggested that O–O bond formation proceeded through a cooperative interaction between the two CuIII centers rather than by the formation of a CuIV=O unit in which a CuIII–O(H) moiety couples with a

Scheme 11 Proposed mechanism for H2O oxidation catalyzed by Cu complex 158.

Fig. 54 CuII complexes 161–164 based on tetradeutate amidate ligands.

Fig. 55 Structure of the dinuclear CuII,II2 complex [CuII,II2(bpman)(μ-OH)3]+ (165). bpman = 2,7-[bis(2-pyridylmethyl)aminomethyl]-1,8-naphthyridine.
μ-oxo unit. The proposed O−O bond forming step is different from the previously reported Cu-based WOCs and could provide new routes for the activation of H$_2$O.

Ding and co-workers recently reported that the Cu-containing POM $[\text{Cu}_5\text{(OH)}_4\text{(OH}_2)_2\text{(A-α-SiW}_9\text{O}_{33})_2]^{10–}$ (166) functions as a catalyst for visible light-driven water oxidation. The Cu-substituted POM 166 (Fig. 56) was originally developed by Kortz and co-workers and consists of two A-α-[SiW$_9$O$_{34}$]$^{10–}$ Keggin moieties that are linked together through two W−O−W bonds and stabilized by a central $[\text{Cu}_5\text{(OH)}_4\text{(OH}_2)_2]^{20+}$ unit. Of the evaluated Cu POMs, POM 166 was the only complex that exhibited water oxidation activity. Using a system consisting of Cu POM 166, [Ru(bpy)$_3$]$^{2+}$ as the photosensitizer and S$_2$O$_8^{2–}$ as the sacrificial electron acceptor afforded a TON of 91 under optimized conditions.

Several molecular Cu-based complexes have been reported to act as precursors to catalytically active heterogeneous materials, highlighting the importance of determining the true nature of the catalytic entity when studying Cu-based WOCs. For Cu POM 166, multiple experiments, including DLS, UV/vis spectroscopy and nanosecond laser flash photolysis experiments, suggest that Cu POM 166 is the dominant species under the studied catalytic conditions and that it operates through a homogeneous mechanism.

Although relatively few water oxidation catalysts based on first-row transition metals had been reported before the early 2000s, recent years have seen a dramatic increase in the number of such catalysts. However, a majority of these catalysts require two-electron oxo-transfer oxidants to drive H$_2$O oxidation. Despite considerable progress in recent years, a limiting feature encountered with these first-row transition metal WOCs is their relative lability compared to catalysts based on second- and third-row transition metals, such as Ru and Ir. Due to the highly oxidizing conditions required to oxidize H$_2$O, the rational design of robust and efficient WOCs based on first-row transition metals still remains a crucial challenge.

A common topic in the discussion of molecular earth-abundant metal WOCs is the nature of the real catalytic entity—their homogeneity. Their propensity to form metal oxide nanoparticles under the catalytic conditions is a feature that requires particular attention. The generation of heterogeneous materials from the initially well-defined metal complexes is facile and highly dependent on the reaction conditions where small changes can affect both the mechanistic pathway and the stability of the examined metal complex.

The structural knowledge gained from studying the Mn$_4$Ca cluster in the OEC has inspired researchers to design small metal-based model clusters of various shapes and nuclearities. These seminal studies have afforded considerable insight for the rational synthesis of closely related artificial metal-based cubane mimics of the natural photosynthetic system. For a long time, Nature has been a great source of inspiration in the development of molecular photosynthetic mimics for solar energy to fuel conversion and will certainly continue to stimulate the design of molecular mimics based on earth-abundant metals.

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