Dalton Transactions



PAPER

View Article Online
View Journal | View Issue



Cite this: *Dalton Trans.*, 2022, **51**, 1206

Influence of the spatial distribution of copper sites on the selectivity of the oxygen reduction reaction[†]

N. W. G. Smits, D. Rademaker, A. I. Konovalov, Da M. A. Siegler and D. G. H. Hetterscheid b *a

Moving towards a hydrogen economy raises the demand for affordable and efficient catalysts for the oxygen reduction reaction. **Cu-bmpa** (bmpa = bis(2-picolyl)amine) is shown to have moderate activity, but poor selectivity for the 4-electron reduction of oxygen to water. To enhance the selectivity towards water formation, the cooperative effect of three **Cu-bmpa** binding sites in a single trinuclear complex is investigated. The catalytic currents in the presence of the trinuclear sites are lower, possibly due to the more rigid structure and therefore higher reorganization energies and/or slower diffusion rates of the catalytic species. Although the oxygen reduction activity of the trinuclear complexes is lower than that of mononuclear **Cu-bmpa**, the selectivity of the copper mediated oxygen reduction was significantly enhanced towards the 4-electron process due to a cooperative effect between three copper centers that have been positioned in close proximity. These results indicate that the cooperativity between metal ions within biomimetic sites can greatly enhance the ORR selectivity.

Received 28th September 2021, Accepted 25th November 2021 DOI: 10.1039/d1dt03296h

rsc li/dalton

Introduction

The development and storage of renewable energy are crucial to limit our fossil fuel consumption while sustaining the demand for energy. Within such a hydrogen society, fuel cell technology plays a central role. 1-3 The limiting factor of such systems lies with the oxygen reduction reaction (ORR), which involves the redistribution of four protons and four electrons with a simultaneous cleavage of the O-O bond. This leads to a complex reaction mechanism with numerous intermediates.⁵ Consequently, a significant overpotential is required for ORR catalysis, which results in a substantial loss of energy. For the ORR, platinum catalysts are typically employed due to their relatively low overpotential, which is still quite substantial with roughly 400 mV.6,7 Moreover, platinum is not a sufficiently abundant material for large scale applications. This raises the demand for catalysts based on more affordable materials to drive the ORR at a low overpotential.

Inspiration for the design of efficient catalysts that catalyse the ORR at a low overpotential and are based on abundant materials can be found in natural systems, particularly in redox metalloenzymes. A prime example is the multicopper enzyme laccase, which belongs to a family of oxidases and can be found in a variety of natural sources. This enzyme couples the oxidation of an organic substrate near a mononuclear Cu site to ORR catalysis at a trinuclear Cu cluster. Electrochemical studies on immobilized laccase have shown that the enzyme catalyzes the ORR close to the equilibrium potential. However, laccase has a low overall efficiency for the ORR in fuel cells due to the instability of the enzyme under fuel cell conditions and slow electron transfer to the active site. Nevertheless, the active site of laccase represents an interesting starting point for the development of new Cubased molecular catalysts for the ORR that operate with a low overpotential and a high efficiency.

Mononuclear Cu complexes have been explored to catalyze the ORR. ^{16–34} Additionally, several dinuclear complexes have been investigated to induce a cooperative effect during ORR catalysis. ^{35–38} We have recently shown that the mononuclear complex [Cu(tmpa)(solv)]⁺ (Cu-tmpa, tmpa = tris(2-picolyl) amine) shows exceptionally high ORR catalytic performance with a turn-over frequency (TOF) of almost 2 million per second. ^{19,39} We showed that the reduction of oxygen to water proceeds *via* a two-step process in which hydrogen peroxide is formed as an obligatory intermediate product. ⁴⁰ The complexes Cu-terpy and Cu-bmpa, which showed a lower denticity and flexibility of the ligand framework than Cu-tmpa, undergo the ORR with a lower activity and with a lower selectivity

^aLeiden Institute of Chemistry, Leiden University, P.O. box 9502, 2300 RA Leiden, The Netherlands. E-mail: d.g.h.hetterscheid@chem.leidenuniv.nl

^bDepartment of Chemistry, Johns Hopkins University, 3400 N. Charles Street, Baltimore, MD 21218, USA

[†]Electronic supplementary information (ESI) available. CCDC 2099322 for Cu₃L1. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/d1dt03296h

Dalton Transactions Paper

towards water (terpy = 2,2':6',2"-terpyridine; bmpa = bis(2-pyridylmethyl)amine).20 Since H2O2 is damaging to fuel cell systems, the production of this compound is an unwanted side-reaction.

In laccase, the cooperativity of the Cu ions in the trinuclear cluster results in the reduction of oxygen to water, without the formation of H₂O₂ as an intermediate product. Inspired by laccase, several trinuclear Cu-based molecular catalysts have been reported for the ORR, of which most are based on ligands bearing alkylamine and pyridylalkyl-amine functional groups. 41-52 In early reports, only oxygen binding and reductive cleavage were investigated. 41,42,44 Later, the ORR activity of several trinuclear Cu complexes was investigated. 46-48 These studies were carried out either by using organic solvents in the presence of sacrificial reagents, 47 or by dropcasting the catalyst as part of carbon paste onto electrodes. Due to the very flexible and dynamic linkers employed to tether the copper sites together, it remains difficult to assess whether under these operative conditions these structures truly function as trinuclear sites.44,47 Consequently, the reported results have been rather inconclusive thus far. Inspired by the active site of laccase, and lessons learned in the previous studies, our study here focuses on a structurally rigid triethylbenzene node that forces all three copper sites linked to the remaining aromatic positions in close proximity to each other (Fig. 1).

Copper complexes with the L1 and L2 ligands have previously been reported for their reactivity with oxygen in organic solution and for their ability to cleave DNA via hydrolysis. 53-56 In these studies, the crystal structures of these trinuclear complexes indicated that in Cu₃L2 all three copper sites are forced into close proximity due to steric repulsion between neighboring groups on the aromatic node, while in Cu₃L1 only two Cu centers will lie in close proximity. ^{55,56} We report here that Cu₃L1 has a similar selectivity for H₂O compared to the parent mononuclear complex Cu-bmpa, and that the close proximity of the three Cu ions in Cu₃L2 induces high selectivity for the selective formation of H₂O.

Results

Characterization of the trinuclear compounds

Synthesis. The L1 and L2 ligands were synthesized via adopting the reported procedures (ESI 2†).55 The consecutive complexation of the ligands with three equivalents of Cu(OTf)₂

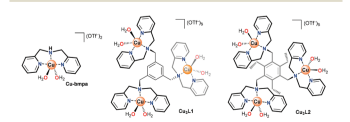


Fig. 1 Structures of Cu-bmpa and trinuclear Cu complexes Cu₃L1 and Cu₃L2.

resulted in the formation of the trinuclear copper complexes Cu₃L1 and Cu₃L2 (Fig. S1†). Both Cu₃L1 and Cu₃L2 were characterized by UV-Vis spectroscopy and superconducting quantum interference device (SQUID) magnetometry. The purity of the samples was confirmed by elemental analysis, while UV-Vis stability studies also indicated that both complexes are stable in an aqueous pH 7 phosphate buffer for at least two days (Fig. S2†).

Single crystal X-ray crystallography

Slow vapour diffusion of Et₂O into a concentrated solution of Cu₃L1 in acetone at 279 K resulted in single crystals which were suitable for X-ray crystallography (Fig. 2 and ESI 3†).

The crystal structure clearly shows an asymmetric distribution of the three Cu-bmpa sites relative to the benzene plane. A similar distribution was published for the crystal structure of [L1(CuCl₂)₃] by Guo et al. in 2006,⁵⁶ who reported square pyramidal geometries for all three Cu^{II} ions. In contrast, two of the three CuII ions in the crystal structure of Cu₃L1 have an octahedral geometry due to the close proximity of triflate counter ions (Fig. 2b). The relatively short Cu1-O19 and Cu3-O13 bond distances of 2.691(3) and 2.650(3) Å, respectively, suggest that the triflate counter ions are weakly coordinated to the two Cu centers.

For Cu₃L2, the various single crystals that were obtained during this study did not diffract well enough for X-ray structure determination. However, the crystal structure of [L2 (CuCl₂)₃] has been reported by Anslyn et al. and showed closer proximity of the three Cu-bmpa sites to each other. 55 All three sites are forced to the same side of the benzene plane due to the steric effect of the three ethyl substituents (Fig. 3).

Magnetic properties

To assess the strength of the spatial interaction of the three paramagnetic Cu^{II} centers, the magnetic properties of complexes Cu₃L1 and Cu₃L2 were investigated using a superconducting quantum interference device (SQUID). In Fig. 4, the inverse of the obtained paramagnetic susceptibility (X_p) is plotted versus temperature. To extract the exchange coupling constants (J) between each pair of Cu^{II} ions, the obtained magnetic data were fitted using the PHI software (ESI 4†).57

For Cu₃L1, this resulted in three J-values, two of which were negligibly small suggesting virtually no magnetic coupling between two pairs of CuII ions. The third constant amounted to +23 cm⁻¹ indicating ferromagnetic coupling (Fig. 5).⁵⁸ In order to be able to couple ferromagnetically, the Cu ions have to be in close proximity to each other which is consistent with the two Cu ions being on one side of the benzene plane and the third one to the other side of the ring as was observed in the acquired crystal structure. For Cu₃L2, the fitting returns one J value of +49 cm⁻¹ confirming the symmetric distribution of the three Cu^{II} sites as expected on the basis of the reported crystal structure for [L2(CuCl₂)₃] (Fig. 5).

Paper Dalton Transactions

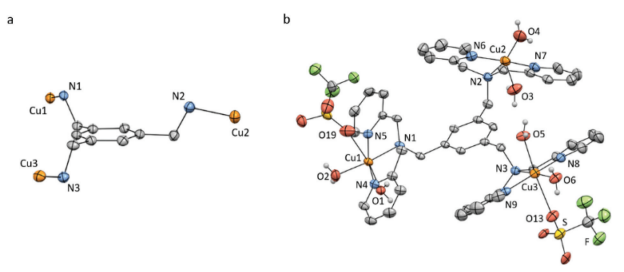


Fig. 2 Displacement ellipsoid plots (50% probability level) of Cu_3L1 at 110(2) K. (a) Orientation of the Cu ions relative to the benzene plane. (b) Full structure of Cu_3L1 . Lattice solvent molecules, four non-coordinating triflate ions, and all hydrogen atoms which are not part of the aqua ligands are omitted for clarity. Selected bond distances and angles are reported in ESI section 3.†

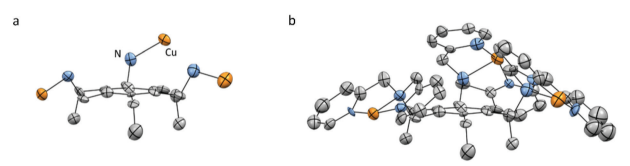


Fig. 3 Reported crystal structure of the cationic part of $[L2(CuCl_2)_3]$ as determined by X-ray crystallography with displacement ellipsoids scaled at the 30% probability level. (a) Orientation of the Cu ions relative to the benzene plane. (b) All hydrogen atoms, lattice solvent molecules, and the chloride ions are omitted for clarity. Adapted with permission from Anslyn *et al.* Copyright (2002) American Chemical Society.⁵⁵

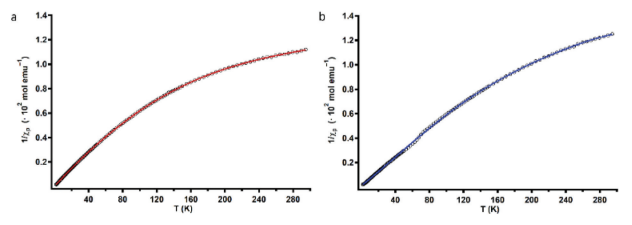


Fig. 4 Variable-temperature magnetic susceptibility plots of Cu_3L1 (a) and Cu_3L2 (b). Black circles depict the experimentally obtained data points, and red and blue lines correspond to the fitted data that were used to obtain magnetic exchange coupling constants (J).

Dalton Transactions Paper

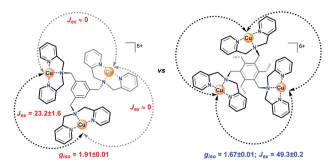


Fig. 5 Spatial distribution of the Cu-bmpa sites relative to the benzene plane for Cu₃L1 (left) and Cu₃L2 (right) as confirmed by the exchange coefficients obtained by fitting of the obtained magnetic data. Black arrows indicate ferromagnetic coupling and grey arrows indicate no magnetic coupling between the two corresponding copper ions.

Structure in solution

Although the X-ray and SQUID analysis confirm the structure in the solid phase, conversion to other conformers may still occur in solution. The structure of hexaethylbenzene and variations thereof have been extensively studied in solution in the past. Both computational and NMR studies showed that for hexaethylbenzene the up-down-up-down-up-down conformation of six substituents is the lowest conformer, with at least 3.46 kcal mol⁻¹ energy difference to the next favorable geometry (i.e. up-down-down-up-down-down). 59,60 Therefore, a large majority of the compounds adopts the alternating updown conformation. We anticipate that the even more bulkier Cu-bmpa substituent will not lower this energy difference between the various conformers, and therefore, it is expected that the alternating up-down conformation is also predominantly adopted by Cu₃L2 in solution. ⁵⁹⁻⁶³ For Cu₃L1 the rotation of the Cu-bmpa substituents around the benzene node will be less prohibited due to the absence of the ethyl groups. Therefore, the distribution between the conformers is expected to be much more random for Cu₃L1.

Electrochemical behaviour of Cu₃L1 and Cu₃L2

Redox couple under an argon atmosphere. The redox behavior of Cu₃L1 and Cu₃L2 was investigated by performing cyclic voltammetry (CV) measurements (Fig. 6). The acquired voltammograms for Cu₃L1 and Cu₃L2 show quite broad cathodic and anodic peaks which are located at a half-wave potential $(E_{1/2})$ of 0.37 and 0.50 V vs. the RHE, respectively. The peak-to-peak potential separation (ΔE_p) amounts to 105 mV for Cu₃L1 and to 90 mV for Cu_3L2 . These relatively large ΔE_p values can be the result of the slow electron transfer and/or partial overlap of multiple electrochemical processes which have a lower redox potential than the preceding electrochemical step. 64 The presence of an oxidative shoulder at more positive potential than the main oxidative process of Cu₃L1 supports the latter hypothesis.

Differential pulse voltammetry (DPV), and linear sweep voltammetry (LSV) were used to further pinpoint the redox behavior of Cu₃L1 and Cu₃L2. In the latter experiment, a resting

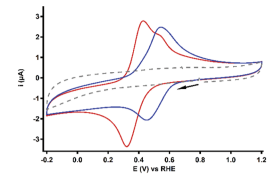


Fig. 6 CV profiles of 0.1 mM Cu₃L1 (red) and Cu₃L2 (blue). For both complexes, only the first scan of the measurement is depicted. The reference voltammogram in the absence of the complex is depicted as a grey dashed line. Conditions: 0.1 M pH 7 PB, 1 atm Ar, r.t., GC WE, 100 mV s^{-1} scan rate.

potential is applied either at a high or low potential before the start of the LSV measurement to ensure that all copper sites are either in the +II or +I oxidation state at the start of the LSV experiment despite the slow electron transfer kinetics.

For Cu₃L1, both anodic LSV and DPV measurements indicated the presence of two anodic peaks at 0.38 and 0.48 V vs. the RHE (Fig. S4†). The separation of the broad anodic peak into multiple oxidative processes has previously been observed for [L1(CuCl₂)₃] by Zhao et al., ⁵⁶ who identified three individual anodic peaks in 0.1 M aqueous KCl. The observation of three separate anodic processes instead of two for Cu₃L1 might be an effect of the presence of a different electrolyte and other counter ions. The cathodic peak could not be resolved in separate reduction processes for Cu₃L1.

For Cu₃L2, a cathodic sweep resulted in a separation of the main cathodic peak into two distinct reductive processes at 0.46 and 0.17 V vs. the RHE (Fig. S5†). Assuming the influence of the magnetic coupling of the Cu^{II} ions on this separation, the initial reduction of one or two Cu^{II} ions could result in a thermodynamically less favorable reduction of the other Cu^{II} ion(s). Not only this electronic coupling, but also the structural changes upon reduction can cause a separation of the cathodic peak. 65-68 Separation of both the cathodic and anodic peak has been reported for $[L2(CuX)_3]$ (X = Br or I) in DCM by Kim et al. 54 In contrast, the DPV of Cu₃L2 did not result in the separation of the main anodic peak into distinct processes.

Oxygen reduction reaction catalysis

The ORR behavior of Cu₃L1 and Cu₃L2 was investigated with CV under 1 atm O2. Under these conditions, the voltammograms of both complexes show a peak-shaped catalytic wave (Fig. 7). For Cu-bmpa, an $E_{\text{cat/2}}$ value of 0.37 V vs. the RHE has been reported with the notion that $E_{\text{cat/2}} > E_{1/2}$ due to substrate depletion near the electrode. ²⁰ However, a relatively low $E_{\text{cat/2}}$ value is observed for both Cu₃L1 and Cu₃L2, amounting to 0.37 V vs. the RHE for Cu_3L1 and 0.33 V vs. the RHE for Cu_3L2 . This means that $E_{\text{cat}/2}$ is equal to the $E_{1/2}$ value for $\text{Cu}_3\text{L1}$ and lower than the $E_{1/2}$ value for Cu_3L2 . This suggests that the Paper **Dalton Transactions**

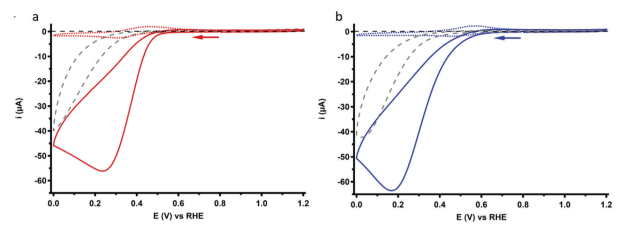


Fig. 7 CV profiles of 0.1 mM Cu₃L1 (a) and Cu₃L2 (b) under 1 atm O₂ (solid lines) or 1 atm Ar (dotted lines). For both complexes, only the first scan of each measurement is depicted. The reference voltammogram in the absence of the complex under 1 atm O2 is depicted as a grey dashed line. Conditions: 0.1 M pH 7 phosphate buffer, 293 K, GC WE, 100 mV s⁻¹ scan rate.

rates for the oxygen reduction reaction are not much influenced by substrate depletion compared to Cu-bmpa.

Active species homogeneity

The homogeneity of the redox behavior of both Cu₃L1 and Cu₃L2 was assessed by performing a scan rate dependence study. From the $i_{p,red}$ vs. $v^{1/2}$ plots depicted in Fig. S6,† a linear relationship is observed between the cathodic peak current and the square root of the scan rate for both Cu₂L1 and Cu₂L2, which is in good agreement with a diffusive species (ESI 6.2†).

A deposition test with CV under 1 atm O2 indicated that after 1 scan, an ORR active deposit was formed (ESI 7.1†). However, the activity of this deposition was significantly lower than that for the catalyst solution. Moreover, electrochemical quartz crystal microbalance (EQCM) experiments were performed to quantify the amount of complex that was deposited during a CV experiment (ESI 7.2†) which showed that only 7.0 and 8.8 pmol cm⁻² of Cu₃L1 and Cu₃L2 are deposited during one scan, respectively. Therefore, the effect of the deposit on the ORR catalysis during CV experiments was considered to be negligible.

Product selectivity determination

The product selectivity of the ORR catalyzed by Cu₃L1 and Cu₃L2 was investigated using a setup with a rotating ring disk electrode (RRDE).20 This was done by performing LSV at the GC disk and chronoamperometry (CA) at the Pt ring while rotating the RRDE at a speed of 1600 RPM. For both Cu₃L1 and Cu₃L2, LSV was performed between 1.0 and -0.15 V vs. the RHE, while CA at the ring was performed at 1.2 V vs. the RHE to be able to oxidize any H₂O₂ that is produced during ORR catalysis (Fig. 8). Since H_2O_2 is the product of the twoelectron ORR, the presence of the Pt ring enables the determination of the product selectivity of the catalyzed ORR (ESI 8.2†).

For the RRDE LSV measurements the ORR onset potential of Cu₃L1 and Cu₃L2 has been defined as the potential at

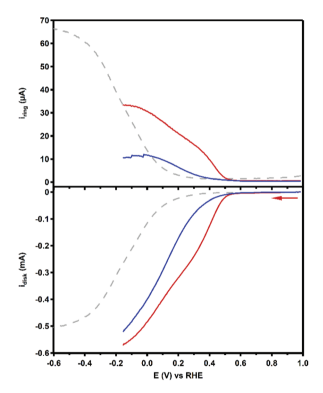


Fig. 8 RRDE LSV curves of 0.1 mM Cu₃L1 (red) and Cu₃L2 (blue) under 1 atm O₂ at 1600 RPM. The reference voltammogram in the absence of the complex is depicted as a grey dashed line. Conditions: 0.1 M pH 7 phosphate buffer, 293 K, GC disk, Pt ring at 1.2 V vs. RHE, 50 mV s⁻¹ scan rate.

which $i_c/i_{\rm GC}$ > 3, in which i_c is the disk current observed during ORR catalysis performed by the catalyst and i_{GC} is the disk current observed in the absence of the catalyst. 19,20 For Cu₃L1 and Cu₃L2, this onset potential is located at 0.55 and 0.52 V vs. the RHE, respectively. Considering the reported ORR onset potential of 0.49 V vs. the RHE for Cu-bmpa, the overpotential for the ORR catalyzed by the trinuclear catalysts is

Dalton Transactions Paper

slightly lower than the overpotential of Cu-bmpa.²⁰ For Cu₃L1, this slight decrease of 0.06 V for the overpotential reflects the slight positive shift in the $E_{1/2}$ value of 0.07 V vs. the RHE compared to Cu-bmpa. However, this is not the case for Cu₃L2; the slight decrease of 0.03 V for the overpotential does not reflect the large positive shift in the $E_{1/2}$ value of 0.20 V vs. the RHE compared to Cu-bmpa.

The RRDE LSV data of Cu₃L1 and Cu₃L2 show maximum catalytic disk current (i_{cat}) values of 0.57 and 0.52 mA at -0.15 V vs. the RHE, respectively (Fig. 8). These values are similar to the reported i_{cat} value of 0.57 mA for Cu-bmpa at -0.15 V vs. the RHE.20 Just like for Cu-bmpa, an increase in the ring current is observed with decreasing applied disk potential for both Cu₃L1 and Cu₃L2. This indicates that both catalysts produce H₂O₂ along the entire potential window in which the ORR takes place.

To quantify the formation of H₂O₂ along the potential regime for ORR catalysis, the percentage of H2O2 produced during ORR catalysis (%H2O2) was determined according to the following equation:

$$\% H_2 O_2 = \frac{2 \times (i_{\text{ring}} / N_{\text{H}_2 O_2})}{i_{\text{disk}} + (i_{\text{ring}} / N_{\text{H}_2 O_2})} \times 100\%$$

where $i_{\rm ring}$ and $i_{\rm disk}$ are the observed ring and disk current, respectively, and $N_{\rm H_2O_2}$ is the collection efficiency of the Pt ring for H₂O₂ (see the ESI 8.1† for a full derivation).⁶⁹ This collection efficiency amounts to 0.125 as we have reported previously for the same RRDE setup.¹⁹

Fig. 9 shows the %H₂O₂ values along the potential regime for ORR catalysis by Cu₃L1 and Cu₃L2 obtained from the RRDE LSV data. For Cu₃L1, the percentage of H₂O₂ produced during ORR catalysis remains relatively stable with a slight decrease from ~76% near the ORR onset potential to ~63% at -0.15 V vs. the RHE (Fig. 9a). These values are comparable to the reported %H₂O₂ values for Cu-bmpa.²⁰ For Cu₃L2, the initial %H₂O₂ of ~58% near the ORR onset potential decreases more rapidly to $\sim 27\%$ at -0.15 V vs. the RHE (Fig. 9b). This

% H_2O_2 value of ~27% observed for Cu_3L2 at -0.15 V vs. the RHE is significantly lower than the observed %H₂O₂ values for Cu₃L1 and Cu-bmpa.

Additionally, the values for %H₂O₂ were determined by RRDE CA measurements as a function of time. These RRDE CA measurements were performed at applied disk potentials of 0.35, 0.30, 0.20 and 0.0 V vs. the RHE for 5 minutes (ESI 8.3†). As shown in Fig. 9b, the %H₂O₂ values obtained from the RRDE CA data of Cu₃L2 correlate well with the values obtained from the RRDE LSV data. However, the %H₂O₂ values obtained from the RRDE CA data of Cu₃L1 at applied disk potentials of 0.0 and 0.20 V vs. the RHE are significantly lower than the values obtained from the RRDE LSV data (Fig. 9a). These observations are concomitant with an increase of the disk current and a decrease of the ring current over time during CA. This points to the formation of Cu⁰ at the electrode surface which catalyzes the 4-electron reduction of dioxygen, as we have observed previously for Cu-bmpa (ESI 8.3†).20 The involvement of Cu⁰ in the LSV experiments can be excluded on the basis of the aforementioned dipping and microbalance experiments (ESI 7†). Overall, it can therefore be concluded that Cu₃L1 has a comparable product selectivity to Cu-bmpa, while Cu₃L2 has a much higher selectivity for the 4-electron process.

H₂O₂ reduction behavior

As discussed in the previous section, the RRDE LSV data of both Cu₃L1 and Cu₃L2 result in %H₂O₂ values above zero along the entire ORR active potential window. This means that both complexes do not catalyze the full four-electron ORR in the investigated potential window. However, since there is also no complete H₂O₂ selectivity, limitations seem to arise after the initial two-electron reduction of O₂ to H₂O₂. Therefore, the H₂O₂ reduction behavior of both Cu₃L1 and Cu₃L2 was investigated by performing rotating disk electrode (RDE) measurements in phosphate buffer containing 1.1 mM H₂O₂ (Fig. 10). The H₂O₂ concentration amounted to 1.1 mM in order to

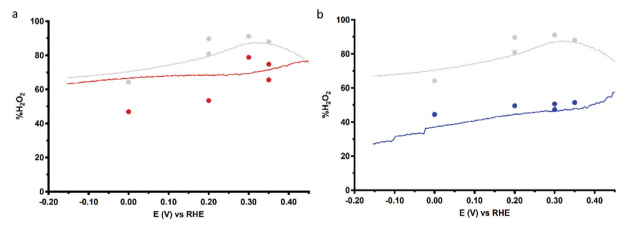


Fig. 9 Percentage of H₂O₂ produced during ORR catalysis (%H₂O₂) obtained from RRDE LSV (lines, 50 mV s⁻¹) and CA (dots) measurements as a function of applied disk potential for 0.1 mM Cu₃L1 (a) and Cu₃L2 (b). The reference %H₂O₂ values for 0.3 mM Cu-bmpa are depicted in grey. Conditions: 0.1 M pH 7 phosphate buffer under 1 atm O2, 293 K, GC disk, Pt ring at 1.2 V vs. RHE, 1600 RPM.

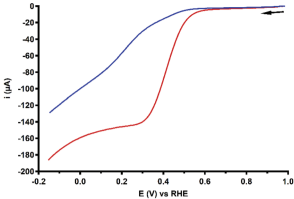


Fig. 10 RDE LSV curves of 0.1 mM $\rm Cu_3L1$ (red) and $\rm Cu_3L2$ (blue) in the presence of 1.1 mM $\rm H_2O_2$ under 1 atm Ar. Conditions: 0.1 M pH 7 phosphate buffer, 293 K, GC disk, 1600 RPM, 50 mV s⁻¹ scan rate.

reproduce the concentration of O_2 in O_2 saturated 0.1 M pH 7 phosphate buffer.^{20,70–72}

The $\rm H_2O_2$ reduction profile obtained by RDE measurements for both $\rm Cu_3L1$ and $\rm Cu_3L2$ did not show the presence of a mass-transport limiting current plateau between 1.0 and -0.15 V νs . the RHE, which we expect to find at 400 μ A according to the Levich equation (ESI 6.3†). This absence of a mass-transport limiting current plateau confirms that the $\rm H_2O_2$ reduction by these trinuclear catalysts must be a relatively slow process, which was shown for $\rm Cu\text{-}bmpa$ and $\rm Cu\text{-}terpy$ previously as well. 20,39

Upon addition of 1.1 mM H₂O₂ to the Cu₃L2 solution in aqueous phosphate buffer, a change in the color and more specifically a change in the UV-Vis spectrum were observed (Fig. S13†). For the triethylbenzene ligand it has been observed previously that oxygen reactivity can induce an aromatic ligand hydroxylation reaction involving an NIH-shift of one of the ethyl substituents on the benzene spacer. 41,42,50 To investigate if this is also induced by H2O2, the ligand was recovered from the complex upon treatment with a strong acid and EDTA to bind the free Cu ions after the exposure of Cu₃L2 to H₂O₂ (ESI 9.2†). Spectroscopy analysis does not suggest a shift of the ethyl substituent as was previously observed. The exact nature of the final structure could not be determined, but the absence of ligand oxidation for Cu₃L1 after the addition of H₂O₂ suggests that the benzylic ethyl-CH₂ substituents of L2 are prone to oxidation upon exposure of Cu₃L2 to high concentrations of H₂O₂.⁷³

An overview of the electrochemical characteristics of Cu_3L1 , Cu_3L2 and Cu-bmpa is given in Table 1.

Discussion

There are four factors that have been considered during this work: the nature of the active species, the selectivity of the reaction, the efficiency of the Cu-mediated ORR, and the catalytic stability of Cu_3L1 and Cu_3L2 .

Table 1 Overview of the electrochemical characteristics of Cu-bmpa, Cu_3L1 , and Cu_3L2

	Cu-bmpa ²⁰	Cu ₃ L1	Cu ₃ L2
$E_{1/2}$ (V vs. RHE)	0.30	0.37	0.50
$\Delta E_{\rm p}$ (mV)	56	105	90
Diffusion coefficient (cm ² s ⁻¹)	2.1×10^{-6}	8.6×10^{-7}	4.8×10^{-7}
$E_{\text{cat/2}}$ (V vs. RHE)	0.37	0.37	0.33
RRDE onset (V vs. RHE)	0.49	0.55	0.52
%H ₂ O ₂ at onset	75	76	58

Active species

The active species during the ORR catalysis by Cu₃L1 and Cu₃L2 are the reduced molecular species of these complexes. The deposition tests and RRDE experiments under Ar illustrate that both Cu₃L1 and Cu₃L2 do deposit on the electrode surface to some extent, yet that the activities of these deposits are negligible. This was confirmed by the low deposited mass found by the EQCM experiments. Also, during CA for several minutes under rotating conditions we did not see an increase of the catalytic activity, unless potentials below 0.2 V vs. RHE were applied. Under these conditions we see a clear build-up of Cu⁰, which is directly visualized by a decrease in %H₂O₂ for Cu₃L1 at these potentials, due to the 4-electron ORR on metallic copper. We do not see these effects in the LSV curves and at prolonged CA above 0.2 V vs. the RHE for both complexes. The formation of substantial amounts of Cu⁰ requires time and negative potentials, which we have reported previously in a study concerning Cu-bmpa. 20 It is therefore likely that the catalytic activity displayed in the LSV curves is due to the ORR mediated by the homogeneous active species of Cu₃L1 and Cu₃L2.

Selectivity

During the ORR three reactions can occur, namely the direct 4-electron reduction of oxygen to water, the 2-electron reduction of oxygen to H_2O_2 , and the subsequent reduction of H_2O_2 to water. ^{19,39} During the direct 4-electron mechanism, no H_2O_2 is evolved, while H_2O_2 is formed as an obligatory intermediate in a [2+2]-stepwise mechanism.

Directly from the onset of the catalytic wave the determined ${}^{\circ}_{3}H_{2}O_{2}$ is significantly lower with $Cu_{3}L2$ compared to that with $Cu_{3}L1$ in the LSV RRDE experiments. In the case of Cu-tmpa we have shown that the build-up of hydrogen peroxide is directly affected by the relative rates between the two electron reduction of dioxygen *versus* the reduction of hydrogen peroxide, and by their relative concentration near the electrode surface. 19,39 In the case of Cu-tmpa this leads to a build-up of hydrogen peroxide, unless the oxygen reduction reaction becomes mass transport limited in oxygen. In the case of $Cu_{3}L1$ and $Cu_{3}L2$ mass transport limitations do not seem to play a role and consequently these catalysts produce hydrogen peroxide over the entire potential domain in the LSV curves. Whereas the oxygen reduction rates of $Cu_{3}L1$ and $Cu_{3}L2$ are fairly similar, there appears to be a significant difference

between the LSV curves of Cu₃L1 and Cu₃L2 in the presence of H₂O₂, with Cu₃L2 being the slower catalyst (Fig. 10).

Dalton Transactions

The slower H₂O₂ reduction by Cu₃L2 is inconsistent with the lower %H₂O₂ observed for this catalyst, compared to Cubmpa and Cu₃L1 (Fig. 9). This indicates that the selectivity must be due to other reasons besides the relative rates of the ORR versus the hydrogen peroxide reduction reaction (HPRR). In other words, Cu₃L2 must carry out the ORR in a different manner compared to Cu₃L1 and Cu-bmpa. The low %H₂O₂ for Cu₃L2 suggests that the selectivity is not a product from freely exchanging H₂O₂ from the coordination sphere of the trinuclear center, but instead must be due to a cooperative effect. Two modes of cooperation may occur. The cooperative effect might be caused by the trinuclear copper site at Cu₂L2 to operate in a similar manner to laccase and facilitate a direct 4-electron reduction reaction leading to a transformation of dioxygen to water without the intermediacy of hydrogen peroxide. However, since H2O2 is still formed along the entire measured potential regime, this is not likely to be the sole form of cooperation. Most likely the improved selectivity of Cu₃L2 towards the overall four electron reduction of dioxygen is that it is difficult for hydrogen peroxide to effectively dissociate from the trinuclear copper site of Cu₃L2, resulting in the alternating reduction of O₂ and H₂O₂ at the catalytic site. We anticipate that this is an effect of the three Cu centers being positioned in close proximity to each other, making the probability of H₂O₂ to diffuse from the catalytic pocket lower.

Efficiency

Due to considerable uncertainty regarding the number of involved Cu centers during ORR catalysis performed by Cu₃L1 and Cu₃L2, quantitative methods to determine the turnover frequencies of the catalysts such as the foot-of-the-wave analysis (FOWA) and the catalytic current enhancement methods without cannot be performed making substantial assumptions.74-78 However, a qualitative description can be put forward by comparison of the RDE LSV profiles obtained during ORR catalysis. Specifically a comparison of the steepness of the ORR profiles provides more insight into the relative catalytic rate. The RDE LSV profiles for ORR catalysis performed by Cu₃L1, Cu₃L2 and Cu-bmpa are depicted in Fig. 11.20 The reductive current for the ORR profile of Cu₃L1 increases faster than for the ORR profile of Cu₃L2, especially between the onset potential and \sim 0.3 V ν s. the RHE. This indicates that the rate for ORR catalysis is higher for Cu₃L1. Additionally, a comparison with the RDE LSV profile for ORR catalysis performed by Cu-bmpa reveals a slower increase in the reductive current for the ORR profiles of the trinuclear catalysts compared to the mononuclear catalyst above ~ 0.2 V vs. the RHE.²⁰ This indicates that the turnover frequencies of the trinuclear catalysts are substantially lower than those of the mononuclear systems reported previously.²⁰

There are several possible explanations for the slow catalysis in these complexes. One reason might be found in the reorganization energy associated with the change of the oxidation state of the Cu ions. ^{79,80} According to the Marcus theory the

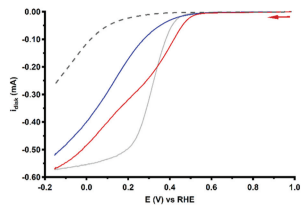


Fig. 11 RDE LSV curves of 0.1 mM Cu_3L1 (red) and Cu_3L2 (blue) under 1 atm O_2 at 1600 RPM. The reference LSV curve of 0.3 mM Cu-bmpa is depicted in grey. The reference voltammogram in the absence of the complex under 1 atm O_2 is depicted as a grey dashed line. Conditions: 0.1 M pH 7 phosphate buffer, 293 K, GC disk, 50 mV s⁻¹ scan rate.

rates of electron transfer reactions are affected by their accompanying reorganization energies.⁸¹ This largely relates to the ability of the ligands to accommodate the metal site at multiple oxidation states, and to switch between the different preferred geometries via facile transitions. It is therefore expected that slow electron transfer kinetics and consecutive slow ORR catalysis are the result of large structural reorganization barriers during the formation of the fully reduced state of Cu₃L1 and Cu₃L2. Due to the steric hindrance by the relatively close proximity of the Cu-bmpa sites of Cu₃L2 to one another compared to the Cu-bmpa sites of Cu₃L1, this effect is more pronounced in Cu₃L2. The lower catalytic currents may also be caused by the lower diffusion rate of the trinuclear complexes compared to for example Cu-bmpa (Table 1). This results in a relatively low number of catalytic sites being reduced by the cathode compared to those in the case of catalysts with higher diffusion constants.

Stability

With UV-Vis spectroscopy we have shown that both Cu₃L1 and Cu₃L2 are stable over prolonged time in a 0.1 M pH 7 phosphate buffer. Moreover, the UV-Vis measurement results before and after ORR experiments remained unchanged. However, at high H₂O₂ concentration, the trinuclear complex Cu₃L2 suffers from intrinsic stability problems. It seems that in particular the ethylene functionalities that force all three copper sites towards the same plane of the aromatic node of L2 are susceptible towards intramolecular oxidation reactions in the presence of millimolar concentrations of hydrogen peroxide. 82,83 However, this structural change for Cu₃L2 was only observed upon addition of large quantities of H2O2 and was not observed during ORR catalysis, where high concentrations of peroxide were avoided. Therefore, this structural change upon H₂O₂ addition is not expected to play a role in ORR catalysis.

Paper

Conclusions

We have studied the effect of cooperativity between two or three copper sites on the catalytic activity and selectivity of the ORR. Although the catalytic currents are lower than those for freely rotating and diffusing single site complexes, our results show that the selectivity of the copper mediated ORR was significantly enhanced towards the overall 4-electron process due to a cooperative effect between three copper sites that have been positioned in close proximity.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

The authors gratefully acknowledge the Magnetism Competence Center at the Jean Lamour Institute in Nancy, France for the support during SQUID measurements.

Financial support was provided by the European Research Council (ERC starting grant 637556 Cu4Energy to D. G. H. H.).

References

- 1 D. G. Nocera, ChemSusChem, 2009, 2, 387-390.
- 2 A. Kirubakaran, S. Jain and R. K. Nema, Renewable Sustainable Energy Rev., 2009, 13, 2430-2440.
- 3 S. Shiva Kumar and V. Himabindu, Mater. Sci. Energy Technol., 2019, 2, 442-454.
- 4 C. Song and J. Zhang, in PEM Fuel Cell Electrocatalysts and Catalyst Layers, ed. J. Zhang, Springer, London, UK, 2008, pp. 89-134.
- 5 Z. W. Seh, J. Kibsgaard, C. F. Dickens, I. Chorkendorff, J. K. Nørskov and T. F. Jaramillo, Science, 2017, 355, eaad4998.
- 6 K. L. Hsueh, D. T. Chin and S. Srinivasan, J. Electroanal. Chem. Interfacial Electrochem., 1983, 153, 79-95.
- 7 Y. Wang, D. F. Ruiz Diaz, K. S. Chen, Z. Wang and X. C. Adroher, Mater. Today, 2020, 32, 178-203.
- 8 P. Giardina, V. Faraco, C. Pezzella, A. Piscitelli, S. Vanhulle and G. Sannia, Cell. Mol. Life Sci., 2010, 67, 369-385.
- 9 H. Komori and Y. Higuchi, J. Biochem., 2015, 158, 293-298.
- 10 E. I. Solomon, A. J. Augustine and J. Yoon, Dalton Trans., 2008, 1, 3921-3932.
- 11 S. M. Jones and E. I. Solomon, Cell. Mol. Life Sci., 2015, 72, 869-883.
- 12 M. R. Tarasevich, A. I. Yaropolov, V. A. Bogdanovskaya and S. D. Varfolomeev, J. Electroanal. Chem. Interfacial Electrochem., 1979, 104, 393-403.
- 13 V. Soukharev, N. Mano and A. Heller, J. Am. Chem. Soc., 2004, 126, 8368-8369.

- 14 M. S. Thorum, C. A. Anderson, J. J. Hatch, A. S. Campbell, N. M. Marshall, S. C. Zimmerman, Y. Lu and A. A. Gewirth, J. Phys. Chem. Lett., 2010, 1, 2251-2254.
- 15 M. J. Moehlenbrock and S. D. Minteer, Chem. Soc. Rev., 2008, 37, 1188-1196.
- 16 C. J. Cramer and W. B. Tolman, Acc. Chem. Res., 2007, 40, 601-608.
- 17 S. Kakuda, R. L. Peterson, K. Ohkubo, K. D. Karlin and S. Fukuzumi, J. Am. Chem. Soc., 2013, 135, 6513-6522.
- 18 S. Itoh, Acc. Chem. Res., 2015, 48, 2066-2074.
- 19 M. Langerman and D. G. H. Hetterscheid, Angew. Chem., Int. Ed., 2019, 58, 12974-12978.
- 20 N. W. G. Smits, B. van Dijk, I. de Bruin, S. L. T. Groeneveld, M. A. Siegler and D. G. H. Hetterscheid, Inorg. Chem., 2020, **59**, 16398-16409.
- 21 H. Oh, S. Choi, J. Y. Kim, H. S. Ahn and S. Hong, Chem. Commun., 2019, 55, 12659-12662.
- 22 P. Vasudevan, Santosh, N. Mann and S. Tyagi, Transition Met. Chem., 1990, 15, 81-90.
- 23 S. Fukuzumi, H. Kotani, H. R. Lucas, K. Doi, T. Suenobu, R. L. Peterson and K. D. Karlin, J. Am. Chem. Soc., 2010, 132, 6874-6875.
- 24 M. S. Thorum, J. Yadav and A. A. Gewirth, Angew. Chem., Int. Ed., 2009, 48, 165-167.
- 25 G. A. Goenaga, A. Belapure, C. Zhang, A. Papandrew, S. Foister and T. Zawodzinski, ECS Trans., 2011, 41, 1193-1205.
- 26 M. Kato, N. Oyaizu, K. Shimazu and I. Yagi, J. Phys. Chem. C, 2016, **120**, 15814–15822.
- 27 R. Venegas, K. Munoz-Becerra, L. Lemus, A. Toro-Labbe, J. H. Zagal and F. J. Recio, J. Phys. Chem. C, 2019, 123, 19468-19478.
- 28 J. Zhang and F. C. Anson, Electrochim. Acta, 1993, 38, 2423-2429.
- 29 C. C. L. McCrory, X. Ottenwaelder, T. D. P. Stack and C. E. D. Chidsey, J. Phys. Chem. A, 2007, 111, 12641-12650.
- 30 L. M. Mirica, X. Ottenwaelder and T. D. P. Stack, Chem. Rev., 2004, 104, 1013-1046.
- 31 E. A. Lewis and W. B. Tolman, Chem. Rev., 2004, 104, 1047-1076.
- 32 S. Hong, Y.-M. Lee, K. Ray and W. Nam, Coord. Chem. Rev., 2017, 334, 25-42.
- 33 C. E. Elwell, N. L. Gagnon, B. D. Neisen, D. Dhar, A. D. Spaeth, G. M. Yee and W. B. Tolman, Chem. Rev., 2017, 117, 2059–2107.
- 34 M. A. Thorseth, C. E. Tornow, E. C. M. Tse and A. A. Gewirth, Coord. Chem. Rev., 2013, 257, 130-139.
- 35 J. A. Halfen, S. Mahapatra, E. C. Wilkinson, S. Kaderli, V. G. Young, L. Que, A. D. Zuberbühler and W. B. Tolman, Science, 1996, 271, 1397.
- 36 B. van Dijk, J. P. Hofmann and D. G. H. Hetterscheid, Phys. Chem. Chem. Phys., 2018, 20, 19625-19634.
- 37 J. Serrano-Plana, I. Garcia-Bosch, A. Company and M. Costas, Acc. Chem. Res., 2015, 48, 2397-2406.
- 38 L. Tahsini, H. Kotani, Y.-M. Lee, J. Cho, W. Nam, K. D. Karlin and S. Fukuzumi, Chem. - Eur. J., 2012, 18, 1084-1093.

Dalton Transactions Paper

- 39 M. Langerman and D. G. H. Hetterscheid, *ChemElectroChem*, 2021, **8**, 2783–2791.
- 40 S. Fukuzumi, Y.-M. Lee and W. Nam, *ChemCatChem*, 2018, **10**, 9–28.
- 41 K. D. Karlin, Q. F. Gan, A. Farooq, S. Liu and J. Zubieta, *Inorg. Chem.*, 1990, 29, 2549–2551.
- 42 K. D. Karlin, Q.-F. Gan and Z. Tyeklár, *Chem. Commun.*, 1999, 1, 2295–2296.
- 43 E. Y. Tsui, M. W. Day and T. Agapie, *Angew. Chem., Int. Ed.*, 2011, **50**, 1668–1672.
- 44 D. Maiti, J. S. Woertink, R. A. Ghiladi, E. I. Solomon and K. D. Karlin, *Inorg. Chem.*, 2009, 48, 8342–8356.
- 45 D. Lionetti, M. W. Day and T. Agapie, *Chem*, 2013, 4, 785–790
- 46 E. C. M. Tse, D. Schilter, D. L. Gray, T. B. Rauchfuss and A. A. Gewirth, *Inorg. Chem.*, 2014, 53, 8505–8516.
- 47 X. Engelmann, E. R. Farquhar, J. England and K. Ray, *Inorg. Chim. Acta*, 2018, **481**, 159–165.
- 48 N. Thiyagarajan, D. Janmanchi, Y. F. Tsai, W. H. Wanna, R. Ramu, S. I. Chan, J. M. Zen and S. S. F. Yu, *Angew. Chem., Int. Ed.*, 2018, 57, 3612–3616.
- 49 E. Salvadeo, L. Dubois and J.-M. Latour, *Coord. Chem. Rev.*, 2018, 374, 345–375.
- 50 H. Ohi, Y. Tachi and S. Itoh, *Inorg. Chem.*, 2006, **45**, 10825–10835.
- 51 E. C. Brown, B. Johnson, S. Palavicini, B. E. Kucera, L. Casella and W. B. Tolman, *Dalton Trans.*, 2007, 1, 3035–3042.
- 52 A. M. Geer, C. Musgrave Iii, C. Webber, R. J. Nielsen, B. A. McKeown, C. Liu, P. P. M. Schleker, P. Jakes, X. Jia, D. A. Dickie, J. Granwehr, S. Zhang, C. W. Machan, W. A. Goddard and T. B. Gunnoe, ACS Catal., 2021, 11, 7223-7240.
- 53 K. D. Karlin, Q. F. Gan, A. Farooq, S. Liu and J. Zubieta, *Inorg. Chem.*, 1990, **29**, 2549–2551.
- 54 C. Walsdorff, S. Park, J. Kim, J. Heo, K.-M. Park, J. Oh and K. Kim, *Dalton Trans.*, 1999, **1**, 923–930.
- 55 M. Komiyama, S. Kina, K. Matsumura, J. Sumaoka, S. Tobey, V. M. Lynch and E. Anslyn, *J. Am. Chem. Soc.*, 2002, 124, 13731–13736.
- 56 Y. Zhao, J. Zhu, W. He, Z. Yang, Y. Zhu, Y. Li, J. Zhang and Z. Guo, *Chem. – Eur. J.*, 2006, **12**, 6621–6629.
- 57 N. F. Chilton, R. P. Anderson, L. D. Turner, A. Soncini and K. S. Murray, *J. Comput. Chem.*, 2013, 34, 1164–1175.
- 58 Z. Boulsourani, V. Tangoulis, C. P. Raptopoulou, V. Psycharis and C. Dendrinou-Samara, *Dalton Trans.*, 2011, 40, 7946–7956.
- 59 D. J. Iverson, G. Hunter, J. F. Blount, J. R. Damewood and K. Mislow, J. Am. Chem. Soc., 1981, 103, 6073–6083.
- 60 J. C. Barnes, J. A. Chudek, G. Hunter, A. J. Blake, P. J. Dyson, B. F. G. Johnson and W. Weissensteiner, J. Chem. Soc., Faraday Trans., 1995, 91, 2149–2153.

- 61 V. Marks, H. E. Gottlieb, A. Melman, G. Byk, S. Cohen and S. E. Biali, *J. Org. Chem.*, 2001, **66**, 6711–6718.
- 62 W. Weissensteiner, A. Gutierrez, M. D. Radcliffe, J. Siegel, M. D. Singh, P. J. Tuohey and K. Mislow, *J. Org. Chem.*, 1985, **50**, 5822–5827.
- 63 S. Choksakulporn, A. Punkvang and Y. Sritana-Anant, J. Mol. Struct., 2015, 1082, 97–102.
- 64 N. Elgrishi, K. J. Rountree, B. D. McCarthy, E. S. Rountree, T. T. Eisenhart and J. L. Dempsey, *J. Chem. Educ.*, 2018, 95, 197–206.
- 65 M. J. Powers and T. J. Meyer, J. Am. Chem. Soc., 1978, 100, 4393–4398.
- 66 J. E. Sheats, R. S. Czernuszewicz, G. C. Dismukes, A. L. Rheingold, V. Petrouleas, J. Stubbe, W. H. Armstrong, R. H. Beer and S. J. Lippard, J. Am. Chem. Soc., 1987, 109, 1435–1444.
- 67 M. M. Morrison and D. T. Sawyer, J. Am. Chem. Soc., 1977, 99, 257–258.
- 68 R. Lomoth, P. Huang, J. Zheng, L. Sun, L. Hammarström, B. Åkermark and S. Styring, *Eur. J. Inorg. Chem.*, 2002, **2002**, 2965–2974.
- 69 R. Zhou, Y. Zheng, M. Jaroniec and S.-Z. Qiao, *ACS Catal.*, 2016, 6, 4720–4728.
- 70 A. Schumpe, I. Adler and W.-D. Deckwer, *Biotechnol. Bioeng.*, 1978, **20**, 145–150.
- 71 D. Tromans, Ind. Eng. Chem. Res., 2000, 39, 805-812.
- 72 W. Xing, M. Yin, Q. Lv, Y. Hu, C. Liu and J. Zhang, *Rotating Electrode Methods and Oxygen Reduction Electrocatalysts*, Elsevier, Amsterdam, 2014, pp. 1–31.
- 73 M. M. Rahman, M. G. Ara, M. S. Rahman, M. S. Uddin, M. N. Bin-Jumah and M. M. Abdel-Daim, *J. Nanomater.*, 2020, 2020, 1–20.
- 74 C. Costentin, S. Drouet, M. Robert and J. M. Saveant, *J. Am. Chem. Soc.*, 2012, **134**, 19949–19950.
- 75 E. S. Rountree, B. D. McCarthy, T. T. Eisenhart and J. L. Dempsey, *Inorg. Chem.*, 2014, **53**, 9983–10002.
- 76 C. Costentin and J. M. Saveant, *ChemElectroChem*, 2014, 1, 1226–1236.
- 77 D. J. Wasylenko, C. Rodriguez, M. L. Pegis and J. M. Mayer, *J. Am. Chem. Soc.*, 2014, **136**, 12544–12547.
- 78 A. M. Appel and M. L. Helm, ACS Catal., 2014, 4, 630-633.
- 79 E. Garribba and G. Micera, *J. Chem. Educ.*, 2006, **83**, 1229–1232.
- 80 E. W. Dahl and N. K. Szymczak, *Angew. Chem., Int. Ed.*, 2016, 55, 3101–3105.
- 81 R. A. Marcus, Rev. Mod. Phys., 1993, 65, 599-610.
- 82 S. Mahapatra, V. G. Young, S. Kaderli, A. D. Zuberbuhler and W. B. Tolman, *Angew. Chem., Int. Ed. Engl.*, 1997, **36**, 130–133.
- 83 S. Mahapatra, J. A. Halfen and W. B. Tolman, *J. Am. Chem. Soc.*, 1996, **118**, 11575–11586.