Ru^III(edta) complexes as molecular redox catalysts in chemical and electrochemical reduction of dioxygen and hydrogen peroxide: inner-sphere versus outer-sphere mechanism†

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The reduction of molecular oxygen (O_2) and hydrogen peroxide (H_2O_2) by [Ru^III(edta)(pz)]^{2-} (edta^{4-} = ethylenediaminetetraacetate; pz = pyrazine) has been studied spectrophotometrically and kinetically in aqueous solution. Exposure of the aqua-analogue [Ru^II(edta)(H_2O)]^{2-}/C_0 to O_2 and H_2O_2 resulted in the formation of [Ru^III(edta)(H_2O)]^{2-}/C_0 species, with subsequent formation of the corresponding Ru^V complex. A working mechanism for the O_2 and H_2O_2 reduction reactions mediated by the Ru^III(edta) complexes is proposed. The role of the coordinated water molecule (by its absence or presence in the primary coordination sphere) in controlling the mechanistic pathways, outer-sphere or inner-sphere, is discussed.

Introduction

The electrochemical oxygen reduction reaction (ORR) proceeds by two-electron two-proton (2e^-/2H^+) partial reduction of O_2 to produce H_2O_2 (eqn (1) in Scheme 1) or direct four-electron four-proton (4e^-/4H^+) reduction of O_2 to 2H_2O (eqn (3) in Scheme 1). Although selective reduction of O_2 directly to H_2O is of continued interest in regard to its application in energy conversion, particularly in the field of fuel cells and metal-air batteries,^1-4 production of H_2O_2 via two-electron two-proton reduction of O_2 is also of considerable importance for environmental application like waste water treatment and chemical feedstocks.⁵,6

Use of transition metal complexes as molecular catalysts as a redox mediator to affect the reduction of oxygen is well documented in the literature.⁷-¹² Many schemes of catalytic processes that affect reduction of O_2 in combination of 2e^-/2H^+ and/or 4e^-/4H^+ pathways, either selectively or sequentially, have been reported. Noteworthy here, is that mechanistic details and kinetic parameters that control the efficiency of the four-electron reduction of O_2 to H_2O versus two-electron partial reduction of O_2 to H_2O_2 are still lacking, even though exhaustive efforts have been devoted for more than the last two-decades in mimicking and understanding the enzymatic activity of cytochrome c oxidase (which catalyses the direct four electron reduction of O_2 to H_2O during the final stage of respiration¹³). Another important aspect is, that although the reduction of hydrogen peroxide to water (eqn (2)) is seemingly easier thermodynamically than the reduction of dioxygen to hydrogen peroxide (eqn (1)), it is kinetically very difficult as it involves the cleavage of the O–O bond.

While mononuclear as well as binuclear complexes of copper, iron, cobalt and manganese, have been exhaustively studied pertaining to the oxygen reduction reaction,⁷-¹² the use of the ruthenium complex in this context is scanty in the literature.¹⁴ In the present work, we set out to examine the ability of the Ru^III(edta) complex (edta^{4-} = ethylenediaminetetraacetate) to affect the reduction reaction of molecular oxygen in aqueous medium. The feature that dominates the chemistry of the
[RuIII(edta)(H2O)]− complex is its lability towards aqua-substitution reactions,13 which affords an advantage of facile and straightforward binding of substrate molecules to the metal centre. In addition, a range of accessible and stable oxidation states made Ru(edta) complexes abidingly important to the catalytic studies for the past two decades. The significance of the Ru(edta) complexes in mimicking enzymatic redox reactions and small molecule activation, have been well established in very recent review articles.16,17

We for the first time explore, that [RuII(edta)(pz)]2− (pz = pyrazine) and its aqua-analogue [RuII(edta)(H2O)]2−, can efficiently mediate the sequential 2e−/2H+ reduction of O2 to H2O2 and further reduction of H2O2 to H2O. We report herein the results of the detailed spectral and kinetic investigation of the reduction of O2 and H2O2 by the above referred RuII(edta) complexes.

Results and discussion

The ‘edta’ ligand functions as a pentadentate ligand towards Ru(u) with a pendant acetate arm.19 The [RuIII(edta)H2O]− complex reacts with pyrazine (pz) to form the [RuIII(edta)(pz)]− complex, through a rapid and straightforward water displacement reaction (k1 = 2 × 104 M−1 s−1 and k−1 = 2 s−1 at 25 °C)19 as outlined in eqn (4).

\[
[RuII\text{(edta)}(\text{H}_2\text{O})]^- + \text{pz} \stackrel{k_1}{\rightleftharpoons} [RuII\text{(edta)}(\text{pz})]^- + \text{H}_2\text{O} \quad (4)
\]

The [RuII(edta)(pz)]2− complex can easily be reduced electrochemically (E1/2 value corresponding to the [RuII(edta)(pz)]− / [RuII(edta)(pz)]2− couple is 0.252 V vs. SHE)19 or chemically (in the presence of reducing agents viz. ascorbic acid)20 to its ruthenium(u) analogue, [RuII(edta)(pz)]2− (Fig. 1). The facile inter-conversion between low-spin RuII/RuIII redox states, does not require significant needs for structural changes that limit electron transfer steps.

The electronic absorption spectrum of the [RuII(edta)(pz)]2− complex in aqueous solution is featureless in the entire visible range, whereas its Ru(u)-analogue, [RuII(edta)(pz)]2− (Fig. 1) displays a strong band in the visible range (λmax = 462 nm, εmax = 11 600 M−1 cm−1) which was assigned to a metal to ligand charge transfer (MLCT) band.19 This huge spectral difference thus offers an amenable way to follow the kinetics of the electron transfer reactions spectrophotometrically.

Reduction of O2 by [RuII(edta)(pz)]2−

Addition of an oxygen-saturated aqueous solution to the deaerated red solution of the [RuII(edta)(pz)]2− complex (see S1 in ESI†) at pH 5.0 (acetate buffer), resulted in a gradual disappearance of the red colour.

The overall spectral changes recorded immediately after mixing the solutions of [RuII(edta)(pz)]2− and aqueous solution of dissolved oxygen, are shown in Fig. 2a. The spectral changes are attributed to the oxidation of [RuII(edta)(pz)]2− to [RuII(edta)(pz)]−, and a typical kinetic trace recorded at 462 nm (decay) is shown in Fig. 2b.

Effect of the concentration of the dissolved oxygen on the rate of the reaction was studied at 25 °C and pH 5.0 (representative kinetic traces recorded at 462 nm are shown in Fig. S1

Fig. 1 Pictorial presentation of [RuII(edta)(pz)]2−.
in ESI†). Under the specified conditions, the rate of the reaction estimated by the maximum slope, increases linearly with increasing concentration of dissolved oxygen (Fig. S2 in ESI†).

The spectral and kinetic observations can be accounted for in terms of the reaction sequence outlined in Scheme 2. The rate-determining step (5) proposed in the mechanism, involves a one-electron transfer from [RuII(edta)(pz)]2−/C0 to the O2 to yield O2− radical species in an outer-sphere manner. In the subsequent and kinetically inconsequential step (eqn (6)), the O2− radical rapidly reacts with another molecule of [RuII(edta)(pz)]2− to produce the peroxide ion (O22−), protonation of which (eqn (7)) results in the formation of hydrogen peroxide (H2O2).

The reduction of O2 with [RuII(edta)(pz)]2− under the specified conditions can be accounted for in terms of the rate-law expressed by eqn (8).

\[
\text{Rate} = k_1[Ru^{II}(edta)(pz)]^{2−}[O_2]
\]  

The value of the second-order rate constant (k1) estimated from the slope of the plot of rate versus [O2] shown in Fig. S2 (in ESI†) is 0.14 ± 0.01 M−1 s−1 at 25 °C. Addition of fresh ascorbic acid to the resultant solution obtained at the end of the aforementioned reaction (experimental conditions given under Fig. 2), resulted in the formation of the [RuII(edta)(pz)]2− almost quantitatively as evidenced by the spectral measurements (Fig. S3 in ESI†). The above observations clearly indicate the existence of a catalytic process in the overall reactions, wherein dioxygen (O2) is reduced to hydrogen peroxide (H2O2) via an electron transfer reaction, and the [RuII(edta)(pz)]− complex acts as a redox relay for electron transmission from ascorbic acid to O2.

**Reduction of H2O2 by [RuII(edta)(pz)]2−**

In order to understand the reaction of [RuII(edta)(pz)]2− with H2O2 formed during the reduction of O2 by [RuII(edta)(pz)]2− as shown in Scheme 2, we performed a detailed kinetic study of the reaction of [RuII(edta)(pz)]2− with H2O2 discretely under similar conditions of pH (5.0) and temperature (25 °C).

In Fig. 3a typical UV-visible spectral changes with time are shown (recorded by using stopped-flow rapid scan, diode array spectrophotometer) that occurred upon mixing an aqueous solution of [RuII(edta)(pz)]2− with the solution of H2O2 (in acetate buffer). The overall kinetic trace (Fig. 3b) derived from the time-resolved spectral changes, exhibited three clear steps (two decay and one growth at higher [H2O2] marked as I, II and III, respectively). The first decay step involves a small decrease in absorbance, not of enough significance (less than 5% as compared to the total absorbance change in the overall reaction time). Kinetic traces (see Fig. S4 in ESI†) pertinent to step I analysed on a shorter time scale as a function of [H2O2], are seemingly exponential in nature and could be fitted with a single exponential function within the precision of experimental data (R > 0.99). The values of the observed rate constant (kobs = 0.18 ± 0.02 s−1 at 25 °C and pH 5.0) so obtained were found to be independent of the H2O2 concentration. The spectral changes involved in step I may be attributed to the formation of a very weak [RuII(edta)(Ac)]3− (Ac = acetate) complex by the reaction of [RuII(edta)(pz)]2− with buffer component.

**Scheme 2** Proposed mechanism for the reduction of O2 by [RuII(edta)(pz)]2−.

$$\begin{align*}
[Ru^{II}(edta)(pz)]^{2+} + O_2 &\rightarrow k_1[Ru^{II}(edta)(pz)]^{2−} + O_2^{−} \quad (5) \\
[Ru^{II}(edta)(pz)]^{2−} + O_2 &\rightarrow \text{fast} [Ru^{III}(edta)(pz)]^{2+} + O_2^{2−} \quad (6) \\
O_2^{2−} + 2H^+ &\rightarrow H_2O_2 \quad (7)
\end{align*}$$
attainment of a limiting rate at higher [H2O2] (Fig. 4), the rate of the disappearance of [RuII(edta)(pz)]2− (Scheme 3) by H2O2 remaining present in excess (see S1 in ESI†) to form [RuIV(edta)O]− (eqn (11)). The following rate-law (eqn (13) and (14)) can be derived from the reactions in Scheme 3 on the basis that the rate-determining step (eqn (10)) involves the disappearance of the [RuII(edta)(pz)]2− complex monitored at 462 nm.

\[
\begin{align*}
-d[[Ru^{II}(edta)(pz)]^{2-}/dt & = k_1k_3[[Ru][H_2O_2]/(1 + K_1[H_2O_2])] \\
-\frac{d[[Ru^{II}(edta)(pz)]^{2-}}{dt} & = 1/k_2[[Ru] + 1/k_3[[Ru][H_2O_2] \\
\end{align*}
\]

A plot of \(-d[[Ru^{II}(edta)(pz)]^{2-}}/dt versus \(1/[H_2O_2]\) was found to be linear (Fig. S5 in ESI†). Considering \([Ru]_0\) (total concentration of ruthenium) is 0.25 mM, the values of \(k_2\) and \(K_1\) calculated from the intercept and slope of the plot (Fig. S5 in ESI†) are 8.24 \times 10^{-3} \text{ s}^{-1} and 60 \text{ M}^{-1}, respectively, at 25 °C and pH 5.0.

As seen in Fig. 3a, after complete disappearance of the peak at 462 nm (attributed to the oxidation of the [RuII(edta)(pz)]2− to [RuIII(edta)(pz)]− under the specified conditions (see Fig. 3a), the reaction is followed by a step that involves the formation of a band at 390 nm. This new band is characteristic of the [RuV(edta)O]− complex \((\lambda_{max} = 390 \text{ nm}; \epsilon_{max} = 8 \times 10^3 \text{ M}^{-1} \text{ cm}^{-1})\). The observed increase in the absorbance at 390 nm with time at higher H2O2 concentration (step III in Fig. 3b), recorded after 95 s at 25 °C and pH 5.0 (acetate buffer). [Ru] = 0.25 mM [H2O2] = 40.

Reaction of H2O2 with [RuIII(edta)(pz)]−

Above observations, necessitated us to perform further kinetic investigations to understand the mechanistic details pertaining to the formation of [RuIV(edta)O]− in the reaction of [RuIII(edta)(pz)]− with H2O2. Noteworthy here, is that the formation of the [RuIV(edta)O]− complex through an oxo-transfer reaction from the precursor oxidant ROOH (ROOH = H2O2, BuOOH and KHSO5) to [RuIII(edta)[H2O2]] was reported by us.25–27 An inner-sphere mechanism involving the formation of the [(edta)RuIV(OOR)]− (intermediates \(R = H, \text{Bu and SO}_3^-\)) in a rapid pre-equilibrium step, followed by the rate-controlling heterolytic cleavage of the O−O bond to produce the [RuV(edta)O]− complex, was proposed.25–27

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In Fig. 6 the UV-vis spectral changes with time that occurred upon mixing aqueous solutions of [RuIII(edta)(pz)]\(^{2-}\) and H\(_2\)O\(_2\), are shown. The observed spectral changes (Fig. 6) are attributed to the oxidation of [RuIII(edta)(pz)]\(^{2-}\) to [RuV(edta)O]\(^{2-}\) under the specified conditions. The kinetic traces at 390 nm, generated from the recorded spectra, are presented in Fig. S6 (see ESI†).

The effect of the H\(_2\)O\(_2\) concentration on the values of the observed rate constant (\(k_{\text{obs}}\)) is shown in Fig. 7a. The observed saturation of the \(k_{\text{obs}}\) values at higher H\(_2\)O\(_2\) concentration, suggests a process that involves a rate-limiting pre-equilibrium step followed by the rate-determining formation of [Ru\(^{V}\)(edta)O]\(^{-}\) species. On the basis of a pre-equilibrium approach, the above kinetic results for the formation of [Ru\(^{V}\)(edta)O]\(^{-}\) in the reaction of [RuIII(edta)(pz)]\(^{2-}\) with H\(_2\)O\(_2\), can be accounted for in terms of the following mechanism proposed in Scheme 4.

Considering H\(_2\)O\(_2\) being a poor nucleophile that cannot efficiently replace pyrazine (a strong aromatic N-heterocyclic \(\pi\)-acidic ligand) to form [Ru\(^{III}\)(edta)(OOH)]\(^{2-}\) species (as observed in case of the reaction of [Ru\(^{III}\)(edta)(H\(_2\)O)]\(^{-}\) with H\(_2\)O\(_2\))\(^{27}\), we invoke the formation of a [Ru\(^{III}\)(edta)(H\(_2\)O\(_2\))(pz)]\(^{-}\) intermediate \(\nu\) via coordination of H\(_2\)O\(_2\) to the metal centre by dislodging the coordinated acetate arm of the ‘edta’ ligand in the pre-equilibrium step (eqn (15)). The dangling acetate arm plausibly stabilizes the proposed intermediate species by forming a transient hydrogen bond between carboxylate oxygen atom of the dangling acetate arm and H-atom of the H\(_2\)O\(_2\) coordinated to the metal centre as shown pictorially in Scheme 4 (eqn (15)). However, no spectral evidence in favour of the formation of the proposed intermediate was observed. Noteworthy here is that a distinct spectral evidence (formation of a shoulder in the 390 nm band at 425 nm) confirming the formation of [Ru\(^{III}\)(edta)(OOH)]\(^{2-}\) intermediate species was observed during the course of the reaction of [Ru\(^{III}\)(edta)(H\(_2\)O)]\(^{-}\) with H\(_2\)O\(_2\))\(^{27}\). In the subsequent rate-determining step, the [Ru\(^{III}\)(edta)(H\(_2\)O\(_2\))(pz)]\(^{-}\) intermediate undergoes heterolytic cleavage of the O-O bond to produce [Ru\(^{V}\)(edta)(pz)O]\(^{-}\) species, with concomitant release of a water molecule (eqn (16)) in a concerted pathway. However, the metal–pyrazine bond at a higher oxidation state of the metal in [Ru\(^{V}\)(edta)(pz)O]\(^{-}\) becomes less stable, and the \(\pi\)-acidic ligand pyrazine, thereby dissociates allowing the dangling carboxylate group to bind to the metal centre again to produce the [Ru\(^{V}\)(edta)O]\(^{-}\) product complex in a kinetically inconsequential step (eqn (17)). The following rate-law (eqn (18) and (19)) can be derived for the reactions in Scheme 4 on the basis of the rate-determining formation of the [Ru\(^{V}\)(edta)O]\(^{-}\) complex.

\[
k_{\text{obs}} = k_3K_2[H_2O_2]/(1 + K_2[H_2O_2]) \quad (18)
\]

\[
1/k_{\text{obs}} = 1/k_3 + 1/k_1K_3[H_2O_2] \quad (19)
\]

The plot of \(1/k_{\text{obs}}\) versus \(1/[H_2O_2]\) is linear (Fig. 7b), and the values of \(k_3\) and \(K_3\) from the intercept and slope are 0.23 ± 0.03 s\(^{-1}\) and 82 ± 2 M\(^{-1}\), respectively, at 25 °C and pH 5. The product...
The above findings taken together with that reported for the reaction of [RuIII(edta)(H2O)]− with H2O2 are suggestive of the fact that the reduction of H2O2 by both [RuIII(edta)(pz)]− and [RuIII(edta)(H2O)]−, resulting in the formation of [RuIII(edta)[O]]2− and H2O as ultimate reaction products, proceeds through an inner-sphere pathway demonstrating similar kinetic features. Although the common suggested mechanism involves heterolytic cleavage of the O–O bond, the ability of H2O2 to bind to the Ru(III)-centre through a ligand substitution process, governs the efficiency of the H2O2 reduction process. In this regard, [RuIII(edta)(H2O)]− due to its unusual lability towards a substitution reaction, has an advantage over [RuIII(edta)(pz)]−.

### Reaction of O2 and H2O2 with [RuIII(edta)(H2O)]−

Noteworthy here, is that the [RuIII(edta)(H2O)]− complex also exhibited a metal based one-electron transfer reaction, and the E1/2 value reported for the [RuIII(edta)[H2O]]2−/RuIII[edta][H2O]− is −0.018 V (vs. SHE),19 which is much more negative than that reported for the [RuIII(edta)(pz)]−/RuIII[edta](pz)2− couple (E1/2 = 0.252 V vs. SHE).19

Though [RuIV(edta)(H2O)]2− is a stronger reductant thermodynamically than [RuIV(edta)(pz)]2− towards O2 reduction, detailed kinetic investigations were practically not feasible because of very insignificant spectral differences between analogous of Ru(m) and Ru(n)-edta complexes. The [RuIII(edta)(H2O)]− in water is almost featureless over the entire visible range of the spectrum, but exhibits a strong absorption band at 280 nm (εmax = 2800 ± 50 M−1 cm−1) and a shoulder at 350 nm (εmax = 680 ± 60 M−1 cm−1) in the UV region.19 The Ru(n) analogue, [RuIV(edta)(H2O)]2− displays almost similar spectral features exhibiting an intense band at 282 nm (εmax = 2900 ± 100 M−1 cm−1) and a weak shoulder at 427 nm (εmax = 260 ± 15 M−1 cm−1).19 Nevertheless, the formation of the [RuIV(edta)[O]]2− complex was evidenced in the spectral changes (Fig. 8) that occurred in the reaction of [RuIV(edta)(H2O)]2− with oxygen. Time resolved spectral changes recorded (Fig. 8) after mixing of the aqueous solution of [RuIV(edta)(H2O)]2− (0.025 mM) with an oxygen saturated aqueous solution, clearly revealed the gradual build-up of the peak at 390 nm (inset of Fig. 8) characteristic for the [RuIV(edta)[O]]2− complex.24 The above spectral observations may be explicable in terms of the following reactions as outlined in Scheme 5.

In the above proposed mechanism (admittedly speculative), the reduction of O2 to H2O2 with concomitant formation of the [RuIII(edta)(H2O)]− takes place in an outer-sphere pathway (eqn (20)). The appearance of the peak at 390 nm in Fig. 8 can be accounted for by the reaction of H2O2 with the one-electron oxidized product [RuIII(edta)(H2O)]− (eqn (21)).

Formation of [RuIV(edta)[O]]2− was also noticed in the direct reaction of [RuIV(edta)(H2O)]2− with H2O2. The absorbance versus time profile (recorded at 390 nm), pertaining to the formation of [RuIV(edta)[O]]2− in the reaction of [RuIV(edta)(H2O)]2− with H2O2, is given in Fig. 9. As it is seen, the kinetic trace exhibits a clear initial induction period. Above observations may be explicable in terms of the following reaction scheme (Scheme 6) proposed for the reduction of H2O2 by [RuIV(edta)(H2O)]2−.

The observed initial induction period (Fig. 9) is explicable in terms of a catalytic cycle wherein the [RuIII(edta)(H2O)]− complex formed concomitantly during reduction of H2O2 by [RuIV(edta)(H2O)]2− (eqn (22)) rapidly undergoes reduction by the ascorbic acid present in excess in the reacting system (see S1 in ESI†), regenerating the [RuIV(edta)(H2O)]2− species in the reaction mixture. Such reaction cycles sustained until the ascorbic acid present in the reacting system are completely

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**Scheme 6** Proposed mechanism for the reaction of [RuIV(edta)(H2O)]2− with H2O2.
analogues) are found effective in reducing H₂O₂ to H₂O, however, through an inner-sphere electron transfer pathway as depicted in Scheme 7. The key-feature is that both the [RuII(edta)(pz)]²⁻ and [RuII(edta)(H₂O)]²⁻ complexes are found effective in reducing H₂O₂ to H₂O, however, through an inner-sphere electron transfer pathway (Scheme 7).

[RuIII(edta)(pz)]⁻ mediated electrochemical reduction of O₂

Being inspired by the aforementioned results, we performed a brief spectro-electrochemical experimentation (see S2 in ESI†), the results of which evidently reveal that [RuIII(edta)(pz)]⁻ could act as a ‘molecular redox catalyst’ for electrochemical reduction of O₂. Reduction of [RuIII(edta)(pz)]⁻ to [RuII(edta)(pz)]²⁻ was achieved electrochemically by carrying out constant potential electrolysis (at −0.05 V vs. Ag/AgCl) of the solution of [RuIII(edta)(pz)]⁻ (in acetate buffer solution at pH 5.0). Spectrum of the solution of [RuIII(edta)(pz)]⁻ (0.5 mM in acetate buffer) is shown in Fig. 11a. Constant potential electrolysis of the solution of [RuIII(edta)(pz)]⁻ leading to the formation of the [RuII(edta)(pz)]²⁻ species was evident spectrophotometrically by the appearance of its characteristic peak at 462 nm (Fig. 11b). The typical current versus time plot pertaining to the above mentioned electrochemical process is shown in the inset of the Fig. 11. After withdrawal of the potential, O₂ was bubbled through the electrolysed solution for 300 s. The spectral changes thereafter (Fig. 11c) show appreciable collapse of the peak at 462 nm, which is consistent with the re-oxidation of [RuII(edta)(pz)]²⁻ to [RuIII(edta)(pz)]⁻ by dioxygen (O₂). Electrolysis of the solution (at −0.05 V) for 500 s again regenerates the [RuIII(edta)(pz)]⁻ species as evident spectrophotometrically (Fig. 11d). However, the intensity of the band at 462 nm as noticed in the spectrum (Fig. 11d) is significantly smaller in comparison to that observed in the spectrum (Fig. 11b). This may plausibly be associated with the fact that the unconsumed dissolved oxygen present in the reacting system prior to the second run of electrolysis, may compete with the electrochemical reduction process for which complete formation of the [RuII(edta)(pz)]²⁻ could not take place within the time period the voltage was on (500 s). Observation of the band (at 462 nm), with a higher absorbance in the spectrum of the electrolysed solution, which was deoxygenated through argon purging prior to the electrolysis (see Fig. S7 in ESI†), supports our above argument.
Based on the above observations, taken together with that reported for the reaction of [RuIII(edta)(pz)]O2− with molecular oxygen (O2) in the preceding section, the role of [RuIII(edta)(pz)]− as an electron transfer redox catalyst in the electrochemical reduction of O2 may be outlined in Scheme 8.

Conclusions

In conclusion, the results of the present study reveal that in the overall reactions, whether chemical or electrochemical, dioxygen (O2) is reduced to hydrogen peroxide (H2O2) via electron transfer reaction, and the [RuIII(edta)(pz)]O2− complex acts as a redox relay for electron transmission. In case of the chemical process, it takes electrons from ascorbic acid to reduce O2 to H2O2, whereas in case of the electrochemical process, it uses the electrons from the working electrode to effect electrochemical reduction of O2 to H2O2 in aqueous acidic solution. The results of our studies further ascertain that both the [RuII(edta)(pz)]2+ and [RuIII(edta)[H2O]2]3− complexes in presence of electron donors can reduce O2 to H2O2 and H2O2 to H2O efficiently in a sequential manner. The redox mediating properties of the aforesaid Ru(edta) complexes along with their wide range of chemically accessible oxidation states (II to V), and their durability in the redox processes, are indeed intriguing and prospective. Our results may shed light towards mechanistic understanding of the homogeneously catalysed reduction of O2 and H2O2 and provide pointers for future research pertaining to the application of such metal complexes in the field of fuel cells and metal–air batteries.

Conflicts of interest

There are no conflicts to declare.

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