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Introduction

Significant, attention has been focused in recent years on the synthesis of transition metal based dendrimer structures owing to their diverse applications in various fields.¹ In particular, these dendrimers, in many cases, allow synergistic interactions between the individual transition metal centers in carrying out a variety of important transformations. The organooxotin clusters are in particular attractive because of the diversity of arrangements that they adopt, such as ladder, O-capped, cube, butterfly, drum, one, two and three-dimensional structures, $(1D, 2D,$ and $3D)$.²⁻⁶ Furthermore, incorporation of redox-active transition-metal centers into the stannox-

Catalytic dioxygen reduction mediated by a tetranuclear cobalt complex supported on a stannoxane core†

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The synthesis, spectroscopic characterization (infrared, electron paramagnetic resonance and X-ray absorption spectroscopies) and density functional theoretical calculations of a tetranuclear cobalt complex Co₄L1 involving a nonheme ligand system, L1, supported on a stannoxane core are reported. Co₄L1, similar to the previously reported hexanuclear cobalt complex Co₆L2, shows a unique ability to catalyze dioxygen (O₂) reduction, where product selectivity can be changed from a preferential 4e⁻/4H⁺ dioxygen-reduction (to water) to a 2e[−]/2H+ process (to hydrogen peroxide) only by increasing the temperature from −50 to 30 °C. Detailed mechanistic insights were obtained on the basis of kinetic studies on the overall catalytic reaction as well as by low-temperature spectroscopic (UV-Vis, resonance Raman and X-ray absorption spectroscopies) trapping of the end-on μ -1,2-peroxodicobalt(III) intermediate 1. The Co₄L1- and Co₆L2-mediated O₂-reduction reactions exhibit different reaction kinetics, and yield different ratios of the 2e[−]/2H⁺ and 4e[−]/4H⁺ products at –50 °C, which can be attributed to the different stabilities of the μ-1,2-peroxodicobalt(III) intermediates formed upon dioxygen activation in the two cases. The deep mechanistic insights into the transition-metal mediated dioxygen reduction process that are obtained from the present study should serve as useful and broadly applicable principles for future design of more efficient catalysts in fuel cells. PAPER

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ane clusters has previously led to the demonstration of important reactivity patterns.^{7,8} For example, an extensive cooperative effect between the Cu centers was observed during the cleavage of supercoiled DNA catalyzed by a hexanuclear Cu– porphyrin complex, supported on a stannoxane core.^{7a} In our group we have previously demonstrated the ability of a nonheme stannoxane based hexanuclear ligand system to undergo O–O bond formation^{7b} and O–O bond cleavage reactions,⁸ when bound to iron(π) and cobalt(π) centers, respectively. In the present manuscript we report the synthesis, characterization and X-ray structure of a tetranuclear stannoxane based non-heme ligand system (L1), and a detailed kinetic study of the catalytic dioxygen reduction reaction mediated by the corresponding cobalt complex Co₄L1. Notably, catalytic reductions of O_2 to water or H_2O_2 have tremendous technological significance. $9-12$ However, in contrast to biology, where cheap and readily available transition-metal complexes of Fe, and Cu are employed for O_2 reduction,¹³ high loadings of a precious metal like platinum is warranted for achieving appreciable reactivity during abiological $O₂$ -reduction reactions.^{12d} Thus the present study is relevant to the ongoing research activities that are being dedicated towards the development of $O₂$ reduction catalysts based on nonprecious

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metals.¹⁴ Furthermore, it provides deep mechanistic insights into the factors that control two- vs. four-electron reductions of $O₂$, thereby providing useful and broadly applicable principles for the future design of more efficient O_2 reduction catalysts.

Results and disscussion

Synthesis and characterisation of L1

The condensation reaction (Schemes S1 and S2†) of equimolar amounts of di-n-butyltin oxide and 4-(1,3-bis(2-pyridylmethyl)-

Fig. 1 Comparison of the distances between the metal binding sites in L1 (top) and L2 (bottom). X-ray crystal structure of L1 and L2 with 30% ellipsoid probability of the atoms. Hydrogen atoms and the n -butyl ("Bu-) groups on the tin atoms have been omitted for clarity. Color code: nitrogen-blue; carbon-grey; oxygen-red; tin-green.

2-imidazolidinyl)benzoic acid in toluene afforded L1 as a pale yellow solid. The molecular structure of L1 shows that a planar $Sn₄O₂$ core supports the four metal-binding sites (Fig. 1: top). This is in contrast to the situation reported earlier for the hexanuclear non-heme ligand system L2, where six metal-binding sites were located in a wheel-like arrangement around a central $Sn₆O₆$ prismane core (Fig. 1: bottom). The stannoxane core in Ligand L1 adopts a ladder framework with two central and two terminal tin atoms. The tetranuclear structure of L1 is maintained in solution. ¹¹⁹Sn NMR spectrum of L1 exhibits two sharp singlets of equal intensity at −210.82 ppm and −213.81 ppm (Fig. S1†), which is the characteristic signature for a planar Sn_4O_2 core.^{2–6} The infrared spectrum shows four vibrations at 1622 cm^{-1} , 1591 cm^{-1} , 1569 cm^{-1} , and 1545 cm⁻¹ for the carboxyl absorptions ($\nu_{\rm{COO}}$), and one strong band at 682 cm⁻¹ assigned to $\nu_{\text{Sn-O}}$ for the Sn₄O₂ core (Fig. S2†).

Synthesis and characterization of $Co₄Li$

The reaction of L1 with 4 equiv. of $Co(CF_3SO_3)_2$ in acetone yields Co₄L1 as a dark yellow powder in 70% yield (Scheme 1). The C, H, and N content of $Co₄Li$, determined by elemental analysis, established the presence of four cobalt atoms per tetrameric ligand, with two triflates associated with each cobalt (see ESI†).

The infrared spectrum of $Co₄Li$ depicts the characteristic vibrations of the Sn_4O_2 core at 1625 cm⁻¹, 1593 cm⁻¹, 1572 cm−¹ , 1549 cm−¹ , and 682 cm−¹ (Fig. S2†). These vibrations are only slightly shifted relative to that of L1, which reveals that the tetranuclear arrangement is also maintained in $Co₄Li$.

Electronic and structural information of $Co₄Li$ were obtained from X-ray absorption spectroscopy (XAS) in conjunction with density functional theory (DFT) calculations. The near edge structure (XANES) was used for determination of the oxidation states, whereas the extended fine structure (EXAFS) unraveled the local site geometries around the Co atoms. The spectra are displayed in Fig. 2, the corresponding fit values are collected in Table 1. The XANES spectrum of $Co₄Li$ (blue trace) is displayed together with spectra from Co reference compounds of known oxidation states $(Co^{2+}, Co^{2.66+}, Co^{3+})$, see Fig. 2a; it is consistent with a Co^{2+} oxidation state in $Co₄Li$

Scheme 1 Synthesis of the tetra-nuclear cobalt(II) complex (Co_4L1) from the tetra-nuclear stannoxane ligand (L1) and the formation of the cobalt(III)–peroxo complex (1).

Fig. 2 (a) Co K-edge XANES spectra of reference compounds (black: $Co(OH₂)₆(NO₃)₂$, dark grey: $Co₃O₄$ spinel, grey: $CoOOH$) and samples (blue: Co4L1, red: 1). The horizontal line at 0.5 is used for determination of the oxidation states. (b) Oxidation states of samples $Co₄LI$ and 1 derived from linear fit lines extracted from the 0.5 edge-rise positions of reference compounds (black line and squares) and derived from the integral method (grey line and squares). Both methods result in equal oxidation state assignments. (c) k -Space EXAFS spectra (k^3 weighted, colored) and respective fits (grey). (d) Fourier transform of the EXAFS spectra (colored) and fits (grey).

Table 1 EXAFS fit parameters for $Co₄ L1$ and 1

Model	Shell	N^a	R(A)	Err	$\sigma(\AA)$	Err
Co ₄ L1	$Co-N$	5	2.14	0.01	0.056	0.007
$R_f = 18.4$	$Co-N$	1	2.34	0.05	0.056	
$E_0 = 0.74$	$Co-C$	3	2.94	0.03	0.056	
d_{av} = 2.17 Å	$Co-C$	2	3.06	0.05	0.056	
1	$Co-N$	$\overline{4}$	1.91	0.01	0.036	0.009
$R_f = 15.0$	$Co-O$	1	2.02	0.03	0.036	
$E_0 = 2.82$	$Co-C$	3	2.78	0.03	0.036	
$d_{av} = 1.93 \text{ Å}$	$Co-C$	$\overline{2}$	2.92	0.05	0.036	

^a Value kept constant in the final refinement. Amplitude reduction factor $S_{02} = 0.95$. *N* represents the EXAFS coordination number, *R* the absorber-backscatter distance and σ the Debye Waller parameter.

(Fig. 2b). The EXAFS of $Co₄Li$ could be well fitted by four shells, with one shorter N-shell with coordination number (N) of 5, a longer N/O-shell with $N = 1$, and two C-shells with $N = 3$ and 2 (Fig. 2c and d). Attempts to fit $Co₄Li$ with a sum of $N = 5$ in the first two shells (instead of 6) significantly worsen the fit parameters. In principle, there are up to nine C-atoms within a radius of 3.5 Å around the Co-atom, however, due to the pronounced inhomogeneity of the Co–C distances, these shells may partially cancel each other out. The average Co–O/N distance is found to be 2.17 Å.

Since there are no X-ray diffraction (XRD) structures available for $Co₄Li$ and the Co atoms are mainly surrounded by O, N and C atoms with similar scattering properties, the EXAFS fits may suffer from non-uniqueness and misinterpretations. In order to reduce this problem as well as to obtain suitable phase functions for the fits, DFT calculations were conducted in the experimentally observed (from EPR; Fig. S3†) $S = 3/2$ spin state for a series of potential structural variants of the monomeric subsection of the organic ligands, starting from the modified XRD structure of the tetrameric stannoxane ligand (see Fig. 3 and S4†). This approach is justified as there are no intra-molecular electronic interactions detectable between adjacent $Co(n)$ sites, as evident from the X-band EPR spectrum of $Co₄Li$, which exhibits a major axial signal with effective g'^{\perp} = 4.01 and $g''^{\parallel} \approx 2.0$ corresponding to the S = 3/2 ground state (Fig. S3†). Structural variants include the

Fig. 3 (a) The DFT calculated coordination environment of the individual Co-centers in Co₄L1, which matches best with the experimental data. The straight green lines reveal the N-capped O_2N_3H -octahedron environment of the central Co(II) center, by taking into account the agostic C–H⋯Co interaction. (b) DFT model of 1 with fixed carboxylate C⋯C atom distance of 6.43123 Å according to the XRD structure of the free ligand. In 1, L1 acts as a tetradentate ligand with one of the nitrogen atoms staying away from the coordination environment of Co.

coordination of triflate (OTf) and/or solvent acetone molecules in cis- or trans-orientations, and with or without inherent molecular symmetry, (see legend of Fig. S4†). Four of the six DFT models show hexa-coordinated Co (n) (no. 1, 2, 5, and 6), and the other two show penta-coordination (no. 3 and 4; see Table S1[†]). Since unrestrained EXAFS fits of $Co₄Li$ clearly indicate hexa-coordination, the corresponding DFT models are considered to be closer to the actual structure of $Co₄Li$. The average Co–X $(X = N, O)$ bond distances, however, vary in the narrow range of 2.09 to 2.19 Å in all six models, which can hardly be discriminated by EXAFS, but all of them are close to the 2.17 Å obtained from the experiments.

The lowest molecular energy is obtained for the hexa-coordinated model 5, followed by penta-coordinated model 3, which is only 5 kJ mol⁻¹ higher in energy. Fine structural details are visible in the EXAFS wave, which promises more insight into the real structure than geometric and energetic considerations alone. Accordingly, EXAFS was calculated for all six small DFT models and compared to the experimental spectra of Co₄L1, see Fig. S5 in the ESI.[†] Here again, model 5 apparently gives the best match to the spectrum of $Co₄Li$, followed by the other hexa-coordinate models 2, 6 and 1, whereas the two penta-coordinate models 3 and 4 give the worst match. Taking all results into account – geometry, energy and EXAFS – DFT model 5 seems to be the closest representative for the structure underlying in the experimental data of $Co₄LI$ we have so far. In this model, the six Oand N-atoms bound to the central Co-atom are aligned in a low symmetrical fashion, which might be described as quadratic-pyramidal (O_2N_3) with one (extra) N-atom below but close to the quadratic plane, (see Fig. 3). However, these results seemingly are in contradiction with the XANES spectrum of Co₄L1, which looks like typical octahedral or trigonal-bipyramidal (i.e. high local coordination symmetry) compounds, e.g. the hexa-aqua $Co(n)$ compound used for reference (black line in Fig. 2a).

The answer to this riddle might be the potentially underrated electronic and steric effect of the agostic proton in DFT model 5, which is part of the carboxylated phenyl group, see Fig. 3. Since the organic ligand system has only limited flexibility, an unoccupied coordination site can be filled by a C–H⋯Co contact, which changes the picture. Taking the H-atom into account, the coordination geometry is rather an N-capped O_2N_3H -octahedron, see green lines in Fig. 3, than a square pyramid. In order to understand this in more detail, the electronic situation of model 5 was thoroughly analyzed by means of Real-Space Bonding Indicators (RSBIs) extracted from the computed electron density (ED, Fig. S6†). Fig. S6b† shows the spin-density, the majority of which is localized at the Co-atom, as expected, and with minor contributions at all six non-H-atoms. Bond topological analysis of the ED according to the Atoms-In-Molecules $(AIM)^{15}$ theory, however, finds bond critical points (bcp) and thus bond paths to all seven O-, N- and H-atoms, see Fig. S6c.† AIM theory also provides atomic basins. Mapping the ED distribution on them discovers bonding regions and strength of chemical interactions. The

AIM atomic Co basin has the basic shape of a cube (typical for octahedral ligand sphere) with one edge cropped by the capping N-atom, see Fig. S6d.† The more interesting point, however, is that the shape of the basin is also flat along the Co⋯H axis, although the agnostic interaction is quite weak (only little ED accumulation on the respective cube face). This "regular shape" of the Co-atom is also visible applying the Non-Covalent interactions Index (NCI) ,¹⁶ which uncovers noncovalent bonding aspects of strong medium and even very weak atom–atom contacts, see Fig. S6e.† Ring-shaped bluecolored NCI basins indicate dominating covalent bonding aspects (one O, one N), whereas disc-shaped blue-colored NCI basins indicate dominating non-covalent bonding aspects (one O, three N). The agostic Co \cdots H contact is represented by a flat and extended greenish-blue colored NCI basin, being typical for weak non-covalent interactions, such as H⋯H or metallophilic contacts. AIM and NCI are complemented by the Electron Localizability Indicator $(ELI-D)$,¹⁷ which dissects realspace into regions/basins of (non-) bonding electron pairs, resembling in a way the Lewis-picture of chemical bonding. An iso-surface representation is shown in Fig. S6f.† Highlighted (solid, green) are the six non-bonding d-electron ELI-D basins of the Co-atom, which altogether form a regular polyhedron in order to minimize electron–electron repulsion to the electron pairs from the electron donating ligand atoms, according to the well-known "key-lock" arrangement in transition metal chemistry. Paper More continuation of criticle (OTI) and/or solvent accessor and entired on 2020. Download by the entired in the case of the selection of the selecti

Co4L1 catalyzed dioxygen reduction reaction

The evaluation of the catalytic activity of $Co₄Li$ towards oxygen reduction was carried out using the Fukuzumi and Guilard's method;¹⁸ decamethyl-ferrocene was employed as a one electron donor, triflic (TfOH) or fluoroboric (HBF₄) acids were used as proton source, and, in their presence, O_2 was set to react with a catalytic amount of $Co₄Li$ in acetone. The occurrence of the oxygen reduction reaction was proved by the formation of decamethylferrocenium ion (Fe^{*+}) with a characteristic absorption band at 780 nm (Fig. 4; $\varepsilon_{780 \text{ nm}}$ = 520 M⁻¹ cm⁻¹).^{19,20} Notably, the rate and yield of formation of Fc^* is not significantly affected by the nature of the proton source (TfOH or HBF_4 Fig. $S7df$), thereby suggesting that the conjugate bases (OTf⁻ or BF₄⁻) play no major role in controlling the efficiency of the O_2 -reduction reactions. However, the concentration of Fc^{*+} formed in the complex $Co₄Li$ -catalyzed reduction of $O₂$ by Fc* is dependent on the temperature at which the reactions were performed (Fig. 4 bottom, S7a–c†). At 30 $\rm{^{\circ}C}$ 0.35 mM of Fc*⁺ ion is generated in the reaction, which corresponds approximately twice that of the O_2 concentration (0.18 mM). Thus, only two-electron reduction of O_2 occurs at 30 °C. With decreasing temperature, the amount of Fc^{*+} generated from O_2 reduction increases, presumably because of the increasing contribution of the four-electron reduction of O_2 . At 25 °C the amount of Fc^{*+} formed is 0.44 mM, which is 2.5 times that of the O_2 concentration. The mechanism shifts predominantly to a four-electron reduction process at -50 °C; the amount of Fc^{*+} gener-

Fig. 4 Top: Spectral changes associated with the catalytic reduction of dioxygen by $Co₄LI$ (0.04 mM) in the presence of Fc* and TFA. Bottom: Time profiles of the formation of Fc^{*+} monitored at 780 nm during the reduction of O_2 (0.18 mM) by Fc* (3 mM) and Co_4L1 (0.04 mM) in the presence of TFA (10 mM) in acetone at 30 °C, 25 °C as well as at −50 °C.

ated is 0.62 mM, which represents 3.45 equiv. relative to the initial concentration of O_2 (0.18 mM).

¹H-NMR spectrum (Fig. 5a) of the reaction mixture of $Co₄Li$ (0.04 mM) , Fc* (3 mM) , TFA (10 mM) and O_2 (0.18 mM) at 25 °C in d_6 -acetone further confirms the change in mechanism from a predominantly 2e[−] reduction of O_2 (to H₂O₂) at 30 °C to a 4e[−] reduction to H₂O at –50 °C. The solution after the Co₄L1 catalysed O_2 reduction at 30 °C shows a signal at 3.88 ppm, whose position is upshifted relative to the signal at 3.93 ppm obtained for an authentic H_2O_2/H_2O mixture (70 weight % H_2O_2 basis) in d₆-acetone at 25 °C. Note that the signal corresponding to a H_2O_2/H_2O mixture undergoes a downshift with increasing amounts of water in the mixture; pure water shows a signal at 4.61 ppm. Thus for a catalytic reaction at 30 °C, a $>70\%$ H₂O₂ concentration can be inferred. When the catalysis is performed at 25 °C the signal gets downshifted to 3.99 ppm, thereby confirming the presence of the higher amount of water as the 4e[−] reduction product. The resultant solution after the catalytic O₂-reduction reaction at −50 °C shows a signal at 4.20 ppm, which lies between the signals at 4.44 ppm and 4.05 ppm obtained for authentic 15% and 30% H_2O_2/H_2O mixtures, respectively. A turnover number (TON) of 28.7 during a lapse of 2000 s was determined in acetone at −50 °C. The TON decreased linearly with increasing temperature to a value of 9 at 25 °C (Fig. S8†).

Reaction of $Co₄LI$ with dioxygen to form 1

An acetone solution of $Co₄Li$, when treated with $O₂$ saturated acetone at −50 °C, results in the formation of an orange

Fig. $5⁻¹H-NMR$ spectra of the products formed during the complex $Co₄LI$ catalyzed dioxygen reduction reactions at 25 °C, 30 °C and −50 °C [Co₄L1 (0.01 mM), Fc* (3 mM), TFA (10 mM) and O₂ (0.18 mM)] in acetone- d_6 and comparison with authentic samples containing 70%, 30%, 15% and 0% (weight percent H_2O_2 basis) H_2O_2/H_2O mixtures. All the ¹H-measurements were performed at 25 °C. The results obtained for $Co₄$ L1 are then compared with that for $Co₆$ L2 in Table 2.

Table 2 Temperature dependence of the catalytic $O₂$ reductions mediated by Co₄L1 and Co₆L2

Temp.	Catalyst used	mМ	Equiv. of O_2 reduced	% of H_2O_2 formed (from NMR)
$-50~^0$ C	Co ₄ Li	0.01	3.46	20%
+25 $\mathrm{^{0}C}$		0.01	2.5	64%
$+30\text{ °C}$		0.01	1.92	74%
$-50~^0$ C	Co ₆ L2	0.01	3.80	Ω
$+25\,^0C$		0.01	1.88	70%

species 1 with an intense absorption maximum λ_{max} (ε_{max} , $\text{M}^{-1} \text{ cm}^{-1}$) centered at 464 nm (12 200 $\text{M}^{-1} \text{ cm}^{-1}$). As the temperature is increased, the absorption band at 464 nm due to 1 is decreased (Fig. 6: top one). This process is reversible in the temperature range −50 to 30 °C. The resonance Raman (rR) spectrum (Fig. 6: bottom one) of 1 in acetone- d_6 displays two isotopically sensitive vibrational bands at 862 (O–O stretching mode of a peroxo ligand) and 595 cm^{-1} (Co–O stretching mode), which are downshifted to 808 and 561 cm⁻¹, respectively, in ${}^{18}O_2$ prepared samples.

XAS studies were also performed to probe the oxidation state and the coordination environment of Co in 1. The XANES spectra of 1 when compared with that of $Co₄Li$ and other reference compounds reveals an almost complete oxidation from Co^{2+} to Co^{3+} during the transformation of Co_4L1 to 1. Additionally, the edge shape of 1 shows minor altera-

Fig. 6 Top: Absorption spectra showing the reversibility of dioxygen binding to Co_4L1 . Bubbling O_2 into an acetone solution of Co_4L1 (0.02 mM) produces 1 (in high yield) at −50 °C (orange, solid line). Increasing the temperature up to 25 °C produces the blue solid spectrum. After recooling to −50 °C the orange solid spectrum can be regenerated. Bottom: Resonance Raman spectra of 1-¹⁶O (red trace), 1-¹⁸O (black trace) with 514 nm laser excitation in acetone-d₆ at −40 °C. Solvent bands are marked by "blue color".

tions in comparison with $Co₄Li$ (rise of the pre-edge – see inset of Fig. 2a, lower steepness of the edge rise, formation of a shoulder in the edge) indicating a slightly lower coordination symmetry and coordination number in 1 (Fig. 2, Table 1). The EXAFS of 1 reveals a 5 coordinate geometry at Co with 4 short Co–N/O distances at 1.91 Å, and one long Co– N/O distance at 2.02 Å. The average Co–O/N distances of 2.17 Å for $Co₄Li$ and 1.93 Å for 1, is a consequence of the different oxidation states and the lower steric requirements of five versus six ligands attached to the Co-atom. Notably, the 470 nm absorption band, ν (O–O) of 862 cm⁻¹ and ν (Co–O) of 595 cm⁻¹ and the Co–O₂ distance of 1.91 Å of 1 matches the spectroscopic properties of the previously reported end-on μ -1,2-peroxo-dicobalt(III) complexes.²¹ DFT calculations also support an $S = 0$ end-on μ -1,2-peroxo-dicobalt(m) assignment of 1; the calculated metrical parameters very well match with the experiments.

Reactivity of 1 with protons and electrons

1 exhibited different reactivities with proton and electron donors depending on the reaction temperature. At −50 °C no reaction of 1 with Fc* was observed in the absence of TfOH.

Fig. 7 Top: Changes in the absorption band associated with the reaction of 1 (0.015 mM) with TFA (3 mM) and Fc* (0.15 mM) at -50 °C; Inset: The pseudo-first order decay of the absorption band at 464 nm as a function of time (left) and the linear dependence (right) of the pseudofirst order rate constants (k_{obs}) on Fc* concentrations (0.13–0.38 mM) that led us to determine the second order rate constant value, k'_2 . Bottom: Changes in the absorption band associated with the reaction of 1 (0.015 mM) with TFA (0.75 mM) at +25 °C; inset: the pseudo-first order decay of the absorption band at 464 nm as a function of time (left) and the linear dependence (right) of the pseudo-first order rate constants (k_{obs}) on TFA concentrations (0.75–2.7 mM) led us to determine the second order rate constant value, k'_2 .

Similarly, no reaction of 1 with TfOH was observed in the absence of Fc*. However, in the presence of both TFA and Fc* 1 underwent fast decay, presumably by a proton coupled electron transfer (PCET) mechanism to form water as the major product (Fig. 7 top). At 25 \degree C in the absence of TFA, no reduction of 1 by Fc* was observed, very similar to our findings at −50 °C. However, in presence of TFA, even in the absence of Fc*, fast decay of 1 was observed (Fig. 7 bottom), with the release of H_2O_2 by a proton transfer (PT) mechanism.

The temperature dependence of the PT and PCET processes will be the controlling factor in determining the temperature dependence of the $4e^-/4H^+$ vs. $2e^-/2H^+$ reductions of dioxygen mediated by 1. We therefore compared the temperature-dependence of the PCET and PT processes of 1 at various temperatures (Fig. 7, 8 and S9†). PCET rates were determined at -50 °C, -40 °C, -30 °C and -20 °C under the condition [1] \ll [Fc*] \ll [TFA] to ensure pseudo first-order kinetics (Fig. 7) top; at these temperatures PT rates are negligible). Similarly, PT rates were determined at 20 °C, 22 °C, 25 °C, 30 °C and 32 °C under the condition $[1] \ll [TFA]$ (Fig. 7 bottom). PT is found to vary with temperature at a much more drastic rate relative to that of PCET, and it becomes the predominant mechanism for the reduction of 1 at temperatures >11 °C (Fig. 8a).

Fig. 8 Comparison of the temperature-dependence of the PCET vs. PT rate constants for (a) 1 and (b) 2.

Conclusions

In our previous study 8 we reported the synthesis and characterization of a hexanuclear cobalt complex $Co₆L2$ involving a nonheme ligand system, L2, supported on a $Sn₆O₆$ stannoxane core (Fig. 1: bottom), whose cobalt complex acts as a unique catalyst for dioxygen reduction, whose selectivity can be changed from a preferential $4e^-/4H^+$ dioxygen-reduction (to water) to a $2e^-/2H^+$ process (to hydrogen peroxide) only by increasing the temperature from −50 to 25 °C. Herein, we report the synthesis and characterization of a tetranuclear Co4L1 complex, supported on the stannoxane core, and compare its dioxygen reduction ability with that of $Co₆L2$ (Fig. 8). The temperature dependence of the product selectivity of the catalytic dioxygen reduction is still observed in $Co₄Li$; however, some subtle differences are noted relative to $Co₆ L2$, which can be attributed to the different nuclearity and Co–Co distances in the two cases (Fig. 1).

In L2 all the six plausible metal binding sites are equidistant from each other at 11.365 Å. This is in contrast to L1, where only two of the four metal binding sites are in close proximity to each other. The more symmetric nature of L2 ensures an efficient cooperative dioxygen binding in $Co₆L2$ relative to $Co₄LI$, which results in the lower stability of 1 compared to that of the corresponding ${[\text{L2}(\text{Co}^{\text{III}}(O_2) \text{Co}^{\text{III}})]_3]}^{12+}$ complex 2 that is formed upon dioxygen activation of $Co₆ L2$.

The faster self-decay rate (1 × 10⁻⁴ s⁻¹ for 2 *vs*. 2 × 10⁻³ for 1 at 25 °C), as well as the 16 cm−¹ downshift in the Co–O vibration energy (ν (Co–O) for 1 is 595 cm⁻¹ and 611 cm⁻¹ for 2)⁸ in 1 relative to 2, is consistent with the lower stability of 1. Accordingly, as previously observed, the high enthalpic stability of 2 makes its formation at −50 °C highly favored that leads to the complete oxygenation of $Co₆ L2$. Complex 2 then undergoes O-O bond cleavage via a PCET mechanism to yield water as the sole product under catalytic turnover conditions. The rate constant of the reaction was found to be independent of the O₂ concentration; the kinetic equation at −50 °C for $Co₆L2$ is

$$
d[Fc*]/dt = k_{obs}[Co_6L2]
$$

$$
k_{obs} = k_{cat}[Fc*][TFA]
$$

where " k_{cat} " is the third-order rate constant for the catalytic 4e⁻-reduction of O₂ by Fc^{*} at −50 °C and k_{obs} is the pseudo first-order rate constant. In contrast, an equilibrium binding of O2 occurs for Co4L1, even at −50 °C, so that the rate of the catalytic reaction shows a linear dependence on the $O₂$ concentration (Fig. $S10\dagger$). The rate equation for Co₄L1 is

$$
d[\text{Fc*}]/dt = k_{\text{obs}}[\text{Co}_4\text{L1}]
$$

$$
k_{\text{obs}} = k_{\text{cat}}[\text{Fc*}][\text{TFA}][\text{O}_2]
$$

where " k_{cat} " is the fourth-order rate constant for the catalytic 4e⁻-reduction of O₂ by Fc^{*} at −50 °C and k_{obs} is the pseudo first-order rate constant. Furthermore, $Co₄LI$ catalysed $O₂$ reduction yields 15–30% H₂O₂ at –50 °C, in contrast to $Co₆L2$ for which no H_2O_2 production could be detected at this temperature. However the rate constant of the two-electron $O₂$ reduction at +25 °C is a fourth-order process for both $Co₆L2$ and $Co₄LI$ (Fig. S11†).

The constraints imposed by the stannoxane core ensure entropic instability of both 1 and 2. This is mainly because of the large reduction in the Co–Co distances that is associated with their formation. Although experimental determination of the Co–Co distances in $Co₄LI$, $Co₆LI$, 1 and 2 was not possible, approximate shortening of ∼2.4 Å (from a distance of 6.82 Å in L1 to the DFT calculated distance of 4.48 Å in 1) and ∼7 Å (from a distance of 11.36 \AA in L2 to the DFT calculated distance of 4.48 \AA in 2) can be predicted for dioxygen binding at $Co₄Li$ and $Co₆Li$ complexes, respectively. This would impose a large strain on the μ -1,2-peroxo-dicobalt (m) cores in 1 and 2, which would attribute to their instability at higher temperatures upon protonation leading to the formation of H_2O_2 as the major product. Thus for both $Co₄Li$ and $Co₆Li$, an equilibrium binding of O_2 will take place at 25–30 °C, such that only a small portion of $Co₄Li$ and $Co₆Li$ will be converted to 1 and 2, respectively. This would also explain the experimentally observed direct correlation of the reaction rates to oxygen concentration at 25–30 °C in both cases.

In summary, the $Co₄LI$ complex like the previously reported $Co₆ L2$ complex is a unique catalyst for dioxygen-reduction reaction, whereby the product selectivity can be changed from

a predominant 4e/4H⁺ reduction process (to water) at -50 °C to a $2e^-/2H^+$ process at 25–30 °C. μ-1,2-peroxo-dicobalt(m) complexes 1 and 2 are proposed as plausible reactive intermediates, which are reduced to $H₂O$ by a PCET mechanism at -50 °C, or to H₂O₂ by a proton transfer mechanism at 25–30 \degree C. For both 1 and 2, the PT rates are found to vary drastically with temperature relative to the PCET rates, and PT becomes the predominant mechanism at 11 °C for 1 and at 19.5 °C for 2. The ∼10 °C reduction in the transition temperature for 1 can be attributed to its reduced stability relative to 2, as also evident from the faster self-decay rate and lower ν (Co–O) vibration energy in 1 relative to 2. This study, therefore, underlines the importance of subtle electronic and steric changes in the reactivity of the biologically relevant metal–dioxygen intermediates, and how they can control the $2e^-/2H^+$ vs. $4e^-/4H^+$ product selectivity in catalytic dioxygen reductions. Paper Maccess Articles. Complete the published on 2021, Characteristic Complex Articles. Ar

Conflicts of interest

There are no conflicts to declare.

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