

Redox Processes Involving Oxygen: The Surprising Influence of Redox-Inactive Lewis Acids

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ABSTRACT: Metalloenzymes with heteromultimetallic active sites perform chemical reactions that control several biogeochemical cycles. Transformations catalyzed by such enzymes include dioxygen generation and reduction, dinitrogen reduction, and carbon dioxide reduction—instrumental transformations for progress in the context of artificial photosynthesis and sustainable fertilizer production. While the roles of the respective metals are of interest in all these enzymatic transformations, they share a common factor in the transfer of one or multiple redox equivalents. In light of this feature, it is surprising to find that incorporation of redox-*inactive* metals into the active site of such an enzyme is critical to its function. To illustrate, the presence of a redox-inactive Ca²⁺ center is crucial in the Oxygen Evolving Complex, and yet particularly intriguing given



that the transformation catalyzed by this cluster is a redox process involving four electrons. Therefore, the effects of redox inactive metals on redox processes—electron transfer, oxygen- and hydrogen-atom transfer, and O–O bond cleavage and formation reactions—mediated by transition metals have been studied extensively. Significant effects of redox inactive metals have been observed on these redox transformations; linear free energy correlations between Lewis acidity and the redox properties of synthetic model complexes are observed for several reactions. In this Perspective, these effects and their relevance to multielectron processes will be discussed.

KEYWORDS: multimetallic effects, redox reactions, metal ion-coupled electron transfer, oxygen transformations, Lewis acids, bioinorganic chemistry

1. INTRODUCTION

The conversion of small molecules such as O2, N2, CO2, H2O, and H₂ into value-added chemicals or liquid energy carriers is important in (i) the context of solar energy conversion and storage, (ii) the efficient use of inexpensive feedstocks, and (iii) the sustainable generation of bulk chemicals. However, transforming these small molecules into useful chemicals is challenging due to the number of e^{-}/H^{+} equivalents transferred and in terms of the necessity for controlling selectivity and the performance of the chemistry at low overpotentials.¹ The biological catalysts for small molecule conversions are proteins that often involve complex inorganic active sites: Mn₄Ca in Photosystem II (PSII) for O₂ evolution;² Cu₃ in laccase and CuFe in cytochrome c oxidase for O₂ reduction;³ Fe₄Ni or CuMo in CO dehydrogenase (CODH) for CO_2 reduction;⁴ Fe₈, Fe₇Mo, or Fe₇V in nitrogenase for N₂ reduction;⁵ Fe₆ or FeNi in hydrogenase, for proton reduction and hydrogen oxidation.⁶ In many of these gas-processing enzymes, two or more types of metals are found in their active sites. Accordingly, the interactions between different metal centers and their influence on

structure and reactivity have been investigated in numerous synthetic systems and have also been the subject of several reviews. $^{7-28}$

Among the enzymes that depend on heteromultimetallic active sites, two systems are unusual for the involvement of metal ions that are redox-inactive, namely (i) the oxygen evolving complex (OEC) of Photosystem II, which contains a Ca^{2+} ion and catalyzes the oxidation of H₂O to O₂, and (ii) Cu/Zn-dependent superoxide dismutase (Cu/Zn-SOD), which catalyzes the disproportionation of superoxide (O₂^{•-}) to H₂O₂ and O₂ at an active site containing a redox-inactive Zn²⁺ ion.^{2,29} Because both enzymes are involved in oxygen chemistry, their structural/activity relationships bear great relevance for energy science and industrial processes. Not

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Figure 1. Lewis acid effects on redox processes involving oxygen. The linear correlation between the pK_a of the metal aqua ion and the "properties" of corresponding metal complexes (center) are universally found among many transformations that involve oxygen including (i) electron transfer (top left), (ii) oxygen-atom transfer (bottom left), (iii) O₂ activation and cleavage (top right), and (iv) O–O bond formation (bottom right).

surprisingly, many investigations have probed the role of these redox-inactive metal ions in catalysis. While a primarily structural role has been ascribed to Zn^{2+} in Cu/Zn-SOD,^{3,30-32} with SOD activity being largely retained upon Zn^{2+} loss,³⁰ the role of Ca²⁺ in the OEC has not been conclusively elucidated. Consequently, synthetic models of the OEC have been targeted with the aim of unraveling fundamental reactivity relevant to biological water oxidation.³³⁻³⁵ Interest in the effects of redox-inactive metals on redox processes involving oxygen has been further piqued by discoveries in the field of heterogeneous catalysis, where mixed-metal oxides containing both redox-active and -inactive components have gained significant attention as electrocatalysts in the oxygen evolution reaction (OER).³⁶⁻⁴⁰ Moreover, employing multiple different metal centers in discrete complexes can likewise result in new physical properties and reactivity pathways beyond oxygen chemistry.⁴¹⁻⁴⁶ As a result, aspects of the effects of redox inactive metals on the redox reactivity of transition metals with oxygen have been addressed in several types of reactions, and this field has seen significant increase in research activity.^{47–53}

Herein, we provide an overview of the literature concerning the behavior of well-defined molecular systems displaying a combination of redox-active and redox-inactive metals in the context of oxygen-related chemistry. Effects of redox-inactive metals on electron transfer (ET), oxygen-atom transfer (OAT), hydrogen-atom transfer (HAT), as well as O-O bond cleavage and formation, will be discussed (Figure 1). Because of their inability to directly participate in redox processes, the basis for controlling reactivity with redoxinactive metals must leverage properties such as their charge, size, polarizability, and/or coordination number. Given the complex interplay between these features, many (though not all) studies in this area have shown that the observed effects on redox properties cannot be ascribed to a single factor, but rather to the broader concept of Lewis acidity invoked by constructing linear free-energy relationships based on experimental properties (e.g., reduction potentials, reaction rates). Because Lewis acidity is not a well-defined quantitative parameter, several alternative scales have been developed in an attempt to provide a semiquantitative measure of Lewis acidity.^{56,37} Methods based on nuclear magnetic resonance $(NMR)^{58,59}$ or fluorescence spectroscopy,^{60,61} for instance, have been successfully implemented in various contexts. A scale of Lewis acidity based on electron paramagnetic resonance (EPR) measurements has also been developed by Fukuzumi and co-workers.⁶² Our work has relied on the pK_a value of metal–aqua ions as a measure of Lewis acidity in part due to the availability of values for mono-, di-, and trivalent ions.^{56,63–65} These values also emerge from the same interplay between distinct factors that makes use of individual parameters often inadequate to explain experimental observations. Overall, useful correlations between Lewis acidity and various types of electron- and group-transfer reactivity exist across many transition metal compounds of varying composition, nuclearity, and complexity.

In this Perspective, correlations between observed properties and Lewis acidity will be presented using the methods for quantification of Lewis acidity described in the original publications, including in cases where other parameters (e.g., ionic radius or charge) were found to provide a more satisfactory correlation. In cases where such correlations were not originally provided, or in cases where a comparison across systems originally relying on different Lewis acidity scales would be particularly insightful, the pK_a of metal-aqua ions will be used as the measure of Lewis acidity. Values for the pK_{a} of a broad series of metal-aqua ions are shown in Table 1. The majority of these values were obtained from the extensive lists reported by Perrin;⁶³ wherever possible, values extrapolated to I = 0 were chosen to best represent thermodynamic values. It should be noted, however, that significant variance exists for the pK_a of metal-aqua ions in different sources, especially for transition metal ions.^{64,66,67} Therefore, where significantly different values ($\Delta p K_a \ge 0.5$) have been cited in other literature sources, these values were also included in the table. For the purposes of identifying correlations between observable properties (reduction potentials, reaction rates, etc.) and Lewis acidity in this Perspective, the Perrin values⁶³ were used (with the exception of the values for K⁺, Cs⁺, and Rb⁺, which were recently extrapolated on the basis of ³¹P NMR experiments).⁵⁶ Nonetheless, the uncertainties inherent in these values make quantitative interpretations of such correlations challenging, and these relationships will therefore be discussed here primarily in qualitative terms.

Table 1. pK_a of $[M(H_2O)_m]^{n+}$ Species

M^{n+}	pK_a^a	M^{n+}	pK_a^a	M^{n+}	pK _a ^a
Mn ³⁺	$0.20 (-0.6)^{b}$	Eu ³⁺	8.03 (8.6) ^b	Ni ²⁺	9.86
Co ³⁺	$0.66 (0.5)^{b} (1.75)^{c}$	Y ³⁺	8.04	Mn ²⁺	10.59
Fe ³⁺	2.19	Dy ³⁺	8.10	Mg ²⁺	11.40
Ga ³⁺	2.92	Tb ³⁺	8.16	Ca^{2+}	12.60
In ³⁺	3.54	Gd^{3+}	8.40 (9.78) ^c	Sr ²⁺	13.18
Al ³⁺	4.90	Nd ³⁺	8.43 (9.0) ^c	Li ⁺	$\begin{array}{c} 13.11 \ (13.6)^{b} \\ (13.8)^{c} \end{array}$
Sc ³⁺	4.93 (4.3) ^b	Co ²⁺	$8.90 (9.7)^{b} (9.85)^{c}$	Ba ²⁺	13.40
Cu^{2+}	7.34	Zn^{2+}	8.96	Na^+	14.80 (13.9) ^b
Lu ³⁺	7.66 (8.2) ^b	La ³⁺	9.06 (8.5) ^b	K^+	16.06^{d} (14)
Pb^{2+}	7.78	Cd^{2+}	9.30 (10.1) ^b	Cs^+	16.29 ^d
Yb ³⁺	8.01	Fe ²⁺	9.30 (6.8) ^c	Rb ⁺	16.34 ^d
-					1.

 ${}^{a}pK_{a}$ values from ref 63 except where otherwise noted. ${}^{b}pK_{a}$ values from ref 67. ${}^{c}pK_{a}$ values from ref 64. ${}^{d}pK_{a}$ values from ref 56.

2. ELECTRON TRANSFER AND LEWIS ACIDS

Lewis acids have been shown to affect reduction potentials and rates of electron transfer in transition metal compounds (Figures 2 and 3). These effects were observed either upon addition of excess redox inactive metal ion to solutions of transition metal complexes displaying a basic site for binding of additional metal ions or by utilizing well-defined heterometallic complexes.

One of the more broadly studied effects of Lewis acids on electron transfer involves the conjugation of redox-active metal complexes to crown-ethers (Figure 2).⁶⁸⁻⁸⁸ Extensive studies on mono- (1) and bimetallic (2) manganese Schiff-base complexes first appeared in the 1990s that aimed to elucidate the properties of manganese-dependent enzymes (including the OEC).⁷⁰⁻⁷² While salen-type related ligands have been shown to be useful for binding or sensing of a wide variety of redox-inactive metals via coordination to O atom donors (e.g., phenoxide),⁸⁹ inclusion of crown-ether motifs enables stronger binding of Lewis acidic ions with stoichiometric control. Exposing complex 1 to Li⁺, K⁺, Ca²⁺, or Ba²⁺ leads to incorporation of a single redox-inactive metal into the crownether moiety, resulting in a positive shift in reduction potential for the Mn^{III}/Mn^{II} redox couple.⁷² Nearly identical behavior was displayed by the μ -oxo dimer of 1, although the observed redox couples corresponded to the two-electron reduction of the [Mn^{III}₂O] core in this instance. A similar effect was observed in a related system (7), where the less rigid aminopropane backbone enables isolation of a bridged (μ -O)₂ complex. Incorporation of redox-inactive metal ions in this $Mn^{IV}-(O)_2-Mn^{IV}$ species leads to positive shifts in potential for the $Mn^{IV}_2/Mn^{III}Mn^{IV}$ and $Mn^{III}Mn^{IV}/Mn^{III}_2$ redox couples, though only redox-inactive metals with a narrow range of Lewis acidities ($\Delta p K_{a}$. = ~2.0) were investigated.⁷¹ A similar



Figure 2. Linear correlation between the reduction potential of crown-ether supported (multinuclear) metal complexes and the Lewis acidity of redox-inactive metal ions. (A) Selection of crown-ether supported metal complexes. (B) Observed linear correlation is shown for complexes 1-M (μ -oxo dimer, pink diamonds),⁷² 2-M (blue dots),⁶⁸ 3-M (dark green dots),⁷⁸ 4-M (light green dots),⁸⁵ 5-M (purple dots),⁸⁴ 6-M (salen-type backbone, teal triangles),⁸⁶ 6-M (modified backbone, orange triangles),⁸⁷ 7-M (two redox events, maroon and gray diamonds, respectively),⁷¹ and 8-M (red dots).⁷⁷



Figure 3. Linear correlation between the reduction potential of (multinuclear) metal complexes and the Lewis acidity of redox-inactive metal ions. (A) Observed linear correlation is shown for Mn_3MO_4 (**10-M**; blue), $Mn_3MO(OH)$ (**11-M**; green), and Fe₃MO(OH) (**12-M**; red) type clusters; Ln = La, Nd, Eu. Gd, Tb, Dy, Yb, Y. (B) Linear correlation observed for the rate of electron transfer between **17-M** and ferrocene. (C) Selection of metal complexes that demonstrate modulation of the reduction potential upon interaction with redox-inactive metal ions. Figure 3B is reproduced from ref **149**. Copyright 2011 American Chemical Society.

set of redox-inactive metals (Na⁺, K⁺, Ca²⁺, Sr²⁺, and Ba²⁺) was screened in structurally related cobalt complexes (**2-M**). In these complexes, a linear correlation between the pK_a of the redox-inactive metal and the reduction potential of the Co^{II/I} redox-couple was observed, though the narrow range of Lewis acidities once again limits generalization of this relationship.⁶⁸ Related nickel,⁷⁸ iron,⁸³ zinc,⁸⁵ palladium,⁸⁴ and vanadyl^{86,87} complexes (**3-M** to **6-M**) have been shown to display similar correlations between the reduction potential of the complex (typically corresponding to a metal-centered ET process, though a ligand-based one for Pd and Zn) and the Lewis acidity of a wider range of cations (e.g., Na⁺, Ca²⁺, Nd³⁺, Y³⁺). Incorporating a more flexible triamine-based backbone enables coordination of these Schiff-base ligands to the uranyl cation $([UO_2]^{2+}; 8)$ and thereby enables the extension of these relationships to actinide complexes. The Lewis acidity of the incorporated redox-inactive metals (K⁺, Na⁺, Ca²⁺, and Nd³⁺) linearly modulates the reduction potential of the U^{VI}/U^V redox couple. Notably, the reversibility of the observed electrochemical processes is also affected by the Lewis acidity of the



Figure 4. Plot of reduction potential vs effective basicity for a series of cubane complexes. (A) Selection of metal-oxo clusters. (B) Linear correlation between reduction potential and effective ligand basicity in $[Mn_4O_4]$ complexes, including 10-Mn (blue circles). Similar trend expected in $[YMn_3O_4]$ complexes, including 10-Y (red circles), with the exception of trisamidate complex 10-Y(triam) due to geometric effects. Graph adapted from ref 119. Copyright 2019 American Chemical Society.

redox-inactive metal ions, with evidence of downstream chemical reactivity for more Lewis acidic divalent and trivalent redox-inactive cations. 77

Heterobimetallic complexes with other ligands containing phenolate moieties have also been reported, although they frequently involve a combination of two redox-active metals.^{90–103} In these complexes, some effects can be attributed to the Lewis acidity of a metal ion, but their properties are mainly dominated by electrostatic effects. Notable exceptions are the rare-earth/alkali-metal heterobimetallic complexes (9) reported by Walsh and Schelter.^{104–106} Binding of redox-inactive metals to cerium complexes supported by binolate ligands modulates the reduction potential for the Ce^{IV/III} couple. Different redox-inactive metals were also found to promote formation of chemical oxidation products featuring diverse binding modes of the Lewis acidic ions as a function of their identity.

Our group has extensively studied the effects of Lewis acidity on reduction potentials in multimetallic cluster complexes (10-M to 12-M). $^{67,107-111}$ Some of these clusters (10-M) closely match the structural aspects of the OEC, for which only a few well-defined structural examples have been reported.¹¹²⁻¹¹⁶ These tetraoxido clusters display a [Mn₃CaO₄] "cubane" structural motif, with the redox-inactive metal situated at the apical position (10-M; Figure 3).^{108,109} In these compounds, the potential for the Mn^{IV}₃/Mn^{III}Mn^{IV}₂ redox couple shifts from -0.94 V (vs the ferrocenium/ferrocene couple, hereafter denoted $Fc^{+/0}$) for Sr^{2+} to +0.29 V for Mn^{3+} , and correlates linearly with the Lewis acidity of the corresponding metal ion. The linear fit is observed over a wide pK_a range ($\Delta pK_a \sim 15$), which allows a broader demonstration of the trend, and displays a potential shift of *ca*. 0.1 V for every pK_a unit (Figure 3A; blue dots). Rather than Lewis acidity, computational studies of the OEC incorporating different redox-inactive cations suggested a correlation between the reduction potentials of the clusters and the charge of these redoxinactive cations.⁶⁷ Notably, the protein environment that encages the OEC cluster in Photosystem II imposes bonding pressures from all directions.^{117,118} This is in contrast with

synthetic cubane compounds, where the carboxylate bridges do not place a similar constraint on the apical, redox-inactive metal center. To address the potential effects of geometric constraints of the protein environment on the OEC cluster, synthetic cubane variants with chelating amidate ligands were prepared (Figure 4). Interestingly, a caveat to the correlation between the reduction potential and the Lewis acidity trend was observed in these systems. In a series of $[Mn_4O_4]$ (10-Mn) and $[YMn_3O_4]$ (10-Y) cubane clusters,¹¹⁹ our group showed that ligand basicity modulates the reduction potentials of the clusters in a similar fashion to reported $[Co_4O_4]$ and $[RuCo_3O_5]$ complexes.^{120,121} However, the tris-amidate substituted YMn_3O_4 (10-Y(triam)) cluster displays a 140 mV positive shift in the potential for the [YMn^{IV}₃]/ [YMn^{IV}₂Mn^{III}] couple relative to the tris-acetate cluster (10-Y), inconsistent with the increased basicity of amidates versus acetates. This exception can be explained by geometric changes in the cluster core due to ligand steric constraints-chelating tris-amidate ligands contract the Y-oxo distances by roughly 0.1 Å compared to the those in the tris-acetate cluster, decreasing the electron density available at the Mn centers. Relevant to the cation size dependence on the activity of the OEC, these results suggest geometric constraints may cause nonlinear changes in reduction potentials and reactivity, engendering more pronounced effects than those predicted on the basis of its lower Lewis acidity. In other words, a larger metal may behave like a smaller, more Lewis acidic one of the same charge, resulting in a correlation of properties with charge, not Lewis acidity.

The role of the number of μ -oxo ligands was also investigated with similar tetranuclear complexes **11-M** and **12-M**,^{107,110} which feature a $[M_3(\mu_4-O)(\mu_2-OH)M']$ core (M = Mn, Fe; M' = redox-inactive metal; Figure 3). The Lewis acidity of the redox-inactive metal influences the potential for the $M^{III}_3/M^{III}_2M^{II}$ couples, which shift from $E_{1/2} = -0.30$ to +0.42 V (vs Fc^{+/0}) for M = Mn (Figure 3A, green dots) and -0.49 to +0.07 V for M = Fe (Figure 3A, red dots) as the Lewis acidity of the incorporated redox-inactive metal increases.

Based on the observations from the two series of $[Mn_3]$ complexes, it appears that reducing the oxo content results in a positive shift (~ 0.5 V) in the reduction potential of the corresponding cluster. Despite the higher Mn oxidation states in cubane-type vs oxo-hydroxo complexes ([Mn^{IV}₃] vs [Mn^{III}₃] in the oxidized forms, respectively), the presence of additional electron-rich O²⁻ moieties increases the electron density around the metal centers, shifting the cluster's reduction potential to more negative values compared to complexes 11-M and 12-M. Intriguingly, the nearly identical reduction potentials engendered by calcium and strontium in each series support the hypothesis that redox-inactive metals might be involved in the tuning of reduction potentials in PSII, as substitution of Ca²⁺ with Sr²⁺ uniquely enables retention of catalytic competence in this system. It must be noted, however, that in computational studies of the OEC, substituting Ca²⁺ for other redox-inactive metals did not produce a linear correlation between the redox properties of the OEC and the Lewis acidity of the redox-inactive metals.⁶⁷

The effects of calcium and strontium on electrochemical properties have also been studied in a series of Fe(14) and Mn(15) complexes supported by tripodal tris(sulfonamido) (MST) ligands.^{122,123} Remarkable rate enhancements in O_2 activation by Fe^{II}/Mn^{II} precursors were observed upon addition of Ca²⁺ and Sr²⁺ to generate Fe^{III}/Mn^{III}-(OH)-M complexes 14-M and 15-M, whereas only limited effects were observed upon the addition of less Lewis acidic Ba²⁺. The reduction potentials of complexes 14-Ba and 15-Ba were also found to be significantly more negative (by $\sim 0.06 - 0.10$ V) than those of the isostructural Ca^{2+} and Sr^{2+} analogues. The observation of only minor differences in reduction potential between Ca^{2+} and Sr^{2+} complexes (e.g., ~0.02 V for 15-Ca vs 15-Sr) led the authors to assert that Lewis acidity alone is not sufficient to explain the observed trends, since a larger difference might be expected for Ca²⁺ vs Sr²⁺. Nonetheless, the difference in Lewis acidity between these congeners is indeed small $(\Delta p K_a < 1)$; accordingly, very similar reduction potentials may be expected for Ca and Sr complexes (vide supra).^{107,108,110} A related trisphosphinamide ligand framework, also developed by the Borovik group, was shown to enable anchoring of a variety of 18-crown-6 encapsulated Lewis acids in the proximity of a terminal iron-oxo motif (16).¹²⁴ Although no electrochemical data were reported for this series, the linear trend between pK_a and the energy of the first electronic absorption band was described, indicating that binding of the Lewis acid modulates the molecular orbital energies of the Fe complexes.

Another important class of compounds that display similar effects on ET are complexes that bear high-valent terminal metal—oxo moieties. In nature, these metal—oxo motifs are ubiquitous as intermediates in many oxidative transformations,^{16,125–131} and have even been suggested in biological water oxidation.^{2,132} Consequently, intensive efforts have been undertaken to study these motifs in synthetic model complexes.^{127,133–145} The well-defined nature of many of these systems has also enabled systematic investigations into the effects of Lewis acids on their properties and reactivity. Although many of these complexes are also competent for OAT, HAT, O₂ activation, and/or O–O bond cleavage, their reactivity in these types of transformations will be discussed in later sections (*vide infra*), and the current discussion will remain focused on electron-transfer reactions.^{146–154}

One of the first studies of the effect of redox-inactive metals on high-valent terminal Fe^{IV}-O complexes was reported with a cyclam macrocycle.^{154,155} (TMC)Fe^{IV}(O)²⁺ (**27**; TMC = N, N', N'', N'''-tetramethylcyclam) behaves as a one electron oxidant toward excess ferrocene;¹⁵⁶ however, addition of Sc³⁺ induces a change in the redox chemistry of $(TMC)Fe^{IV}(O)^{2+}$ and a second electron transfer to complex 27 from Fc is observed.¹⁵⁴ Kinetic studies confirmed that the reaction proceeds via a rate-determining 1e⁻ reduction, followed by a fast second ET step. Interestingly, Ca²⁺ induces a similar shift in the reactivity of complex 27, albeit only when present in large excess (40 equiv vs 2 equiv for Sc^{3+}).¹⁵⁴ These observations are consistent with a more positive reduction potential of the Fe^{IV}=O complex upon binding of Lewis acids. Direct evidence for binding of Sc³⁺ to terminal iron-oxo moiety was provided by X-ray diffraction studies on an isolated complex assigned as [(TMC)Fe^{IV}(O)-Sc(OTf)₄(OH)]. Recent reports based on computational modeling and Mössbauer and EPR data, however, have cast doubt on this assignment, suggesting the complex $[(TMC)Fe^{III}-(OH)-Sc^{-1}]$ $(OTf)_4(H_2O)$] as the most likely identity of the species characterized by crystallography.^{157,158}

Peroxide complexes of the form $[(TMC)Fe^{III}(O_2^{2-})M]^{n+}$ $(26-M, M = Ca^{2+}, Sr^{2+}, Zn^{2+}, Lu^{3+}, Y^{3+}, Sc^{3+})$ can be generated from the same Fe^{II} precursor used in preparation of 27 (a process which is itself dependent on the identity of the redoxinactive metal, see Section 4). Electrochemical investigations indicated that the irreversible 1e⁻ reduction potentials correlate linearly with the Lewis acidity of the redox-inactive metal ions. Additionally, the rate for ET between complexes 26-M and 1e⁻ reductants (e.g., bromoferrocene) is also dependent on Lewis acidity, with a plot of log(k) vs Lewis acidity displaying a linear correlation (Figure 6B, blue dots). In the same series, the lower Lewis acidity of Ca²⁺ and Sr²⁺ additionally engendered a shift in the potential required for the oxidation of the complex to values sufficiently negative to enable chemical oxidation resulting in O2 evolution (see Section 5 for details on this reactivity), while complexes containing stronger Lewis acids did not show evidence for electrochemical oxidation. Notably, peroxo complex 26 can itself be oxidized (both electrochemically and with chemical oxidants) to generate 25 and O_{2} ; therefore, the unique behavior of the Ca²⁺ and Sr²⁺ complexes is most accurately ascribed to these ions being the only ones in this series whose Lewis acidity is sufficiently low for the relevant oxidation to occur at moderate potential, while stronger Lewis acids shift the potential for this oxidation too far positively for analogous behavior to be observed.

Complex $[(N4Py)Fe^{IV}(O)]^{2+}$ (17; Figure 3C)), supported by a tetrapyridine ligand, also exhibited intriguing behavior in the presence of Lewis acidic metal ions.¹⁴⁹ The redox properties of this complex were shown to be affected by Sc³⁺ ions, enabling the oxidation of $[Fe^{II}(bpy)_3]^{2+}$, whose potential $(E_{1/2} = +1.06 \text{ vs SCE})$ is normally too positive to allow oxidation by 17 ($E_{red} = +0.51$).¹⁵⁶ Additional experiments suggested that the reduction potential for 17-Sc ranges between +1.26 and +1.35 V (vs SCE), an increase of ~0.8 V compared to $[(N4Py)Fe^{IV}(O)]^{2+}$. Investigation of the effects of Sc³⁺ concentration on ET suggested that two Sc³⁺ ions bind to the Fe^{IV}—oxo moiety at high Sc³⁺ concentrations. Rate acceleration for ET is also observed in the presence of Sc³⁺ and other redox-inactive metal ions (Y³⁺, Lu³⁺, Zn²⁺, Mg²⁺, and Ca²⁺), and ET rates correlate linearly with Lewis acidity



Figure 5. Linear correlation between Lewis acidity of redox-inactive metal ions and oxygen atom transfer (OAT) reactions. (A) Selection of metal complexes where the rate of oxygen atom transfer (OAT) is regulated by the Lewis acidity of bound redox-inactive metals. (B-D) Observed linear correlation between the rate of OAT and Lewis acidity of the redox-inactive metal for oxidation of methanol by permanganate (B), and oxidation of triphenylphosphine to triphenylphosphine oxide by **21** (C) and **22** (D). Figure 5B–D was generated by analyzing the data reported in refs 173, 180, and 194.

(Figure 3B). In the isostructural Mn complex [(N4Py)- $Mn^{IV}(O)$]²⁺ (18) the reduction potential shifts from +0.80 V, to +1.31 V, and finally to +1.42 V (vs SCE) upon binding one and two Sc³⁺ ions, respectively.^{146,152} The effect of other Lewis acidic redox-inactive metals, however, has not been reported. Similarly, [(dpaq)Mn^{IV}(O)]⁺-Mⁿ⁺ (Mⁿ⁺ = Ca²⁺, Mg²⁺, Zn²⁺, Lu³⁺, Y³⁺, Al³⁺, and Sc³⁺) complexes (dpaq = 2-[bis(pyridin-2-ylmethyl)]amino-N-quinolin-8-yl-acetamidate) (19) showed a linear dependence between ET rate constant and Lewis acidity of Mⁿ⁺, with increasing Lewis acidity enhancing reactivity.¹⁵⁹

In studies on a related Cr complex containing a superoxide fragment, $[Cl(TMC)Cr^{III}(O_2^{\bullet-})]^+$, ET from 1e⁻ reductants to Cr was enabled by the presence of redox-inactive metals (Ca²⁺, Mg²⁺, Y³⁺, Al³⁺, Sc³⁺), akin to the behavior exhibited by complex 17.¹⁶⁰ Furthermore, consistent with the results for complex 17, a linear correlation was observed between the logarithm of the rate of electron transfer and the Lewis acidity of the redox-inactive metal, highlighting the generalizability of these effects in complexes containing high-valent metal centers bound to oxygen-derived ligands.

Lewis acid binding was also found to tune the reduction potential of a corrolazine-supported Mn-oxo species studied by Goldberg and co-workers.¹⁵⁰ Using the octakis(*p-tert*-

butylphenyl)corrolazinato (TBP₈Cz)³⁻ ligand, a Mn^V-oxo complex (13; Figure 3C), a diamagnetic species that is unreactive towards reduction by ferrocene, was isolated. Addition of one equivalent of $Zn(OTf)_2$ results in conversion to a paramagnetic species whose spectroscopic features are consistent with formation of a corrolazine π -cation radical. Due to Zn^{2+} being redox-inactive, oxidation of the corrolazine ring indicates concurrent reduction of the Mn^V center via intramolecular electron transfer. The resulting species was therefore assigned as the valence tautomer complex $[(TBP_8Cz^{\bullet+})Mn^{IV}(O)-Zn^{2+}]$ (13-Zn). Treatment of 13-Zn with 1,10-phenanthroline sequesters the Zn²⁺ cation and regenerates 13, confirming the association of Zn^{2+} to the metal complex and the chemical reversibility of the intramolecular ET event. No dependence on Zn²⁺ concentration (beyond 1 equiv) was observed, suggesting that a single zinc ion is bound to the Mn complex. 13-Zn is reduced by ferrocene or 2,4-ditert-butylphenol, yielding a Mn^{IV} complex with a closed-shell corrolazine ring. This contrasts the electrochemical behavior of precursor 13 and aforementioned 27-Sc, which are 2e⁻ oxidants.¹⁵⁴ Moreover, 13-Zn is unique in its ability to form a 1:1 adduct with Zn exclusively, whereas other analogous

 Mn^{IV} –O complexes appear to be able to bind multiple redoxinactive metals.^{146,152}

Valence tautomerism was also suggested during the formation of a metastable Co^{IV}-oxo complex (24) generated by addition of PhIO to $[(TAML)Co^{III}]^{1-}$ (TAML = tetraamido macrocyclic ligand) (Figure 5).¹⁵¹ The presence of Lewis acids (e.g., Sc^{3+} , Y^{3+} , and Zn^{2+}) stabilizes complex 24, which was predicted to be otherwise highly reactive due to population of the metal-oxo π^* orbitals. The presence of a Co^{IV}-oxo was established by extended X-ray absorption spectroscopy (EXAFS), which indicated a Co-O distance of 1.67 Å – slightly shorter compared to other reported Co^{IV} -oxo moieties.^{148,161} DFT calculations carried out on complex 24 predict a doublet ground state with significant spin density on the oxygen atom, giving rise to a radical "oxyl"-type moiety and, consequently, a formal assignment of 24 as [(TAML)- $Co^{III}(O^{\bullet})]^{2-}$. However, in the presence of scandium ions, the unpaired-electron spin is mainly located on cobalt, suggesting transfer of one electron from Co^{III} to oxygen leading to a [(TAML)Co^{IV}(O)Sc]⁺ assignment, highlighting the role of Sc³⁺ in valence tautomerism. Earlier studies also reported on the stabilizing effect of scandium ions on high-valent Co^{IV}-oxo moieties, allowing the observation of the scandium adduct in $[(TMG_3tren)Co^{V}(O)Sc(OTf)_3]^{2+}$ $(TMG_3tren = tris[2-N-tetramethylguanidyl)ethyl]amine).^{148}$ However, subsequent investigations on this system have suggested that a $[(TMG_3tren)Co^{III}(OH)Sc(OTf)_3]^{2+}$ assignment might be more accurate, ^{160,162} highlighting the challenges in accurately determining the oxidation state of transition metals and protonation state of bridging oxo ligands.

Taken together, these reports provide a clear insight into how redox-inactive metal ions influence the reduction potential and the rate of ET of a broad spectrum of mono- and multimetallic transition metal complexes. In most cases, experimentally determined values (e.g., reduction potential) display linear correlations with Lewis acidity, enabling extrapolation of these relationships and prediction of the properties of hitherto unexplored compounds. These effects have also been successfully modeled, in cases where binding of the redox-inactive ions is an equilibrium process, on the basis of the Nernst equation, which can be used to quantify the shift in reduction potential as a function of the differential binding of these ions to the oxidized and reduced forms of the redoxactive fragment. This model, originally developed to analyze the shift in reduction potentials engendered by redox-inactive metal ions in organic fragments (e.g., flavins, quinones, and other carbonyl compounds),^{50,163,164} has been successfully applied to several of the transition metal systems described in this section, especially in the context of high-valent metal-oxo complexes 17, 18, and 19.146,149,159,165 Additionally, the Marcus equation of outer-sphere electron transfer has been applied to the analysis of reactions between high-valent metaloxo compounds and substituted ferrocenes as 1e⁻ donors in order to outline the dependence of ET rate on the driving force for electron transfer.¹⁴⁹

Alternative models have been proposed to rationalize the effects of redox-inactive metal ions on the potential of transition metal complexes to those, discussed in this section, leveraging Lewis acidity to outline correlations with experimental reduction potentials and ET rates. In particular, the effects of redox-inactive metals on the potential of Co, Ni, and Fe complexes supported by salen-type ligands with appended crown ether motifs (such as those shown in Figure 2) have

been ascribed to electrostatic effects engendered by the redoxinactive ions.^{81,83,103,166} According to this model, the electric field generated by the Lewis acidic ions, which depends on the ionic charge and on the distance between redox-active and -inactive sites, is the primary source of shifts in reduction potential in systems in which the incorporation of redoxinactive ions does not significantly alter the ligand field at the redox-active metal center; in these cases, the redox-inactive ions can be modeled as point-charges, though quantitative analyses are complicated by the necessary estimation of dielectric constants for calculations using Coulomb's law¹⁰³ or the need for significant computational work.¹⁶⁶ On the other hand, in cases where Lewis acid binding perturbs the ligand field at the transition metal site more significantly (e.g., when the geometry of bridging ligands is more substantially affected by ion incorporation), both electrostatic and inductive effects are operative and must be considered for adequate analysis. It should be noted that the electrostatic model has been applied primarily in systems where only narrow ranges of redox-inactive ions were investigated (e.g., K⁺, Na⁺, and Ba²⁺, $\Delta p K_a \sim 2.6$; therefore, broader series likely need to be determine to fully explore the true scope of these effects.

One common feature of the systems described in this section (and in this Perspective overall) is the absence of significant metal-metal interactions between redox-active and -inactive centers. In complexes where such metal-metal bonding interactions are present, deviations can occur as a result of the orbital overlap between adjacent metal centers.¹⁶⁷ In a representative example, Lu and co-workers found that the reduction potential of a series of nickel-M complexes (M = Al³⁺, Ga³⁺, In³⁺) supported by tripodal ligands featuring N and P donors varies as a function of heterometal identity. However, a stronger correlation between reduction potential and the size or charge density of the redox-inactive ions, rather than their metal-aqua pK_a or other parameters was observed.¹⁶⁸ These findings were ascribed to the larger redox-inactive metals displaying increased overlap with the Ni⁰ orbitals, resulting in a stronger effect despite the lower Lewis acidity. As metal-metal bonding involves a more direct interaction between redoxinactive and -active metals, it is perhaps unsurprising that in this and analogous cases a stronger correlation of experimental properties with parameters other than Lewis acidity (metal size, charge, or other factor) could be observed. Since metalmetal interactions have not been identified in the OEC, synthetic systems leveraging metal-metal bonds will not be discussed further in this Perspective.

3. LEWIS ACID EFFECTS ON OXYGEN ATOM TRANSFER

As evident from the previous examples, redox modulation by Lewis acids is important for various processes involving electron transfer. Lewis acid effects are also observed in the O atom transfer reactivity of high-valent metal—oxo species. In fact, some of the earliest reports of the effects of redox-inactive metals on redox processes focused on high-valent transition metal—oxo species and their reactions with various substrates in the presence of redox-inactive metals. Ruthenate $([Ru^{VO}_2(OH)_3]^{2-}),^{169}$ chromate $([Cr^{VI}_2O_7]^{2-}),^{170}$ permanganate $([Mn^{VII}O_4]^{-}),^{170-173}$ ferrate $([Fe^{VI}O_4]^{2-}),^{174}$ and nitridoosmate $([Os^{VIII}O_3(N)]^{-})^{175,176}$ were shown to be more effective oxidants in the presence of a variety of Lewis acids, including several Group 1 and 2 metals. In their seminal work, Lau and co-workers reported enhanced oxidation of the

unactivated C-H bonds in cyclohexane under mild conditions by ruthenate, perruthenate, chromate, permanganate, and ferrate. ^{169,170,174} Formation of cyclohexanone is accelerated by addition of Zn^{2+} , Al^{3+} , Fe^{3+} , Cu^{2+} , Mg^{2+} , and Li^+ salts, an effect that is also mildly anion-dependent. Even at this early stage, rate acceleration was proposed to involve a shift in the reduction potential due to binding of a Lewis acid to the M-O moieties of the oxoacid. Electrostatic effects were ruled out based on the observation that trifluoroborane-a coordinating but neutral Lewis acid-leads to rate acceleration while the positively charged, noncoordinating tetraphenylphosphonium cation has no effect on the reaction rates.¹⁷⁷ Reaction of KMnO₄ with methanol was also observed to be greatly accelerated (up to 10^7 -fold) in the presence of Lewis acids (e.g., Ba^{2+} , Ca^{2+} , Zn^{2+} , Sc^{3+} , or BF_3).¹⁷³ Our analysis of the reported kinetic data reveals a clear linear correlation between reaction rate and the Lewis acidity of the redox-inactive metal (Figure 5B). Recently, Lau and co-workers further demonstrated cooperative effects of mildly Lewis acidic Ca2+ and acetic acid, a weak Brønsted acid, in activating RuO4- and MnO₄⁻ toward alkane oxidation.^{178,179}

Cross-bridged cyclam-supported manganese complexes $(Me_2EBC)Mn^{11}Cl_2$ (20; $Me_2EBC = 1,1$ -dimethyl-1,4,8,11tetraazabicyclo[6.6.2]-hexadecane) and (Me₂EBC)- $Mn^{IV}(OH)_2(PF_6)_2$ (21) display another example of reactivity modulation by Lewis acidic ions.¹⁸⁰ Complex 20 is a sluggish catalyst for oxygen atom transfer to thioanisole. However, addition of redox-inactive metal salts (Na⁺, Mg²⁺, Ca²⁺, Al³⁺, Sc³⁺, Y³⁺, and Yb³⁺) leads to more efficient catalysis, displaying accelerated oxygenation rates and an overall increase in conversion and yield.¹⁸⁰ Stoichiometric studies with 21 and Al³⁺ likewise showed a moderate increase in activity. Mechanistic studies suggested initial formation of an Scentered radical via metal-coupled electron transfer (MCET), a well-established mechanism for high-valent terminal metaloxo species involving Fe and Mn (vide infra).¹⁶⁴ In the same report,¹⁸⁰ the stoichiometric oxygenation of triphenylphosphine by 21 afforded a direct comparison of the rate of oxygenation across a diverse series of Lewis acidic metals. A plot of log(rate) vs the pK_a of the metal-aqua species based on the reported data displays a clear linear correlation (Figure 5C).¹⁸⁰ In subsequent studies by the same authors, the activity of high-valent Mn-oxo species toward the epoxidation of cyclooctane was enhanced with the increasing Lewis acidity of redox-inactive metal additives.^{181–183} A graphical analysis, however, could not be performed since quantitative rate data were not reported. In this system, the catalytically competent species is presumed to contain a $[Mn]-O-LA^{n+}$ moiety (LA = Lewis acid) in analogy to other published systems. However, the exact binding mode of the Lewis acid in these complexes has not been corroborated experimentally.

The presence of Lewis acids was also found to affect the O atom transfer reaction from 17 to sulfides. The rate of oxidation of thioanisoles by 17 is greatly accelerated (>10²-fold) by Sc^{3+} .^{184,185} In fact, the mechanism of oxidation was found to shift from direct O atom transfer in the absence of Sc^{3+} (a $2e^{-}$ step), to MCET followed by fast O atom transfer (two 1 e^{-} processes) in the presence of Sc^{3+} . This effect was ascribed to the increased driving force for electron transfer resulting from the positive shift in reduction potential of the Fe complex upon binding of Sc^{3+} . When this driving force is lowered, e.g., in reactions with more electron-poor thioani-soles, the extent of rate acceleration by Sc^{3+} is diminished.¹⁸⁵

ET to the Fe complex was found to be the rate-limiting step in the sulfide oxidation reaction,¹⁸⁴ which is in contrast with the mechanism of sulfide oxidation via proton-coupled electron transfer (PCET) reactions, where protonation of the metal-oxo complex was found to be rate-determining.^{184,186} Oxidation of thioanisole with the analogous Mn complex **18** was also reported; binding of Sc³⁺ to **18** resulted in a marked increase in the rate of the reaction (~2200 fold), consistent with a change from direct OAT to a mechanism that involves electron transfer followed by transfer of the oxygen atom (MCET-OT).¹⁶⁵

Oxygen atom transfer to PPh_3 was reported with Mn-corrolazinato complex 13. ¹⁸⁷ This reaction is influenced both by coordination of axial donors to the manganese metal center¹⁸⁸ and by tautomerization of 13 to the radical cation $[(TBP_8Cz^{\bullet+})Mn^{V}(O)]$.¹⁸⁹ As discussed above, addition of Zn²⁺ induces valence tautomerism to the corresponding complex 13-Zn (Figure 3C).¹⁵⁰ While in the thus far described examples, Lewis acid binding resulted in an increase in OAT, somewhat surprisingly, here a decrease $(>10^3)$ in the rate of OAT from 13 to PPh₃ is observed upon binding of Zn^{2+} or other Lewis acids (i.e., H^+ and $B(C_6F_5)_3$).¹⁹⁰ Mechanistic studies indicated that OAT from $[(TBP_8Cz^{\bullet+})Mn^{IV}(O)-Zn^{2+}]$ to PPh₃ proceeds via the same pathway as for [(TBP₈Cz)- $Mn^{V}(O)$], involving direct nucleophilic attack of PPh₃ on the electrophilic Mn-oxo moiety. Additional investigations also revealed that oxygen atom transfer is influenced by the steric environment of the corrazoline ligand and substitution pattern of the triphenylphosphine. Electrochemical experiments indicated that the reduction potential of 13 shifts positively by ~0.70 V upon binding of Zn^{2+} . Nonetheless, the reduction potentials of -0.05 V and +0.69 V (vs SCE) for 13 and 13-Zn, respectively, are not sufficiently positive to enable one-electron transfer from PPh₃ ($E_{ox} = +2.97$ V vs SCE),^{190,191} ruling out a shift to an ET-OT mechanism as observed in the case of the stronger 1e⁻ oxidants 17 and 18. It is speculated that the two-electron electrophilicity for $[(TBP_8Cz^{\bullet+})Mn^{IV}(O)-Zn^{2+}]$ is reduced by Zn²⁺ binding (potentially due to steric effects), resulting in a slower concerted OAT.¹⁹⁰ DFT calculations and experimental studies furthermore suggest that the spin state of the Mn-oxo moiety, which is known to influence reactivity,^{192,193} might also be responsible for the observed behavior in this system.¹⁹³

An alternative approach to binding of redox-inactive ions to metal-oxo motifs under dynamic equilibrium binding conditions is to provide a dedicated binding site for redoxinactive metal ions on the same multidentate ligand scaffold.¹⁹⁴ This approach has been successfully employed in complexes bearing high-valent Mn^V-oxo moieties that are supported by tetraanionic macrocyclic ligands (TAML; 22-M; Figure 5A). Generally, Mn^V-oxo species supported by TAML ligands display diminished reactivity due to the stabilization of the high-valent Mn center by the electron-rich framework.¹⁹⁵ The presence of a pyridine donor in the TAML ligand in 22, however, allows binding of a wide variety of redox-inactive metal ions (Na⁺, Li⁺, Ba²⁺, Zn²⁺, Mg²⁺, and Sc³⁺) to a secondary site, which drastically affects its reactivity in oxygen atom transfer.¹⁹⁴ Binding of Lewis acids to the backbone pyridine site, confirmed via infrared (IR) spectroscopy, accelerates the rate of PPh3 oxygenation upon addition of the above-mentioned redox-inactive metals, and our analysis showed that the log(rate) of PPh₃ oxidation correlates linearly with the Lewis acidity of the respective ions (Figure 5D).



Figure 6. Activation of $[(TMC)Fe^{II}]$ by dioxygen and the linear correlation between the Lewis acidity of redox-inactive metal ions and reactivity from the resulting iron-peroxo species $[(TMC)Fe^{II}(O_2)]$. (A) Reactivity of $[(TMC)Fe^{II}]$ (25) in the presence of redox-inactive metals and dioxygen and subsequent transformations upon oxidation and/or reduction of the resulting iron-peroxo species. (B) Linear correlation between the Lewis acidity of redox-inactive metals in 26-M (M = Ca²⁺, Sr²⁺, Zn²⁺, Lu³⁺, Y³⁺, and Sc³⁺) and the rate of (i) electron transfer with bromoferrocene (blue dots/traces), (ii) hydrogen atom abstraction from cyclohexadiene (black dots/traces), and (iii and iv) deformylation of 2-phenylpropionaldehyde (red dots/traces) or cyclohexane carboxaldehyde (green dots/traces). Figure 6B is reproduced with permission from ref 229. Copyright 2017 Wiley.

When a sufficiently strong Lewis acid is used (e.g., Sc^{3+}) with an analogous TAML system lacking the peripheral pyridine motif, binding of redox-inactive ions is proposed to occur at the one of the amide carbonyl moieties of the macrocyclic ligand (**23-Sc; Figure 5A**).¹⁵² Binding of Sc^{3+} greatly increases the rate (>10⁴) of oxygen atom transfer to PPh₃ and shifts the potential for oxidation by ~0.70 V as measured via spectrophotometric titration. In a related Co^{III} – oxyl complex, binding of Sc^{3+} , Ce^{3+} , Y^{3+} , and Zn^{2+} is proposed to occur at the Co–oxo moiety.¹⁵¹ This binding results in valence tautomerism to a $[Co^{IV}-O-M^{n+}]$ species (**24**, Figure **5**A) and in acceleration of OAT rates to thioanisoles, with the more Lewis acidic ions engendering the largest rate acceleration.¹⁵¹

Oxygen atom transfer from multimetallic clusters has also been reported. The well-defined nature of these clusters is a key advantage in studying the effects of redox-inactive metals on reactivity over strategies that rely on dynamic binding of Lewis acidic metal ions in solution. Heterometallic cubane clusters $[LMn_3MO_4]$ (10-M; M = Sc³⁺, Mn³⁺, Gd³⁺, Co³⁺) react with trimethylphosphine (PMe₃) to generate the corresponding trioxo complexes [LMn₃MO₃] via transfer of a single O atom to PMe₃. The most Lewis acidic apical metal (Mn^{3+}) displayed the fastest rate of OAT. Intriguingly, the oxidation state of the basal manganese metal centers also influenced the rates of OAT: whereas OAT occurs readily with complexes possessing [Mn^{III}Mn^{IV}₂] core, 1e⁻ oxidation to [Mn^{IV}₃] shuts down oxygen atom transfer almost completely, even in the presence of strongly Lewis acidic Sc³⁺. Computational studies revealed large kinetic barriers to dissociation of bridging carboxylates from Mn^{IV} ions. Since carboxylate dissociation is required for substrate access to the cluster O atoms and was calculated to be the first step of the OAT process, slower dissociation rates unsurprisingly lead to slower OAT processes. Labeling studies with ¹⁸O confirmed that OAT occurs from one of the "top" μ_3 -oxygen atoms-near the dissociating carboxylate on the cubane face-and that the resulting trioxo species are structurally dynamic, resulting in ¹⁸O scrambling and isolation of complexes in which the "empty" oxo site is located at the bottom of the cluster.

The more oxidizing oxo-hydroxo clusters 11-M exhibited much faster rates of OAT. Similar to their cubane analogues, these clusters displayed OAT rates that were dependent on the Lewis acidity of the redox-inactive metal, where the most Lewis acidic metal (Y^{3+}) displayed faster rates of phosphine oxidation. It is perhaps unsurprising that these tetranuclear metal complexes display faster OAT, since the μ_2 -oxygen atom is much more accessible and does not require dissociation of any additional ligands. In contrast to the cubane series, however, oxygen atom transfer is not reversible in these metal clusters, and the metal-containing products of these reactions were not characterized. OAT reactivity of the corresponding iso-structural iron complexes (12-M) was analogously affected by the Lewis acidity of the redox-inactive metal¹¹¹. Overall, when comparing Fe₃ and Mn₃ complexes that incorporate redox inactive metals of similar Lewis acidity (e.g., Gd³⁺, Y³⁺, or La^{3+} , which possess similar metal-aqua pK_a's) OAT reactivity is in line with the one-electron reduction potentials of the metal clusters, with more positive reduction potentials correlating with faster OAT rates, though not sufficiently oxidizing to support single electron transfer chemistry.¹¹¹

As indicated by the reports highlighted in this section, the OAT reactivity of redox-active complexes in the presence of Lewis acidic additives has been studied extensively, with clear correlations emerging between reaction rates and Lewis acidity, providing a key tool for modulating the reactivity of highvalent metal complexes in O atom transfer reactions. As binding of cations to metal-oxo motifs results in marked effects on reduction potential, significant shifts are often engendered in the mechanism of group transfer reactions and dramatic effects on rate have been observed. Many of the discussed transformations for which mechanistic studies are available, including thioanisole and phosphine oxygenation, share a similar rate-determining step, MCET, that is analogous to well-established PCET processes. These results highlight the ability of redox-inactive moieties to profoundly affect the transfer of oxygen atoms even without direct participation in the transfer of electrons.



Figure 7. Heterobimetallic effects in the activation of dioxygen. (A) Reactivity of complexes 14 and 15 with dioxygen in the presence of redoxinactive metals. (B) Formation of complex 31, upon reduction of superoxo species 28 with potassium metal, followed by transmetalation by zinc. (C) O–O bond cleavage in 32 mediated by Sc^{3+} . (D) Reactivity of monometallic copper complex 34 with dioxygen in the absence and presence of Sc^{3+} . (E) Heterobimetallic Co porphyrin complexes with a variety of bound redox-inactive metals, based on bis-porphyrin scaffold with a biphenylenyl (36-M) and anthracene (35-M) backbone.

4. LEWIS ACID EFFECTS ON O₂ ACTIVATION

Electron transfer coupled with dioxygen activation frequently results in the formation of highly reactive metal-superoxo, peroxo, or -oxo species within the active sites of many enzymes.^{3,196–201} As a result, significant efforts have been undertaken to synthesize metal complexes that bear such moieties in order to better understand the chemistry behind activating and/or breaking the O - O bond.^{9,15-17,127,128,134,136,139-141,143,144,202-220} As discussed for ET and OAT, Lewis acids can significantly influence the chemistry behind dioxygen activation. For instance, it has been demonstrated that many Lewis acids can form adducts with the superoxide $(O_2^{\bullet-})$ anion in solution.^{221,222} Binding of Lewis acids to this radical anion promotes electron transfer from metal complexes to O_{22} a process that has been extensively investigated using $[(TPP)Co^{II}]$ (TPP = tetraphenylporphyrin dianion) as the reductant.^{62,223} In the absence of Lewis acids, $[(TPP)Co^{II}]$ does not engage in ET with O₂, in good agreement with its positive reduction potential (E_{ox} ([(TPP)-Co^{III/II}]) = 0.35 V, E_{red} ($O_2^{0/\bullet-}$) = -0.86 V vs SCE).^{223,224} Addition of redox-inactive metal salts (e.g., Sc³⁺) results in 1e⁻ reduction of O₂ as monitored by UV-vis and EPR spectroscopies. The rate of electron transfer increases dramatically as the Lewis acidity of the metal-ion is increased. A plot of log(k) vs metal-superoxide binding energy reveals a striking linear correlation, which highlights the rate-accelerating effects of stronger binding of M^{n+} ions to superoxide.^{62,223,225} Similar effects have been observed in ET between metal complexes and organic electron acceptors (e.g., quinones).⁶² As a result of these studies, the metal-superoxide binding energy has been implemented as a quantitative measure of Lewis acidity in many subsequent investigations

especially by the groups of Fukuzumi and Nam. 55,62,149,153,159,160,184,221,223,226-230

Lewis acids also coordinate directly to metal-superoxo, and -peroxo species, facilitating their isolation and character-ization.²³¹⁻²³⁴ In particular, the reactivity of metal-peroxo complexes with the TMC ligand has been investigated (Figure 6). $^{227-229,235,236}$ It has been shown that superoxide complex $[(TMC)Fe^{III}-(O_2^{\bullet-})]$ (26) is metastable in solution, but decays to Fe-oxo complex $[(TMC)Fe^{IV}(O)]$ (27) in the presence of both a reducing agent and a source of protons (Figure 6).²³⁷ Similar reactivity can be induced by Lewis acids $(Sc^3 \text{ or } Y^{3+})$ in the presence of a reducing agent $(Ph_4B^- \text{ or }$ Fc).^{227,236} Upon conversion to 27, a distinct intermediate was observed by UV-vis spectroscopy that is stable at -40 °C for several days. Based on characterization via Raman, EPR, ESI-MS, and EXAFS spectroscopy, this intermediate was unequivocally assigned as $\text{Fe}^{\text{III}} - (\mu - \eta^2 : \eta^2 \cdot O_2^{2^-}) - M^{3^+}$ (26-M, M = Sc or Y), where the Lewis acid is bound in a side-on fashion to the metal-peroxo moiety. Binding of the Lewis acid is crucial for the observed reactivity: in the absence of Lewis acids, reduction of 26 does not lead to O-O bond cleavage and formation of the high-valent Fe^{IV}-oxo species.^{227,236} The rate of reduction is dependent on the nature of the Lewis acid, as the more Lewis acidic Sc³⁺ reacts three times faster than the corresponding Y³⁺ analogue. Furthermore, screening of a series of reductants shows that the ET reorganization energy is not affected by the presence of a second metal ion, indicating that the differences in ET rate between 26 and 26-M are derived solely from the positive shift in reduction potential (estimated from the same analysis to be ~ 0.24 V).

In subsequent studies, the reactivity and redox properties of complexes **25** and **26** were reported in further detail.^{228,229} It

was found that upon photoirradiation of an O2-saturated solution containing 25, dimeric N-benzyl-1,4-dihydronicotinamide $[(BNA)_2]$, and a redox-inactive metal salt, different products are formed depending on the nature of the added Lewis acid.²²⁸ With Ca²⁺ and Sr²⁺, the Lewis acid-bound ironperoxo species 26-M is formed, whereas with Zn^{2+} , Lu^{3+} , Y^{3+} , and Sc^{3+} the high-valent iron-oxo species 27 is generated. As previously discussed, the latter species is only formed after reduction of $[(TMC)Fe^{III}(O_2^{\bullet-})]$, which is followed by O–O bond cleavage.^{227,236} Clearly, the redox properties of the intermediate 26-M are modulated by the Lewis acid. These observations are consistent with the 1e⁻ reduction potentials of the independently synthesized complexes 26-M, which correlate linearly with the Lewis acidity of the redox-inactive metal ions (see Section 2). Addition of $[(BNA)_2]$ to these complexes resulted in reduction of 26-M only when $M = Zn^{2+}$, Lu^{3+} , Y^{3+} , or Sc^{3+} , establishing that the Lewis acidity of the redox-inactive metal directs the reactivity of complex 25.228,229

An effect of Ca2+ and Sr2+ on dioxygen activation was observed in Fe and Mn complexes supported by the MST tripodal framework (14 and 15, respectively).^{122,123} In the absence of any additives, O₂ activation by these complexes is sluggish and no oxidized species could be isolated following treatment with O2. Upon addition of Group 2 metals, remarkable rate enhancements were observed for Ca²⁺ and Sr^{2+} (~30-40-fold), whereas for Ba^{2+} only modest (~10-20fold) rate enhancements were observed. A clear difference between Fe and Mn complexes was also noted, as the rate of dioxygen activation differs by 1 order of magnitude in favor of Fe.^{122,123} In this system, the role of the secondary coordination sphere is pivotal in the rate acceleration by these Lewis acids. Not only do the sulfonamido moieties serve as hydrogen bonding acceptors to the generated M^{III}-OH species (M = Mn and Fe), but they can also coordinate the group 2 metal ions, establishing a unique M^{II} -(μ -OH)-M' core (M = Fe and Mn, and M' = Ca^{2+} , Sr^{2+} , and Ba^{2+}) that is in several aspects akin to the OEC (Figure 7A). It is proposed that initial binding of these Lewis acids to [M^{II}MST]⁺ complexes facilitates electron transfer to O2, generating an intermediate MII-(superoxo)-M' species, which-via multiple steps-decomposes to the final M^{III}-OH species.¹²³ Although no definite experimental evidence is provided for the existence of such superoxo intermediates, the Lewis acid-induced rate enhancements are consistent with the previously discussed 1e⁻ O₂ activation by cobalt porphyrin complexes in the presence of Lewis acids (*vide supra*), where superoxo intermediates have unequivocally been identified.^{62,223} Cobalt complexes supported by analogous ligand scaffolds have also appeared in the literature, although no reactivity with O_2 was reported.²³⁸

Addition of redox-inactive metals has also been observed to affect the reactivity of other metal—peroxo and —hydroperoxo species. Peroxo-cobalt(III) compound $[Co^{III}((TBDAP)-(O_2^{2^-})]^+$ (TBDAP = N,N'-di-*tert*-butyl-2,11-diaza[3.3](2,6)pyridinophane) reacts with nitriles to form a hydroximato species. In the presence of redox-inactive metals a peroxyimidatocobalt(III) species is isolated instead,²³⁹ and the logarithm of the rate of this reaction is linearly correlated to the Lewis acidity of the redox-inactive metals. In contrast, more Lewis acidic trivalent redox-inactive metals were found to hinder nucleophilic reactivity of a separate alkylperoxocopper-(II) system with cyclohexylcarboxaldehyde.²⁴⁰ Addition of Sc³⁺ (or HClO₄) also enables rapid hydroxylation of cyclohexane and benzene by a mixture of $[Fe(\beta-BPMCN)(CH_3CN)_2]^{2+}$ (BPMCN = N,N'-bis(pyridyl-2-methyl)-N,N'-dimethyl-trans-1,2-diamino-cyclohexane) and H_2O_2 , for which it is hypothesized that the Lewis acidic additives enhance O–O bond cleavage to yield a highly reactive Fe^V–oxo intermediate.²⁴¹

Activation of O2 with complexes that form distinct homobimetallic $[M-(\mu-O)_2-M]$ cores (M = Fe, Ni, andCu) have been extensively investigated, and their chemistry has been described in a series of excellent re-views.^{3,9,16,17,207,211,213,214,216,242} Heterobimetallic examples are more rare, ^{15,243,244} especially those containing redox-inactive metals.²⁴⁵ A prominent example is the $\mu - \eta^2$: η^2 peroxo-bridged β -diketiminate (L) complex [LNi-(O₂)- $K(THF)_n$ (29), generated upon reduction of the superoxo complex $[LNiO_2]$ (28) with potassium metal.²⁴⁵ While nickel-potassium complex 29 is a stable peroxide species, transmetalation of potassium with the more Lewis acidic Zn²⁺ (supported by an additional diketiminate ligand) results in fast H atom abstraction to generate the complex $[LNi-(\mu-OH)_2-$ ZnL] (31), featuring two bridging hydroxide ligands. The differences in reactivity between complex 29 and the putative $[LNi-(O_2^{2-})-ZnL']$ (30) intermediate generated upon addition of $[LZn^{2+}]$ were explained based on DFT calculations. For peroxo complex 30, a triplet ground state was predicted with significant spin density on the bridging oxo-ligands, indicating (at least partial) O-O bond scission. In contrast, the O-O bond in 29 remains intact, and the ground state is a singlet. Exchange of K⁺ for the more Lewis acidic Zn²⁺ thus results in a higher degree of O_2 activation, from $(\mu - O_2^{2^{-}})$ to $(\mu$ -O²⁻)₂, generating a [LNi-(O)₂-ZnL] intermediate, which is only transient and rapidly undergoes HAT (from the solvent or other exogenous sources) to give 30. Replacement of Zn^{2+} in these systems with Fe^{2+} , Co^{2+} , Ni^{2+} , or Cu^{2+} resulted in nucleophilic reactivity pathways (vs electrophilic reactivity for Zn),²⁴⁶⁻²⁴⁸ highlighting that despite similar Lewis acidities (see Table 1), redox-inactive metals can engender distinct reactivity manifolds from their redox-active counterparts in dioxygen transformations including O-O bond cleavage.

In a related dinuclear copper system, the degree of O₂ activation was also found to be dependent on the presence of Lewis acid ions.²⁴⁹ Exposing $[(MeAN)Cu^{I}][BAr_{4}^{F}]$ (MeAN = *N*-methyl-*N*,*N*-bis[3-(dimethylamino)propyl]amine) to O₂ at low temperatures results in formation of a stable peroxodicopper(II) complex (32). Complex 32 displays a characteristic $[M-(\mu-O_2^{2^-})-M]$ core and exhibits no discernible reactivity toward external substrates.²⁵⁰ However, in the presence of Sc³⁺, 32 generates a species whose spectroscopic features are consistent with a dioxo-bridged copper(III) dimer (33), where the O–O bond has been fully cleaved.²⁴⁹ The presence of the well-precedented $[Cu^{III}-(\mu-O)_2-Cu^{III}]$ core was confirmed by resonance Raman spectroscopy and supported by DFT calculations. In contrast to several of the previously discussed peroxo- and bis- μ -oxo-complexes, in which coordination of the relevant redox-inactive metal ions to the O2-derived ligands is typically observed, calculations on this dicopper system suggested that scandium does not coordinate via the oxygen atoms, but rather to one nitrogen donor from each MeAN ligand.²⁴⁹ Sc³⁺ coordination thus promotes dissociation of one of the nitrogen ligands from copper, resulting in a change in the coordination geometry at Cu (Figure 7B). This change influences the orbital overlap between the MeAN nitrogen donors and the copper d-orbitals; the resulting increase in electron density at Cu enables a higher degree of backbonding to the σ^* orbitals on the ${O_2}^{2-}$ motif,

which facilitates O–O bond scission. A unique feature of this system is that the O–O bond can be reformed upon sequestering the Lewis acid with other bidentate, nitrogen-based ligands, restoring the original coordination geometry of the copper complex. Along with influencing the reversible formation and cleavage of the O–O bond, the binding of Sc³⁺ also alters the reduction potential of the dioxo-bridged copper(III) complex **33**, which in contrast to peroxodicopper(II) complex **32**, is more reactive toward weak C–H and O–H bonds.²⁴⁹

In other copper complexes, oxygen reduction to water (a 4e⁻ process) or to peroxide (a 2e⁻ process) can be regulated by a variety of redox-inactive metal ions.²⁵¹ In the absence of a Lewis acid, O2 reduction to H2O was observed with $[(tmpa)Cu^{II}(MeCN)][ClO_4]_2 (34; tmpa = tris(2$ pyridylmethyl)amine) in the presence of decamethylferrocene (as sacrificial reductant) and triflic acid (as H⁺ source; Figure 7D). This four-electron reduction process proceeds via the [(tmpa)Cu^{II}(O₂²⁻)Cu^{II}(tmpa)]²⁺ peroxide intermediate.²⁵² Performing the same reaction in the presence of Sc^{3+} does not yield H₂O; rather, formation of a $[Sc(O_2^{2-})]^+$ species is indicated by EPR spectroscopy.²⁵¹ Kinetic studies suggest that, unlike in the case of O_2 reduction to water, no bis(copper-(II))-peroxo dimer is formed, and a monomeric superoxocopper complex $[(tmpa)Cu^{II}(O_2^{\bullet-})]^{\scriptscriptstyle +}$ is generated instead.^{251} This copper(II)-superoxo complex is rapidly reduced by decamethylferrocene in the presence of Sc^{3+} to produce $[Sc(O_2^{2-})]^+$ and regenerate 34. Other Lewis acids also facilitate an analogous two-electron reduction to the corresponding metal-peroxide. The rates of these reactions correlate with the Lewis acidity of the redox-inactive metal, where the most Lewis acidic metal ion gave the highest activity.²⁵¹ A quantitative measure of these effects (e.g., a linear correlation), however, could not be generated due to the absence of quantitative rate data in the original publication.

Analogous four- and two-electron reduction processes involving dioxygen were also investigated with a series of a cofacial bisporphyrin "Pacman" complexes containing one Co^{II} center to enable dioxygen binding, along with a Lewis acidic metal in the second porphyrin to control O_2 reduction chemistry (35-M and 36-M, Figure 7E).^{253,254} The catalytic activity of each heterobimetallic complex was compared to that of the known homobimetallic catalyst bisporphyrin-Co2, which facilitates the $4e^{-1}$ reduction of O_2 to H_2O .²⁵⁵ It was proposed that, in this system, dioxygen binds to the mixed valent state of the catalyst (bisporphyrin $-Co_2^{III}Co^{II}$) at the Co^{II} center, and that the second Co site (Co^{III}) acts as a Lewis acid to facilitate the overall reductive transformation. The reactivity of the isostructural heterobimetallic complexes was studied by rotating-disk voltammetric methods;²⁵³ from these studies, complexes 35-M (M = Ti^{4+} , Ga^{3+} , In^{3+} , Lu^{3+} , Sc^{3+}) were found to reduce dioxygen at potentials close to that for Co-monoporphyrin at +0.55 V (vs NHE), with a limiting current indicating a two-electron mechanism.²⁵³ In contrast, complexes 35-Lu, 36-Lu, and 36-Sc reduce dioxygen at a potential close to +0.7 V (~100 mV more negative compared to the homometallic $[Co]_2$ catalyst and 300 mV more positive than a Co-monoporphyrin complex). Moreover, their limiting currents are larger than those observed for the Comonoporphyrin, and only slightly smaller than for the $[Co_2]$ complex. Taken together, these observations suggest that, while bisporphyrin-Co₂ selectively reduces O₂ by fourelectrons to generate H₂O, 35-Lu, 36-Lu, and 36-Sc might

reduce O₂ via two concurrent but separate 2- and 4e⁻ mechanisms to produce H₂O₂ and H₂O, respectively. H₂O₂ was indeed detected at the ring electrode upon dioxygen reduction in the latter cases, providing additional support for the operability of the 2e⁻ pathway.²⁵³ Together with the nature of the Lewis acid, the distance between the metal centers in these bimetallic assemblies also appears to be an important factor. While the identity of the redox-inactive metal played a key part in the observed transformations, the exact role of these metal ions in directing reactivity and enabling different mechanistic pathways for O2 reduction (e.g., 4- vs 2e⁻ reduction) remains unclear. These studies nonetheless demonstrate that, together with accelerating O2 activation, Lewis acidic metal ions are also able to control mechanistic aspects of O₂ reduction, engendering a shift between four- and two-electron reduction processes.

5. LEWIS ACID EFFECTS ON O₂ EVOLUTION

The role of Lewis acids in O₂ evolution is a fundamental aspect of the mechanism of water oxidation by the OEC,^{67,250} which contains calcium and manganese ions $(CaMn_4O_4)$ in a cuboidal arrangement in its active site.^{132,259-263} The influence of Lewis acids on water oxidation has mainly been studied on heterogeneous surfaces of mixed-metal oxides, which have attracted significant interest as promising water oxidation catalysts.²⁶⁴ In 2008, a heterogeneous cobalt-oxide water oxidation catalyst was reported that self-assembles from Co^{II} ions in solution.²⁶⁵ Energy-dispersive X-ray (EDX) analysis of the electrodeposited film indicated that, besides cobalt, potassium and phosphate are also present in substantial amounts. Based on additional X-ray absorption near-edge structure (XANES) and extended X-ray absorption fine structure (EXAFS) studies, it was suggested that potassium might be involved in forming distinct KCo₃O₄ cubanes,²⁶⁶ although other studies have debated such a role for potassium.²⁶⁷ The influence of redox-inactive metals on water oxidation has been studied more extensively in manganese oxides that are structurally similar to the birnessite family of minerals.³⁶⁻⁴⁰ These manganese oxides form layers of edge-sharing MnO₆ octahedra, in which redox-inactive metals can intercalate.³⁷ Studies on these synthetic mixed-manganese oxides revealed that, in the presence of redox-inactive metals, a marked increase in O_2 evolution was observed compared to plain manganese oxide.^{36,37} The O_2 evolution activity followed the trend $Ca^{2+} > Sr^{2+} \gg Mg^{2+} \sim Na^+$, very similar to the activity profile of the OEC. A similar trend was also observed upon incorporating Ca²⁺ and Sr²⁺ in polymeric cobalt cyanide complexes.²⁶⁸ Computational studies into the electronic structure of various redox-inactive metal-substituted birnessites suggested that a suitable band gap for water splitting was observed when Ca^{2+} or Sr^{2+} were intercalated between the manganese oxide layers.²⁶⁹ Incorporation of redox-inactive metals into Ni-Fe oxides have also been demonstrated to enhance their water oxidation capabilities.^{270,271}

In contrast to heterogeneous materials, fewer reports have described in detail the effects of Lewis acids on O_2 evolution from molecular complexes. In one of the few detailed studies, the role of Ca^{2+} and Sr^{2+} in enabling O_2 evolution from Fe– peroxide complexes **26-M** was investigated. As discussed in Section 2, potentials required for both reduction and oxidation of complexes **26-M** depend on the Lewis acidity of the redoxinactive metal M; while reductive reactivity, facilitated by *more* Lewis acidic cations (Zn²⁺, Lu³⁺, Y³⁺, Sc³⁺), leads to generation

of Fe^{IV}-oxo complex 27, oxidation of 26-M is inaccessible at moderate potentials except in the presence of the *less* Lewis acidic cations Ca²⁺ and Sr²⁺. Such a unique behavior for Ca²⁺ and Sr²⁺ is reminiscent of the OEC in PSII, where only in the presence of these two redox-inactive metals is O₂ evolution observed.²⁵⁸ However, as Fe^{III}-peroxide complex 26 can also be oxidized to release O₂ in the absence of *any* redox-inactive metals, it is possible that both in these systems and in the OEC, the reduction potentials engendered by Ca²⁺ and Sr²⁺ for complex/cluster oxidation are sufficiently negative as to not *prevent* O₂ release, rather than actively *promote* it.²²⁸

In an attempt to further elucidate the factors governing the structure and reactivity of the OEC, the effects of water coordination to redox inactive metals in 26-M (M = Ca^{2+} , Zn^{2+} , and Sc^{3+}) were investigated, ²³⁵ as water is known to play a prominent role in the primary coordination sphere of the OEC. $^{132,259-263}_{\rm Addition}$ Addition of water (0–2.8 M) to 26-Zn resulted in a marked change in the UV-vis spectrum of the complex that was found to be consistent with binding of two water molecules to Zn²⁺. A similar value was found for Ca²⁺ but not for Sc^{3+} , where addition of water resulted in formation of redox-inactive metal-free 26, indicating loss of Sc³⁺ The coordination of water to Zn²⁺ had marked effects on the physicochemical properties of the resulting iron-peroxo species $[(TMC)Fe^{III} - (\mu - \eta^2; \eta^2 - O_2) - Zn^{2+}(OH_2)_2]^{.235}$ Most notably, the binding of water resulted in a decrease of the Lewis acidity of Zn^{2+} , estimated from the g_{zz} values determined by EPR spectroscopy in accordance with the previously utilized approach for determination of Lewis acidities by EPR.⁶² The "observed" Lewis acidity for Zn^{2+} under these conditions in the presence of water (0.57 eV), is more in line with those reported for Ca^{2+} (0.59 eV) and Sr^{2+} (0.53 eV), and is reflected in the reactivity $[(TMC)Fe^{III} - (\mu - \eta^2: \eta^2 - O_2) - Zn^{2+}(OH_2)_2]$, which is akin to that observed for $[(TMC)Fe^{III}-(\mu-\eta^2:\eta^2-\eta^2)]$ O_2)-Ca²⁺] (26-Ca): indeed, while CAN does not oxidize $[(TMC)Fe^{III} - (\mu - \eta^2 \cdot \eta^2 - O_2) - Zn^{2+}]$ to give 25 and $O_{2\mu}$ $[(TMC)Fe^{III}-(\mu-\eta^2:\eta^2-O_2)-Zn^{2+}(OH_2)_2]$ readily evolves one equivalent of O2 upon one-electron oxidation. The coordination of water thus modulates the Lewis acidity of Zn^{2+} leading to a change in the redox properties of the corresponding iron-peroxo complex, resulting in reactivity that is analogous to that reported for the related Ca²⁺ and Sr²⁺ complexes in the absence of coordinated water.²³⁵ Coordination of water also induces a negative shift in the reduction potential of $[(TMC)Fe^{III}-(\mu-\eta^{\tilde{2}}:\eta^2-O_2)-Zn^{2+}(OH_2)_2]$, and results in lower ET rates from 1,1'-dimethylferrocene. It is remarkable that a single system has been successfully leveraged to probe both the influence of redox-inactive metals and of water in the (secondary) coordination sphere on both O₂ activation and evolution, all factors and contexts that are proposed to be highly relevant in tuning the structure and reactivity of the OEC in PSII.

The effect of Group 2 metals on O_2 evolution by the inorganic oxidant potassium ferrate (K_2FeO_4) has also been studied. At low pH, potassium ferrate readily oxidizes water to dioxygen, in part due to its positive potential for oxidation.^{272,273} The stability of K_2FeO_4 is pH dependent, ^{272,273} and at pH 9–10 minimal decomposition is observed (2% h⁻¹) in the absence of any redox-inactive metal additives.²⁷⁴ However, addition of excess Ca^{2+} to solutions of K_2FeO_4 at pH 9–10 resulted in O_2 evolution from the reaction mixture. Substituting Ca^{2+} for Sr^{2+} or Mg^{2+} did not change the yield or kinetics of the O_2 evolution reaction. A DFT study

into the mechanism of this reaction suggests mainly a structural role for Ca²⁺ (or Sr²⁺ or Mg²⁺), where a bound Ca²⁺ ion bridges two ferrate anions, bringing two distinct oxygen substituents in close proximity, facilitating O–O bond formation, a mechanism supported by isotopic labeling experiments. A similar mechanism was proposed for water oxidation by K₂FeO₄ under acidic additions, as proton-induced dimerization of FeO₄²⁻ results in O₂ evolution.²⁷²

Overall, even the limited studies that are available suggest that Ca^{2+} and Sr^{2+} occupy a special niche in facilitating O_2 evolution, consistent with observations of the OEC, and a growing body of work has begun to outline the potential modes of their participation in this key transformation. Nonetheless, additional studies are necessary both on synthetic complexes and on the OEC itself to further elucidate the exact role that these metal ions surprisingly have on O_2 evolution.

6. LEWIS ACID EFFECTS ON HYDROGEN ATOM TRANSFER

H atom transfer (HAT) reactions are closely linked to O_2 derived reactivity, including biologically essential organic substrate activation by metal-oxo species. HAT reactions represent another set of processes on which heterometallic effects have been extensively investigated. Many of the complexes described in the previous sections are also active for HAT, and where appropriate some of the effects of redoxinactive metals have already been alluded to in conjunction with their other properties. A handful of additional systems have not yet been discussed and will be the focus of this section.

Fe^{III}-peroxo complexes **26-M** have already been discussed in the context of both ET (Section 2) and O₂ activation (Section 4). Further investigations into these complexes also revealed remarkable linear correlations between the Lewis acidity of the redox-inactive metal and the activity of these complexes in HAT reactions, as well as in aldehyde deformylation.²²⁹ Notably, plots of log(*k*) for various transformations (ET, HAT, aldehyde deformylation) vs Lewis acidity reveal linear correlations (Figure 6B); furthermore, the slope of the linear fit in these correlation plots indicates the direction of charge transfer: reactions proceeding via electrophilic mechanisms (ET and HAT) produce a positive slope, whereas nucleophilic mechanisms show the opposite trend.²²⁹

In the series of complexes of the pentadentate ligand dpaq $[(dpaq)Mn^{IV}O]^+ - M^{n+}(19- M, M = Ca^{2+}, Mg^{2+}, Zn^{2+}, Lu^{3+},$ Y³⁺, Al³⁺, Sc³⁺; Figure 3) discussed in Section 2, contrasting observations were reported on the effects of Lewis acidity on the rate of ET/OAT vs HAT reactions. Specifically, while the rate of ET and OAT reactions displayed a linear, positive correlation with Lewis acidity, HAT rates decreased with increasing Lewis acidity. These results were taken as an indication that the Lewis acidity of the redox-inactive metal is involved in modulating the *basicity* of the metal-oxo moiety. The more Lewis acidic ions make the Mn-oxo motif less basic, affecting the concerted HAT to the oxo-motif, resulting in an overall slower reaction. On the other hand, reactions in which HAT proceeds via an initial ET step due to increased driving force for ET, and for which ET is therefore rate-determining, become faster as the reduction potential of the complex shifts to more positive values (i.e., with more Lewis acidic ions).¹⁵⁹ Analogous studies with triflic acid (HOTf), while not involving redox-inactive metals, further indicate that interactions between the Mn-O moiety and Lewis acidic centers (H⁺ or



Figure 8. Hydrogen abstraction properties of various metal complexes upon binding redox inactive metals. (A) Differences in the rate of hydrogen atom abstraction from cyclohexadiene with complex 18 upon binding of one and two scandium ions. (B) Reactivity trends of a manganese corrolazine complex in oxygen atom (OAT) and hydrogen atom transfer (HAT) reactions in the absence (13) and presence of Lewis acids (13-LA; LA = $B(C_6F_5)_{3/}$ HBF₄, and $Zn(OTf)_2$). (C) Cobalt and vanadium complexes involved in HAT reactions.

 M^{n+}) modulate the chemical and physical properties of the Mn–O group.²³⁰ It should be noted that the assignment of the active species in these studies as $[(dpaq)Mn^{IV}O]^+-M^{n+}$ (with Lewis acidic ions interacting directly with the metal–oxo moiety) has recently been challenged for room-temperature studies on the basis of ¹H NMR and electronic absorption spectroscopies, concurrent with a proposed new role for redoxinactive metal ions in promoting protonation of the precursor $[(dpaq)Mn^{III}(OH)]^+$ complex to the corresponding aqua complex $[(dpaq)Mn^{III}(OH_2)]^{2+.275}$

The effects of redox-inactive metals on HAT by Mn complex 18 were studied in more detail than for the isostructural Fe complex 17 (Figure 8).¹⁶⁵ While the addition of Sc^{3+} to 18 resulted in an increase in the rate of OAT, the opposite effect is observed for H atom transfer, where binding of Sc³⁺ results in a decrease in reaction rate. Reaction of 18 with 1,4-cyclohexadiene (CHD) was found to be 5 and 180 times faster than for its 1:1 and 1:2 Sc^{3+} adducts 18-Sc and 18-Sc₂, respectively.¹⁶⁵ This change in reactivity is ascribed to the different steric requirements for the two reactions. As determined via kinetic studies, OAT reaction rates correlate with the ET driving force-i.e. OAT to thioanisole proceeds via a rate-determining ET step.¹⁴⁶ This step is largely uninfluenced by the steric constraints imposed by Sc3+ binding, and is followed by a fast O atom transfer. In the case of H atom transfer, however, significant interaction between substrate and Mn-O species is proposed to be necessary, and binding of Sc³⁺ sterically hinders this interaction leading to reduced reaction rates.¹⁶⁵ A HAT mechanism is thus proposed for 18, yet no kinetic isotope effects are reported for the reaction of 18 with CHD, so a classical HAT step might not be ratedetermining.

In contrast, studies with complex 17 have mainly focused on C–H bond cleavage in toluene derivatives.¹⁸⁴ Complex 17 itself only displayed slow reactivity with toluene analogues, and the large kinetic isotope effect (KIE = 31) suggested a classical H atom transfer mechanism followed by oxygen rebound to yield benzylic alcohols. Addition of Sc(OTf)₃ resulted in ~2–100-fold increase in observed reaction rates.¹⁸⁴ In parallel work using the HOTf in place of Sc(OTf)₃, small KIEs were

observed at high concentrations of HOTf, indicating a shift in mechanism.¹⁸⁴ As for the OAT to thioanisole (vide supra), this new mechanism is believed to involve ET-OT steps, and the reaction rate is thus not susceptible to the sterically hindered environment around the Fe-oxo moiety, as the ET step is likely rate-determining. Interestingly, a shift in mechanism was also observed in the oxidation of a variety of benzyl alcohols. Addition of Sc³⁺ to 17 results in a 120-fold increase in the rate of oxidation of 2,5-dimethoxybenzyl alcohol without any observable KIE. Indeed, mechanistic studies confirmed that this reaction proceeds via a rate-limiting ET step. On the other hand, reduction of benzyl alcohols with more positive oxidation potentials proceeded with large KIEs, suggesting HAT as the rate-determining step. Consequently, the overall mechanism of benzyl alcohol oxidation depends on the oxidation potential of the substrate and discriminates between HAT and PCET (or MCET) pathways.²⁷⁶ Unfortunately, a direct comparison with the HAT properties of 18 is not possible, since hydrogen atom abstraction from CHD with complex 17 is not reported.¹⁸⁴

More detailed studies into the hydrogen atom abstraction reactivity with complexes 13 and 13-Zn were also performed (Figure 8).²⁷⁷ Addition of xanthene (BDE = 74 kcal mol⁻¹) to complexes 13 and 13-Zn resulted in rapid formation of $[(TBP_8Cz)Mn^{IV}(OH_2)]$ and $[(TBP_8Cz)Mn^{IV}(OH)]$, respectively. A clear difference is thus observed upon coordination of Zn to 13. Whereas for 13 two H atom transfers are observed, only a single H atom transfer is reported for 13-Zn. It is proposed that binding of Zn²⁺ to Mn^{IV}-OH species generated by an initial HAT process prevents transfer of a second hydrogen atom. A 4-fold enhancement of reaction rate is also observed upon Zn binding.²⁷⁷ Rate enhancements were also observed upon addition of other Lewis acids such as HBAr^F₄ and $B(C_6F_5)_3$. Kinetic studies revealed that H atom abstraction is the rate-limiting step in the reaction of xanthene with 13 and 13-LA, as both displayed large kinetic isotope effects (KIE = >20; LA = HBAr^F₄ or B(C₆F₅)₃). Compared to 18-M, steric inhibition by the Lewis acids complexes 13-M is negligible.²⁷

A direct HAT mechanism was also proposed for 24-Sc in the presence of xanthene, 9,10- dihydroanthracene (BDE = 76 kcal



Figure 9. Redox state of remote cations affect HAT reactivity of metal-oxo clusters. Oxidation state and charge of remote Fe_3 sites have significant effect on HAT reactivity of terminal $Mn-OH_2$ moiety. Figure reproduced from ref 282. Copyright 2019 American Chemical Society.

mol⁻¹), CHD (BDE = 77 kcal mol⁻¹), and triphenylmethane (BDE = 81 kcal mol⁻¹). Reaction rates correlate linearly with substrate BDE, and a large KIE (13) was observed upon addition of $[D_2]$ -xanthene to **24-Sc**. Although the addition of Lewis acids stabilizes high-valent Co–oxo complex **24-Sc**, HAT to **24** was not reported, so the effect of the Lewis acid cannot firmly be established.

HAT was reported with high-valent cobalt complex $[(TMG_3tren)Co^{IV}(O)Sc(OTf)_3]^{2+}$ (37-Sc) in the presence of 9,10-dihydroanthracene.¹⁴⁸ Compared to the corresponding iodosobenzene adduct $[(TMG_3tren)Co^{II}(OI^sPh)]^{2+}$ (37), the rate of HAT to 37-Sc is much slower. Although no explanation was given for the observed rate difference, the nature of the ligand (TMG_3tren) and the presence of Sc³⁺, might render the reactive Co^{IV}—oxo sterically inaccessible to result in efficient HAT. Nonetheless, further studies are required to determine the mechanism of HAT with 37-Sc. As previously discussed, other studies have suggested that for 37-Sc an assignment of $[(TMG_3tren)Co^{III}(OH)Sc(OTf)_3]^+$ might be more reasonable, which would significantly alter the driving force and mechanism for HAT by this complex.¹⁶²

The effects of redox-inactive metals on HAT to a series of $V^{\rm IV}-{\rm oxo}$ complexes were studied. 278 In particular, for $[(TPA)]V^{IV}(O\hat{)}Cl]~(38)$ the addition of a variety of redox inactive metals (Na⁺, Mg²⁺, Ca²⁺, Ba²⁺, Zn²⁺, Y³⁺, Yb³⁺, Al³⁺, and Sc3+) greatly accelerated HAT from CHD. During the reaction, formation of the peroxide species [(TPA)V^V(O)- $(O_2^{2^-})$]⁺ was observed by UV-vis spectroscopy. Interestingly, both Lewis acid and CHD are necessary in order to observe this intermediate. The redox-inactive metal most likely facilitates the dissociation of the chloride ligand from VIV opening a coordination site on the metal center for binding and subsequent activation of O_2 .²⁷⁸ O_2 activation might also be further enhanced by coordination of the Lewis acid to the V^{IV}-oxo moiety. Upon 1e⁻ oxidation of the metal center, a transiently stable $[(TPA)V^{V}(O)(O_{2}^{\bullet-})]^{2+}$ species is formed, which abstracts a hydrogen atom from CHD to form $[(TPA)V^{V}(O)(OOH)Cl]^{2+}$. This hydroperoxo species decomposes, possibly via multiple pathways, to regenerate [(TPA)- $V^{V}(O)(O_{2}^{2-})]^{+}$, which is again able to abstract a hydrogen atom from CHD to form $[(TPA)V^{V}(O)(OOH)]^{+}$. The latter HAT process is facilitated by Lewis acids, as in their absence $[(TPA)V^{V}(O)(O_{2}^{2^{-}})]$ is incapable of undergoing HAT reactions. Unfortunately, no direct correlation between the rate of HAT and the nature of the Lewis acid could be

established due to the lack of quantitative rate information in published works, although it appears that the conversion of CHD to benzene qualitatively increases as the Lewis acidity of the redox-inactive metal increases.²⁷⁸

These representative examples highlight the importance of redox-inactive metals and their impact on the mechanism of C-H bond cleavage. Whereas Lewis acids typically accelerate the rate of OAT, their effects on C-H bond cleavage are more complex. In systems where concerted H atom abstraction is rate-limiting, rates are often unchanged or even lower in the presence of Lewis acids.¹⁶⁵ In these systems, typically involving substrates with very positive reduction potentials for their 1e⁻ oxidations, large KIEs are observed.^{165,276} Decreases in HAT rates can be ascribed to the presence of the redox-inactive metal sterically restricting access to the high-valent metal-oxo moiety.²⁷⁷ In contrast, rate enhancements are generally observed if a large driving force for ET is present (e.g., for substrates that can be oxidized at sufficiently negative potentials or metal complexes with particularly positive reduction potentials). In these instances, MCET/PCET mechanisms are proposed where ET is the rate-determining step and negligible KIEs are observed.^{184,276} The fine balance between these two mechanisms is elegantly demonstrated in the oxidation of benzyl alcohol by complex 17.²⁷⁶

In contrast, ligand design can restrict the Lewis acid from the site of HAT, thereby preventing steric effects from influencing the observed reactivity. Salophen ligands with crown ether moieties discussed previously have been shown to support terminal Mn-nitride complexes (supported by analogous ligands as 1-Mn, Figure 2).^{80,279} Redox-inactive metals bind exclusively at the crown ether site, and modulate the reduction potentials as well as HAT reactivity of this complex. Addition of Lewis acidic metals (Na⁺, K⁺, Ba²⁺, Sr²⁺, La³⁺, Eu³⁺) significantly affects both the Mn^{VI}/Mn^V reduction potentials (range of >700 mV) and pK_a of the corresponding Mn^{V} -imido complex(by ~9 pK_a units), with a higher charge cation resulting in lower pKa and higher $E_{1/2}$. This compensatory relationship results in a relatively small change in BDFE, spanning a range of only ~6 kcal/mol, as constrained by the Bordwell equation.²⁸⁰ Conceptually related with respect to effects of the charge of remote metal centers, a series of pyrazolate supported $Fe_3MO(OH_x)$ clusters (**39**; M = Mn, Fe) display more significant changes in O-H BDFE (~16 kcal/ mol) upon modulation of charge and oxidation state of the distal iron centers (Figure 9).^{281,282} These results highlight how properties of the entire cluster or bimetallic system, including at distal metal centers, can significantly affect substrate activation and reactivity.

7. CONCLUSIONS

Lewis acidic metal centers can profoundly affect redox processes even without the intrinsic ability to directly participate in electron transfer. Upon coordination to redox active metal complexes, these Lewis acids impact a range of properties, including the reduction potential of transition metal centers, the electrophilicity and nucleophilicity of metal-oxo, -hydroxo, -peroxo, and -superoxo species, and the steric profile of these motifs. Through these effects, redox-inactive metals influence the electron and group transfer reactions of mono- and multimetallic redox-active complexes including the kinetics of ET, OAT, and HAT reactions. These effects can contribute to the enhancement of activity of otherwise unreactive species, prompt shifts in the mechanism of certain transformations, and help direct product selectivity in various chemical reactions. The growing number of examples of these multimetallic effects highlights the general nature of these influences and shows that they can serve as crucial tools to modulate the reactivity of redox-active complexes. Furthermore, many studies have highlighted systematic effects of Lewis acidic metals of potential relevance to biological active sites such as the oxygen-evolving complex and to mixed-metal heterogeneous catalytic systems for water oxidation and O₂ reduction. While many questions about the role played by redox-inactive metals in the actual mechanisms of these complex transformations remain open, the expanding and outstanding studies in this field continue to provide the insights and tools to attempt to answer them.

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Notes

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