Review

Reactive Cobalt–Oxo Complexes of Tetrapyrrolic Macrocycles and \( N \)-based Ligand in Oxidative Transformation Reactions

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Received: 28 November 2018; Accepted: 25 December 2018; Published: 26 December 2018

Abstract: High-valent cobalt–oxo complexes are reactive transient intermediates in a number of oxidative transformation processes e.g., water oxidation and oxygen atom transfer reactions. Studies of cobalt–oxo complexes are very important for understanding the mechanism of the oxygen evolution center in natural photosynthesis, and helpful to replicate enzyme catalysis in artificial systems. This review summarizes the development of identification of high-valent cobalt–oxo species of tetrapyrrolic macrocycles and \( N \)-based ligands in oxidation of organic substrates, water oxidation reaction and in the preparation of cobalt–oxo complexes.

Keywords: cobalt–oxo complex; tetrapyrrolic macrocycles; \( N \)-based ligand; identification

1. Introduction

In biological systems, metalloenzymes, typically containing Mn, Fe and Cu centers, are known to catalyze a wide range of reactions including aliphatic and aromatic C–H hydroxylation, epoxidation, desaturation, and heteroatom (S, N or O) dealkylation or oxidation [1,2]. It is well known that iron-oxo species are the reactive oxidants in the catalytic cycle of heme [3] and non-heme iron enzymes [4]. Similarly, manganese–oxo complex has been suggested the key intermediate in oxygen-evolving center of photo-system II (PSII) [5–7]. The transition metal–oxo complexes of iron and manganese involved in artificial oxygen transfer and C–H bond activations reactions have been extensively reviewed [8–13]. Except for the early transition metal–oxo complexes, high-valent metal–oxo complexes of late transition metals, particularly cobalt–oxo complexes, are also highly reactive transient intermediates in cobalt-catalyzed C-H bond activation and O-O bond formation reactions [14–16], and they are considered to be more reactive then related iron-oxo species due to a weak metal–oxygen bond [17,18]. Currently, clean energy production by maneuvering natural photosynthesis in water oxidation reactions to develop artificial photosynthesis [19–21] for efficient water splitting is a hot topic of research [22–24]. In particular, the cobalt oxides are often used materials for water oxidation to generate molecular oxygen [25–28]. The high-valent cobalt–oxo complexes of \( N \)-based ligand can be implicated as reactive species in the O–O bond-forming event during water oxidation [29,30]. Furthermore, cobalt complexes based on tetrapyrrolic macrocycles are often used in mimicking the peroxidase-like activity for the selective oxidation of organic substrates via high-valent cobalt(IV)–oxo intermediates [31,32]. Obviously, in the study of the reactive oxidants in these catalytic reactions it is essential to provide insight into their mechanism of reaction, allowing us to probe the critical step in these challenging reactions. However, the isolation and identification of these transient intermediates is considerable challenge. The cobalt–oxo complexes are not stable because cobalt has large number of
d-electrons which produces strong electronic repulsion between the rich electron oxygen and cobalt center. Also, the strong oxidative environment will cause oxidative degradation of the ligand, making the high-valent cobalt–oxo complexes unstable. To reduce the electronic repulsion between cobalt and oxygen, and to avoid oxidative degradation, tetrapyrrolic macrocycles and N-based ligands with a different electronic environment were implemented to increase the stability of cobalt–oxo complexes (Figure 1). Mostly, these can only be characterized in situ by electron paramagnetic resonance (EPR) [33], X-ray absorption [34] and time-resolved Fourier-transform infrared (FT-IR) [16] spectroscopic methods. Computational studies were also carried out to understand the nature of species involved in water oxidation [35]. Recently, the isolation and/or identification of high-valent cobalt–oxo complexes has become a key topic in order to develop and understand the mechanism of artificial photosynthesis and to replicate enzymatic process in artificial reactions.

This review comprehends the high-valent cobalt–oxo complexes of tetrapyrrolic macrocycles and N-based ligands reported to date, along with outlooks in this intriguing research area. It has been divided into three sections: identification of cobalt–oxo species involved in oxidation of organic substrates; identification of cobalt–oxo species involved in heterogeneous and homogeneous water oxidation reactions; and preparation of high-valent cobalt–oxo complexes.

2. Cobalt–Oxo Species Involved in Oxidation of Organic Substrates

Cobalt–oxo species are involved in many of oxidative and C-H bond activation reactions. The ligands used to generate cobalt–oxo species play a key role in stabilizing cobalt–oxo species. Also, to mimic the enzymes-like environment, different types of support are used as protein backbone for example cellulosic fiber and multiwall carbon nanotubes. These supports cannot alter the reaction mechanism however, precisely control the generation of reactive intermediate, which also determines the activity, durability and stability of the complexes [36–38].

Nam et al. reported [39] the catalytic oxidation of alkene and alkane using cobalt-substituted polyoxotungstate and employed different oxidants such as iodosylbenzene, potassium monopersulfate and m-CPBA. Cobalt-substituted polyoxotungstate was proved to be a good catalyst. They proposed the involvement of different cobalt–oxo species with the different oxidants. Two possible species may form with iodosylbenzene, high-valent cobalt(V)–oxo 1 and cobalt–iodosylbenzene adduct 2.

Figure 1. Some of the most used tetrapyrrolic macrocycles and N-based ligands used to stabilize high-valent cobalt–oxo complexes.
They suggested that complex 2 is responsible for oxygen transfer because cobalt cannot be obtained in +5 oxidation state. H2O18O and m-CPBA predict the involvement of cobalt(III)–oxygen adducts as oxygen transfer complex. Isotopically labeled water (H218O) is a useful experimental tool to investigate the involvement of high-valent cobalt–oxygen species in cobalt-mediated oxygen atom transfer reactions, but all the attempts to obtain 18O-labeled products have failed. Furthermore, porphyrins are extensively used to get stable metal–oxygen complexes [13]. Therefore, porphyrins with a different electronic environment were used to stabilize cobalt–oxy species [40,41]. Cobalt(IV)–oxy [40] and cobalt(IV)–oxy porphyrin radical [41] were proposed to be involved in C-H bond activation reaction. These species are quite reactive towards the oxidation of alkane and alcohol, respectively. However, there is no experimental evidence to support presence of cobalt–oxy species due to instability. Likewise, a cobalt(IV)–oxy species was reported [42], based on the tetraanionic cobalt(II) complex of (8HBA-Et)H4, N,N′-(ethane-1,2-diyl)bis(5-bromo-2-hydroxybenzamide), that provides a strong ligand field. Consequently, this specie was stable enough to be characterized by EPR and ESI-MS spectroscopy analysis. Also, the presence of high-valent cobalt(IV)–oxy porphyrin was reported during the oxidation of alcohol to benzaldehyde by molecular oxygen in the presence of isobutyraldehyde, using bifunctional hybrid catalyst originated from cobalt tetra(4-sulfonatophenyl)porphyrinate anion [43] and a cationic meso-tetakis (1-methyl-4-pyridyl) cobalt porphyrin immobilized in montmorillonite interlayers [44]. The presence of a cobalt(IV)–oxy specie was predicted by an 18O-labeled experiment of product [43]. The turnover frequency and catalytic yield was higher in the prior case. Later, the cobalt(IV)–oxy porphyrin was generated [45] by the oxidation of cobalt porphyrin Co(TPFPP)(CF3SO3) utilizing m-CPBA as oxidant in solvent mixture of CH3CN and CH2Cl2. Incorporation of H218O in the catalytic oxidation demonstrated the presence of 18O-labeled alcohol in the product, which is evidence for the presence of cobalt–oxy species. Furthermore, cobalt(V)=O and cobalt(IV)=O were generated [46] by the oxidation of a mononuclear non haem cobalt(III) [Co((bpc)Cl2][Et4N] (H2bpc=4,5-dichloro-1,2-bis(2-pyridine-2-carboxamido)benzene) complex of a tetradentate ligand containing two deprotonated amide moieties with PhIO. Oxidation of the cobalt(III) complex generated cobalt acylperoxo intermediate, which on the heterolytic and homolytic cleavage of O-O bond generated respective cobalt(V)=oxo and cobalt(IV)=oxy species. These species are also not enough stable to be characterized spectroscopically. Similarly, a cobalt(IV)=O specie based on isindole-core ligand was proposed [47] as a reactive intermediate, during the stereoselective oxidation of alkane using m-CPBA as oxidant. The kinetic isotopic effect and 18O-labeled experiment predict the involvement of cobalt–oxy species. Recently, the involvement of a high-valent cobalt(IV)=O radical cation was proposed [48] during the reduction of O2. The dianionic pentadentate ligand system based on bis-pyrazolyl diaryl borate arms attached to a 2,6-substituted pyridyl frame was used to stabilize the cobalt(IV)=oxy intermediate. Cobalt(IV)=O radical cation was generated by the cleavage of Co-O bond, and examined theoretically and experimentally. A density functional theory (DFT) calculation suggests the presence of maximum electron density at oxygen 70% with Co-O bond length of 1.67 Å.

(Scheme 1). They suggested that complex 2 is responsible for oxygen transfer because cobalt cannot be obtained in +5 oxidation state. H2O18O and m-CPBA predict the involvement of cobalt(III)–oxygen adducts as oxygen transfer complex. Isotopically labeled water (H218O) is a useful experimental tool to investigate the involvement of high-valent cobalt–oxy species in cobalt-mediated oxygen atom transfer reactions, but all the attempts to obtain 18O-labeled products have failed. Furthermore, porphyrins are extensively used to get stable metal–oxygen complexes [13]. Therefore, porphyrins with a different electronic environment were used to stabilize cobalt–oxy species [40,41]. Cobalt(IV)–oxy [40] and cobalt(IV)–oxy porphyrin radical [41] were proposed to be involved in C-H bond activation reaction. These species are quite reactive towards the oxidation of alkane and alcohol, respectively. However, there is no experimental evidence to support presence of cobalt–oxy species due to instability. Likewise, a cobalt(IV)–oxy species was reported [42], based on the tetraanionic cobalt(II) complex of (8HBA-Et)H4, N,N′-(ethane-1,2-diyl)bis(5-bromo-2-hydroxybenzamide), that provides a strong ligand field. Consequently, this specie was stable enough to be characterized by EPR and ESI-MS spectroscopy analysis. Also, the presence of high-valent cobalt(IV)–oxy porphyrin was reported during the oxidation of alcohol to benzaldehyde by molecular oxygen in the presence of isobutyraldehyde, using bifunctional hybrid catalyst originated from cobalt tetra(4-sulfonatophenyl)porphyrinate anion [43] and a cationic meso-tetakis (1-methyl-4-pyridyl) cobalt porphyrin immobilized in montmorillonite interlayers [44]. The presence of a cobalt(IV)–oxy specie was predicted by an 18O-labeled experiment of product [43]. The turnover frequency and catalytic yield was higher in the prior case. Later, the cobalt(IV)–oxy porphyrin was generated [45] by the oxidation of cobalt porphyrin Co(TPFPP)(CF3SO3) utilizing m-CPBA as oxidant in solvent mixture of CH3CN and CH2Cl2. Incorporation of H218O in the catalytic oxidation demonstrated the presence of 18O-labeled alcohol in the product, which is evidence for the presence of cobalt–oxy species. Furthermore, cobalt(V)=O and cobalt(IV)=O were generated [46] by the oxidation of a mononuclear non haem cobalt(III) [Co((bpc)Cl2][Et4N] (H2bpc=4,5-dichloro-1,2-bis(2-pyridine-2-carboxamido)benzene) complex of a tetradentate ligand containing two deprotonated amide moieties with PhIO. Oxidation of the cobalt(III) complex generated cobalt acylperoxo intermediate, which on the heterolytic and homolytic cleavage of O-O bond generated respective cobalt(V)=oxo and cobalt(IV)=oxy species. These species are also not enough stable to be characterized spectroscopically. Similarly, a cobalt(IV)=O specie based on isindole-core ligand was proposed [47] as a reactive intermediate, during the stereoselective oxidation of alkane using m-CPBA as oxidant. The kinetic isotopic effect and 18O-labeled experiment predict the involvement of cobalt–oxy species. Recently, the involvement of a high-valent cobalt(IV)=O radical cation was proposed [48] during the reduction of O2. The dianionic pentadentate ligand system based on bis-pyrazolyl diaryl borate arms attached to a 2,6-substituted pyridyl frame was used to stabilize the cobalt(IV)=oxy intermediate. Cobalt(IV)=O radical cation was generated by the cleavage of Co-O bond, and examined theoretically and experimentally. A density functional theory (DFT) calculation suggests the presence of maximum electron density at oxygen 70% with Co-O bond length of 1.67 Å.

Scheme 1. Proposed mechanism for oxidation of alkene by cobalt polyoxotungstate using PhIO as oxidant [39].
Moreover, to mimic the enzyme activity for controllable catalytic oxidation, researchers made extensive efforts to develop and discover functional materials having properties intrinsic to enzymes. Many transition-metal complexes were prepared [49–51] to mimic the expected features of enzymes, such as selectivity and steric accessibility, but these do not present the said features due to the non-natural environment. A catalyst which is a replication of enzyme should possess a suitable cavity or cleft for accessibility of substrates and introduction of functional groups that act as active sites within the cavity [52,53]. Enzymatically inspired catalytic system was prepared by using cobalt tetraaminophthalocyanine (CoTAPc) as a catalyst supported by ordered-mesoporous-carbon (OMC) for controllable activation of hydrogen oxide (H₂O₂) to generate stable cobalt–oxo intermediate [32]. Ordered-mesoporous-carbon provides the steric environment for a substrate to attach with active sites and protects the active sites against the external interface. However, a disadvantage of hydrogen peroxide is the formation of hydroxyl radical that is highly reactive, so it decreases the selectivity. A fifth ligand dodecylbenzenesulfonate (LAS) is employed to inhibit the production of hydroxyl radical. This fifth ligand also helps to generate high-valent cobalt(IV)–oxo specie by heterolytic cleavage of peroxide O-O bond. The involvement of cobalt–oxo specie was corroborated by the results of semiempirical quantum-chemical PM6 calculations. Similarly, a modification in the tetrapyrrolic macrocycle of cobalt tetraaminophthalocyanine (CoTAPc) was made by the attachment of epoxy compound 2,3-epoxypropyl triethylammonium chloride (EPTAC), to obtain a new catalyst with positively charged quaternary ammonium salt chain (OMC-CoTAPc-EPTAC) [31]. The modified catalyst displays high catalytic activity especially for negatively charged substrates. The free radical trapping EPR analysis using 5,5-Dimethyl-1-pyrroline N-oxide (DMPO) as a free radical scavenger did not detect DMPO-·OH and DMPO-·OOH signal, ruling out the free radical type mechanism. That is why, the cobalt(IV)–oxo complex was proposed as a reactive intermediate due to the heterolytic cleavage of O-O bond of peroxide. Moreover, cellulose fiber could play the role of the protein backbone in enzymes, providing an enzyme-like environment with enhanced regioselectivity to remove organic dyes and improve the stability of intermediate generated. A catalyst was developed based on cellulose fiber-bonded cobalt phthalocyanine catalytic entity to activate hydrogen peroxide in order to generate cobalt–oxo specie [54]. The reaction channel was controlled by linear alkylbenzene sulfonate (LAS). High-valent cobalt(IV)–oxo specie 4 was generated by the heterolytic cleavage of peroxide O-O bond and homolytic cleavage generate cobalt(III)–oxo specie 3 (Figure 2).

Figure 2. Possible pathway for the formation of active species in cellulosic fiber-bonded cobalt phthalocyanine (CoPc) H₂O₂ system. (A) Generation of hydroxyl radicals without ligand dodecylbenzenesulfonate (LAS) by the homolytic cleavage of the peroxide O-O bond; (B) Generation of cobalt–oxo with LAS by the heterolytic cleavage of the peroxide O-O bond [54].
The in situ X-band EPR analysis was conducted at room temperature in the presence of LAS demonstrating a signal at \( g_{\text{eff}} = 2.099 \) identifying the presence of Co\(^{IV} \) with spin state \( S = \frac{1}{2} \). Usually, metal–oxo species have been detected at low temperature \( [55,56] \); a high oxidation state of \( \text{PcCo}^{IV}=\text{O} \) was observed at room temperature, presenting high stability of complex, auto-oxidation protected by cellulose matrix. Cellulosic fiber bonded cobalt phthalocyanine (CoPc) can also activate peroxymonosulfate \([57]\). The oxidation activity of catalyst is remarkably enhanced in the presence of bicarbonate ion \( (\text{HCO}_3^-) \) due to the generation of \( (\text{PcCo}^{IV}=\text{O}) \) by the heterolytic cleavage of O–O bond of peroxymonosulfate. Later, the same group \([58]\) employed the multiwall carbon nanotubes (MWCNTs) as protein-like backbone anchored on cobalt phthalocyanine (CoPc) for peroxidase-like activation of hydrogen peroxide. The anchoring of catalytic entity on MWCNTs decreases the diffusional mass transfer process (DMTP) and enhances the resistance of CoPc-MWCNTs oxidative decay. The introduction of linear alkylbenzene sulfonates (LAS) facilitates the heterolytic cleavage of O–O bond of peroxide to generate cobalt(IV)–oxo species. Furthermore, pyridine functionalized MWCNTs were produced and axially coordinated on the cobalt phthalocyanine (CoPc), generating a catalyst with increased catalytic activity and stability \([59]\). The heterolytic cleavage of O–O bond of hydrogen peroxide to produce cobalt(IV)–oxo without presence of any fifth ligand. The high-valent cobalt(IV)–oxo was analyzed by in situ ESI-MS and density functional theory. The DFT calculated bond length of Co–O bond is 1.806 Å and unpaired electron spin populations are mainly on the oxygen. Cobalt(IV)–oxo 5 was generated at pH = 10, the catalytic cycle starts with the coordination of \( \text{OOH}^- \) with \( \text{Co}^{II} \), and the heterolytic cleavage of O–O occurs with the release of \( \text{OH}^- \) (Figure 3a). Moreover, non-haem cobalt(III) oxamate anion could also be used to stabilize high-valent cobalt(IV)–oxo species \([60]\). The oxidation of industrial contaminants was reported \([61]\) by immobilizing non haem cobalt(III) complex \([\text{Co}^{III}(\text{opbaX})]^- (\text{opbaX} = 4-X\text{-o-phenylenebis(oxamate)}) \) on pyridine-modified MWCNTs, where pyridine acts as a fifth ligand. Similarly, cobalt(III) complexes of \([\text{Co}^{III}(\text{opbaX})]^- (\text{opbaX} = 4-X\text{-o-phenylenebis(oxamate)}, X = \text{H, NO}_2, \text{CH}_3) \) with different substituents was reported \([62]\) for accelerating heterolytic cleavage of hydrogen peroxide to imitate the essential and general principle of natural enzymes without using any fifth ligand (Figure 3b). An ESI-MS and EPR trapping technique revealed the presence of cobalt(IV)–oxo reactive specie. The generation of cobalt–oxo species depends on the electronic environment of substituent. In Scheme 2 pathway (b) the electron rich cobalt complex favors the homolytical cleavage of the hydroperoxide O–O bond while the electron deficiency favors the heterolytic cleavage with generation of \( \text{Co}^{IV}=\text{O} \). The tendency of generation of \( \text{Co}^{IV}=\text{O} \) was in order \( 8 > 7 > 9 \). Density functional theory also demonstrated that electron withdrawing group helps in pulling electron and lowering the corresponding energy levels. Keeping in mind the concept of the “oxo wall” \([63]\), another pathway (a) also proposed, heterolytic cleavage of O–O generate the ligand based radical intermediate \( \text{OH}^-\text{Co}^{III}(\text{opbaX}) \), in which ligand transfers one electron to cobalt and cobalt transfers it to oxygen.

Our group recently \([64]\) reported the catalytic oxidation of alkene using four cobalt(III) corroles of different electronic environment \( \text{F}_0\text{C-Co}, \text{F}_5\text{C-Co}, \text{F}_{10}\text{C-Co} \) and \( \text{F}_{15}\text{C-Co} \) employing various oxidants. The in situ ESI HR-MS analysis of styrene oxidation with \( \text{KHSO}_5 \) predicts the presence of high-valent cobalt(V)–oxo complex as active intermediate. The in situ X-band CW EPR analysis revealed a signal at \( g = 2.0135 \) for the presence of cobalt–oxo specie.
was carried out in the presence of photosensitizer [Ru(bpy)₃]²⁺ (bpy = 2,2′-bipyridine) that creates a hole and S₂O₈²⁻ as an electron acceptor. The FT–IR characterization revealed the involvement of two intermediates with absorption band at 1013 cm⁻¹ and 840 cm⁻¹. The band at 1013 cm⁻¹ was assigned to Co(III)OO (fast site) group with a neighboring hydroxyl group. Incorporation of H₂¹⁸O shifts the peaks at 995 cm⁻¹ and 966 cm⁻¹. The shifting of frequency agrees well with the presence of superoxide moiety on a metal-oxide surface [72,73]. The superoxide surface intermediate causes the three-electron water oxidation. The band at 840 cm⁻¹ was assigned to Co⁴⁺=O (slow site) surface species. No change in the spectrum was observed by the incorporation of H₂¹⁸O, ruling out the presence of any peroxide intermediate. From the mechanistic point of view, Co⁴⁺=O was

3. Cobalt–Oxo Species Involved in Water Oxidation Reaction

Water oxidation is a process that involved the four-electron-four-proton oxidation of water to evolve O₂. In natural photosynthesis, sunlight is converted to chemical energy by the oxidation of water [20]. As a consequence, understanding nature’s water oxidation mechanism in photosystem II has been the focus of research for the development of artificial water oxidation catalyst. The development of efficient water oxidation catalysts with minimal cost is a challenge [65–68]. Various water oxidation catalysts were developed to understand the O-O bond formation event in natural photosynthesis to evolve oxygen. Cobalt is the most abundant and cheap earth metal. Cobalt oxide materials are among the most promising catalyst for water oxidation [69–71] and cobalt–oxo species depend on the electronic environment of substituent. In Scheme 2 pathway (b) the electron richness favors the homolytical cleavage of the hydroperoxide O−O bond while the electron deficiency favors the heterolytic cleavage with generation of (Co IV=O) species. A high oxidation state of PcCoIV=O has been detected at low temperature [55,56]; a high oxidation state of PcCoIV=O corresponds to 7H+/3e− process. Further, oxidation of 15 afforded 16. A key step to evolve oxygen. Cobalt is the most abundant and cheap earth metal. Cobalt oxide materials are among the most promising catalyst for water oxidation [69–71] and cobalt–oxo species are involved in the O-O forming event of water oxidation.

Frei et al. reported [16] the photocatalytic water oxidation using cobalt oxide. The water oxidation was carried out in the presence of photosensitizer [Ru(bpy)₃]²⁺ (bpy = 2,2′-bipyridine) that creates a hole and S₂O₈²⁻ as an electron acceptor. The FT–IR characterization revealed the involvement of two intermediates with absorption band at 1013 cm⁻¹ and 840 cm⁻¹. The band at 1013 cm⁻¹ was assigned to Co(III)OO (fast site) group with a neighboring hydroxyl group. Incorporation of H₂¹⁸O shifts the peaks at 995 cm⁻¹ and 966 cm⁻¹. The shifting of frequency agrees well with the presence of superoxide moiety on a metal-oxide surface [72,73]. The superoxide surface intermediate causes the three-electron water oxidation. The band at 840 cm⁻¹ was assigned to Co⁴⁺=O (slow site) surface species. No change in the spectrum was observed by the incorporation of H₂¹⁸O, ruling out the presence of any peroxide intermediate. From the mechanistic point of view, Co⁴⁺=O was
generated by the oxidation of surface group Co(III)-OH. At the fast site catalytic species, catalytic turn
over frequency is at least 10 times more than slow site catalytic species, because it has no neighbor
hydroxyl group. Furthermore, Stahl et al. reported [74] the water oxidation employing cobalt oxide as
an electrocatalyst, and proposed the involvement of (CoIV-O) as reactive specie. The EPR analysis with
signals at g-values 2.59, 2.17 and 1.99 revealed the presence of multiple paramagnetic species during
water oxidation, possibly arising from (CoIV-O) sites in the catalyst with a different coordination
environment. The mechanism of water oxidation is pH dependent, at acidic pH homogeneous
catalysis leading to H2O2 production, while at pH above 3.5 heterogeneous catalysis takes place,
generating O2 from four-electron water oxidation (Scheme 3). The oxidation of 12 produced 14
(12→13→14) corresponds to a 3H+/e− process. Subsequently, 1e− oxidation generated specie 15
corresponding to 7H+/3e− process. Further, oxidation of 15 afforded 16. A key step to evolve oxygen
is the nucleophilic attack of water at 16 to produce 17 [75–77]. Under the acidic pH, the PCET-mediated
formation of 11 was prevented (Scheme 3). The oxidation of 10 produced 18 that dissolve from surface.
The intermediate specie 18 invoked the homogeneous oxidation of water to H2O2 [78]. Similarly,
bringing cobalt(IV)-oxo [79] and terminal cobalt(IV)-oxo radical [80] species were proposed as reactive
catalytic sites for water oxidation, employing amorphous cobalt oxide. X-ray absorption near the edge
provide the insight that the generation of high-valent (CoIV-O) depended on the potential applied
and pH. The edge position of the spectra was taken at pH = 7 and pH = 9 differs by about 1.0 eV
by keeping potential constant at 0.95 V, and edge position of the spectra were taken at pH = 7 by
increasing electrode potential from 0.95 to 1.34 V differs by about 1.2 eV [79]. However, the study
of cobalt–oxo species involved in water oxidation was difficult because in oxygen evolving catalysis
(OEC), large number of spectroscopically active backgrounds species are present which limits their
detection and characterization.

Scheme 3. Mechanism for oxidation of water by cobalt oxide under acidic and basic conditions [74].

Consequently, suitable catalysts to reduce background active species are needed to design.
N-based ligands have attractive properties to be used as homogenous molecular water oxidation
catalysts [81]. Recently, a significant number of catalysts are developed based on single site and
multinuclear transition metal including Mn, Fe, Co, Cu, Ru and Ir [82–87]. The biggest challenge is to
find a suitable coordination environment because the metal–ligand bond opposite a metal–oxygen bond
can be compromised at higher redox level leading the catalyst to be susceptible to degradation [88].
So, single site N-based ligand homogeneous catalysts of cobalt were developed utilizing stable
pentadentate ligand environment of 2,6-(bis(bis-2-pyridyl)methoxy-methane)-pyridine [89] and
6-(bis(bis-2-pyridyl)-methoxy methane)pyridine [30] for water oxidation. The electrochemical studies
revealed that over pH range 7.6–10.3 an oxidation event was observed at +1.43 V vs. NHE corresponding
to $[\text{Co}^{IV}\cdot\cdot\text{OH}]^{3+}/[\text{Co}^{III}\cdot\cdot\cdot\text{OH}]^{2+}$ with significant rise in current. This signal is not classified as PCET because $E_{1/2}$ is static over this pH range. A pH dependent step was observed at pH $> 10.3$ corresponding to $[\text{Co}^{IV}\cdot\cdot\cdot\text{O}]^{2+}/[\text{Co}^{III}\cdot\cdot\cdot\text{OH}]^{2+}$ which is consistent with PCET. High-valent $[\text{Co}^{IV}\cdot\cdot\cdot\text{O}]^{2+}$ species evolves $\text{O}_2$ by the nucleophilic attack of $\text{H}_2\text{O}$ \cite{89,90}. An alternative pathway proposed that the attack of $\text{OH}^-$ at $[\text{Co}^{IV}\cdot\cdot\cdot\text{OH}]^{3+}$ in the rate determining step will evolve $\text{O}_2$ \cite{30}. Likewise, $[\text{Co}^{IV}\cdot\cdot\cdot\text{O}]$ specie was proposed as reactive intermediate during water oxidation at basic pH using cobalt-porphyrins as catalyst \cite{91}. Similarly, Groves and Wang reported \cite{92} the single site homogeneous water oxidation catalyst, employing a series of cobalt porphyrins \cite{19,20 and 21 (Scheme 4)}. A high-valent $\text{Co}^{IV}\cdot\cdot\cdot\text{porphyrin cation}$ radical acts as reactive intermediate. The electrochemical experiment provides the evidence for the formation of high-valent $\text{Co}^{IV}\cdot\cdot\cdot\text{O}$ specie. The redox event at 250 mV vs. Ag/Cl reference represents the resting state of catalyst $\text{H}_2\text{O}\cdot\cdot\cdot\text{Co}^{III}\cdot\cdot\cdot\text{OH}$ \cite{23}. The observed anodic features at $\sim 1$ V demonstrates the oxidation of $\text{Co}^{III}$ porphyrin to $\text{Co}^{III}$ porphyrin radical cation ($+\text{P}\cdot\cdot\cdot\text{Co}^{III}\cdot\cdot\cdot\text{OH}$) \cite{24}. As first oxidation occurred before the onset potential of WOC catalytic current, so $+\text{P}\cdot\cdot\cdot\text{Co}^{III}\cdot\cdot\cdot\text{OH}$ is not the reactive oxidant in this system. The second oxidation at 1320 mV generates a reactive high-valent $\text{Co}^{IV}\cdot\cdot\cdot\text{O}$ porphyrin radical cation \cite{25}. The key step for O-O bond formation is the nucleophilic addition of $\text{H}_2\text{O}$ to $+\text{P}\cdot\cdot\cdot\text{Co}^{IV}\cdot\cdot\cdot\text{O}$ \cite{25} to form $\text{Co-hydroperoxo}$ or peroxo which further oxidized to evolve $\text{O}_2$ as shown in Scheme 5. Likewise, photo-induced generation of $\text{Co}^{IV}=\cdot\cdot\cdot\text{O}$ as active oxidant for the water oxidation was reported \cite{93} based on a cobalt(II) complex of salophen ligand. Moreover, a high-valent $\text{Co}^{IV}\cdot\cdot\cdot\text{O}$ complex isoelectronic to $\text{Co}^{IV}\cdot\cdot\cdot\text{O}$ was reported \cite{29} to act as active species to generate $\text{O}_2$ based on a cobalt(III) complex of N-based ligand bTAML (bTAML = biuret-modified tetraamidomacrocyclic) ligands. The complex $[\text{Co}(\text{O})(\text{bTAML})]^1$ cannot be characterized by spectroscopic techniques due to the non-innocent nature of the ligand except UV-vis spectra. The same specie was generated by the one electron oxidation using cerium ammonium nitrate in the presence of $\text{ZnCl}_2$. The HR-MS analysis revealed the $m/z = 497.026$ corresponding to $[\text{Co}^{IV}(\text{O})\cdot\cdot\cdot\text{Zn})(\text{bTAML})\cdot\cdot\cdot\text{(H}^+)\cdot\cdot\cdot\text{]}$. Further, Nocera et al. reported \cite{94} the dicobalt oxidized site $\text{Co}^{(III)}_2\text{Co}^{(IV)}_2$ during water oxidation using cobalt cubane modified by pyridine ligands that can stabilize tetracobalt core. This pyridine-modified cobalt cubane has molecular nature and termed as molecular cubane. Electrochemical investigation demonstrated two reversible oxidation events at $E_0(1) = 0.3$ V and $E_0(2) = 1.25$ V corresponding to $\text{Co}^{(III)}_2(\text{IV})/\text{Co}^{(III)}_4$ and $\text{Co}^{(III)}_2(\text{IV})_2/\text{Co}^{(III)}_3(\text{IV})$. X-ray absorption spectroscopy also confirms the presence of $\text{Co}^{(III)}_2(\text{IV})_2$ specie. The adjacent terminal $\text{Co}^{IV}\cdot\cdot\cdot\text{O}$ species in cubane provide a site for direct O-O bond formation by radical coupling to evolve $\text{O}_2$. Likewise, the proton-coupled electron transfer generation of $\text{Co}^{IV}\cdot\cdot\cdot\text{O}$ was also reported \cite{95} using molecular model cubane, $[\text{Co}_2\text{O}_4(\text{CO}_2\text{Me})_2_2(\text{bpy}_4)]$. Furthermore, molecular cobalt cubane $\text{Co}_2\text{O}_4(\text{OAc})_4\cdot\cdot\cdot\text{bpy}_4$ \cite{26} and a series of modified molecular cobalt cubane with electron rich and electron poor groups \cite{97} were reported to understand the nature of high-valent cobalt-oxy species involved in the water oxidation reaction. The electrochemical studies of \cite{26} revealed the presence of only one fully redox couple from pH 4 to pH 10 at $E_{1/2} = 1.25$ V corresponding to $\text{Co}^{(III)}_4/\text{Co}^{(III)}_3(\text{IV})$ redox. The increase of pH to 12 produced a significant anode wave current and bubble formation, consistent with the oxidation of hydroxide to oxygen. No change in the current intensity was observed in the presence of EDTA, ruling out the possibility of heterogeneous water oxidation due to the presence of $\text{Co}^{III}$ oxide. The ESI-MS analysis by incorporating 97% enriched $\text{Na}^{18}\text{OH}$ observed the presence of 90% $^{36}\text{O}_2$. No evidence for the exchange of $^{18}\text{O}$-oxygen between \cite{26} revealing that only terminal oxo/hydroxide specie was involved in O-O bond formation. The reaction of protonated $\text{26}^+$ and hydroxide showed the importance of the cobalt(IV) oxidation state in $\text{O}_2$ formation. The generation of $\text{cobalt}^{(V)}(\text{V})=\cdot\cdot\cdot\text{O}$ \cite{26} was proposed by PCET before the evaluation of $\text{O}_2$ as shown in Scheme 6 \cite{96}. The protonated $\text{26}^+$ reacts with hydroxide ion to produce $\text{26(O)O}^-$ which further oxidized to $\text{cobalt}^{(V)}(\text{V})=\cdot\cdot\cdot\text{O}$ \cite{26}. The specie $\text{26(O)O}$ had acted as reactive intermediate to evolve $\text{O}_2$. Involvement of high-valent $\text{cobalt}^{(V)}(\text{V})=\cdot\cdot\cdot\text{O}$ complex during water oxidation was also theoretically proposed \cite{98–101}. Corroles are analogous of porphyrin which have one carbon less than porphyrin and can stabilize metals in a higher oxidation state. A high-valent $\text{Co}^{V}=\cdot\cdot\cdot\text{O}$ specie suggested \cite{102} to act as reactive specie during water oxidation by using series of cobalt.
corroles with different axial ligand. Electrochemical study of cobalt corroles represent two reversible oxidation events at $E_{1/2} = 0.75$ and $E_{1/2} = 1.32$ V vs. NHE corresponding to Co$^{IV}$/Co$^{III}$ and Co$^{V}$/Co$^{IV}$ redox couples, respectively. Nucleophilic attack of the water at Co-O bond to generate Co-hydroperoxo specie is the key step to evolve O$_2$. Cobalt corroles with electron-donating ligands are more reactive because it causes the Co-O bond to be weaker and nucleophilic attack become easier.

Scheme 4. Molecular structures of cobalt porphyrins employed as water oxidation catalyst.

Scheme 5. Proposed mechanism of water oxidation catalyzed by cobalt porphyrins [92].

Scheme 6. Generation of high-valent Co$^V$=O 26(O) during water oxidation by molecular cobalt cubane Co$_4$O$_4$(OAc)$_4$py$_4$ [96].
4. Preparation of Cobalt–Oxo Complexes

The isolated preparation of cobalt–oxo complexes have two major problems (1). Ligands used to stabilize cobalt–oxo complexes are prone to oxidation (2). Electronic repulsion forces between the d-electron of cobalt and electron of the oxygen. Chemists are focusing on how to overcome these problems to prepare cobalt–oxo complexes.

Ray et al. reported [103] the first preparation and isolation of terminal cobalt(IV)–oxo complex using the N-based tetradentate tripodal ligand TMG$_3$tren (tris[2-([N-tetramethylguanidyl]ethyl]amine). The [Co–O] unit was stabilized by the Lewis acid interaction with Sc$^{3+}$ ion, generating [Co-O-Sc]$^{3+}$ unit. The complex 29 was obtained by two electron oxidation of 27-OTf in the presence of Sc(OTf)$_3$ (Scheme 7). The complex 29 was characterized by electrospray mass spectrum, EPR and X-ray absorption spectroscopy, and was reactive towards oxidation of triphenylphosphine and dihydroanthracene. The same group two years later reported [104] the square pyramidal cobalt(IV)–oxo with enhanced stability based on the tetraamido macrocyclic ligand (TMAL). The electrochemical study of 30 gave a reversible oxidation peak at 1.00 V vs. a saturated colomel electrode. This reversible oxidation peak suggests that Co$^{IV}$ state is thermally and kinetically accessible. The one electron oxidation of 30 in the presence of cerium ammonium nitrate (CAN) afforded a blue-colored complex 31-Sc with a half-life of 20 min. This blue complex can also be obtained by the oxidation of 30 with PhIO in the presence other redox-inactive metals like Sc$^{3+}$, Y$^{3+}$ and Zn$^{2+}$ (Scheme 8). The complex 31-M was characterized by cold-spray ionization time-of-flight mass spectrometry (CSI-TOF MS), X-band EPR spectrum, and X-ray absorption spectroscopy. All attempts to obtain resonance Raman spectrum have failed. The 31-Sc complex demonstrated high reactivity in the hydrogen abstraction reaction and oxygen atom transfer reactions. The first fully spectroscopically characterized high-valent Cobalt(IV)–oxo complex 33 was generated [105] by the two electron oxidation of a cobalt complex of 13-TMC (2 mM) 32 by PhIO (3 equiv.) following conventional method in the presence of triflic acid (CF$_3$SO$_3$H, HOTf; 1.2 equiv.) in acetone (Scheme 9). The transient complex had a half-life of 3 h and was characterized by CSI-TOF MS, EPR and X-ray absorption spectroscopy. Resonance Raman spectroscopy considered as authentic technique to confirm the presence of metal–oxo complex [106,107]. The resonance Raman spectrum of 33 showed a band at 770 cm$^{-1}$ which shifts to 736 cm$^{-1}$ upon $^{18}$O-labelling of 33. Recently, preparation of Co$^{III}$≡O complex was reported [108] by using tris-(imidazol-2-ylidene)borate ligand PhB(tBuIm)$_3$$. This complex was characterized by infrared (IR) and X-ray diffraction (XRD) spectroscopy. The length of Co-O bond determined by XRD was 1.68 Å. DFT calculations revealed two Co-O $\pi^*$ interactions with highest lying d$_{xy}$ and d$_{yz}$ orbitals. These orbitals support the presence of two $\pi$-bonds. This complex was thermodynamically unstable with half-life of 8 h.

![Scheme 7. Preparation of high-valent [Co$^{IV}$-O-Sc$^{3+}$] complex 29.](image)

![Scheme 8. Oxidation of complex 30 to 31-M by CAN/H$_2$O or PhIO/M$^{n+}$.](image)
5. Summary and Outlook

High-valent cobalt–oxo complexes are implicated as key intermediates in many of the oxidative transformation reactions and the water oxidation process. Identification of cobalt–oxo species in water-splitting reactions have been extensively studied. However, the transient nature of cobalt–oxo complexes limits their characterizations to in situ EPR, XAS and mass spectroscopy. Although different strategies, such as using ligands with different electronic environments or MWCNT supports, have been adopted to stabilize cobalt–oxo complexes, until now only one example of Raman characterization for cobalt (IV)=O complex using 1,4,7,10-tetramethyl-1,4,7,10-tetraazacyclotridecane ligand has been available. The isolation and identification of high valent cobalt–oxo species remains a great challenge. The design of a suitable N-based ligand which can stabilize coordinated cobalt atom in high oxidation might be the key step for the preparation of higher valent cobalt–oxo complexes, which will allow the full characterization and “slow motion picture” study of the factors controlling its reactivity.

Author Contributions: H.-Y.L. coordinated the whole work and provided technical guidance. A.A. and W.A. collected the references and wrote the paper.

Funding: This work was funded by the National Natural Science Foundation of China (NNSFC) under Grant (21671068).

Conflicts of Interest: The authors declare no conflict of interest.

Abbreviations

| Abbreviation | Full Form |
|--------------|-----------|
| TPFP | Meso-tetrakis(pentafluorophenyl)porphinato dianion |
| CoTAPc | cobalt tetraaminophthalocyanine |
| EPTAC | 2,3-epoxypropyl triethylammonium chloride |
| Pc | Phthalocyanine |
| TAPc | Tetraaminophthalocyanine |
| MWNTs | Multiwall carbon nanotubes |
| DMPO | 5,5-Dimethyl-1-pyrroline N-oxide |
| F9C-Co | Co(III) complex of 5,10,15-triphenylcorrole |
| F9C-Co | Co(III) complex of 5,15-bis(phenyl)-10-(pentfluorophenyl)corrole |
| F10C-Co | Co(III) complex of 5,15-bis(pentafluorophenyl)-10-phenylcorrole |
| F15C-Co | Co(III) complex of 5,10,15-tris(pentafluorophenyl)corrole |
| TMGtren | (tris[2-(N-tetramethylguanidyl)]ethyl)amine |
| *PhIO | 2-((tert-butylsulfonyl)iodosylbenzene |
| TMAL | Tetraamido macrocyclic ligand |
| 13-TMC | 1,4,7,10-tetramethyl-1,4,7,10-tetraazacyclotridecane |
| Co-TDMImP | Co(III) complex of 5,10,15,20-tetraakis(1,3-dimethylimidazolium-2-yl)porphyrin |
| Co-TM4PyP | Co(III) complex of 5,10,15,20-tetraakis(N-methylpyridinium-4-yl)porphyrin |
| Co-TTMAP | Co(III) complex of 5,10,15,20-tetraakis(N,N,N-trimethylanilinium-4-yl)porphyrin |
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