

LJMU Research Online

Loughrey, JJ, Patmore, NJ, Baldansuren, A, Fielding, AJ, McInnes, EJL, Hardie, MJ, Sproules, S and Halcrow, MA

Platinum(ii) complexes of mixed-valent radicals derived from cyclotricatechylene, a macrocyclic tris-dioxolene

http://researchonline.ljmu.ac.uk/id/eprint/8151/

Article

Citation (please note it is advisable to refer to the publisher's version if you intend to cite from this work)

Loughrey, JJ, Patmore, NJ, Baldansuren, A, Fielding, AJ, McInnes, EJL, Hardie, MJ, Sproules, S and Halcrow, MA (2015) Platinum(ii) complexes of mixed-valent radicals derived from cyclotricatechylene, a macrocyclic trisdioxolene. Chemical Science. 6 (12). pp. 6935-6948. ISSN 2041-6520

LJMU has developed LJMU Research Online for users to access the research output of the University more effectively. Copyright © and Moral Rights for the papers on this site are retained by the individual authors and/or other copyright owners. Users may download and/or print one copy of any article(s) in LJMU Research Online to facilitate their private study or for non-commercial research. You may not engage in further distribution of the material or use it for any profit-making activities or any commercial gain.

The version presented here may differ from the published version or from the version of the record. Please see the repository URL above for details on accessing the published version and note that access may require a subscription.

For more information please contact researchonline@ljmu.ac.uk

http://researchonline.ljmu.ac.uk/

Chemical Science

EDGE ARTICLE



Cite this: Chem. Sci., 2015, 6, 6935

Received 29th July 2015 Accepted 18th August 2015 DOI: 10.1039/c5sc02776d

www.rsc.org/chemicalscience

Introduction

Metal/dioxolene complexes exhibit complicated redox chemistry and spectroscopy, reflecting the accessible catecholate ('cat'), semiquinone ('sq') and quinone ('q') oxidation states of the dioxolene group.¹⁻⁴ Charge transfer between a metal ion and dioxolene is often facile,¹ leading to intense visible/NIR absorptions that could be of use in solar energy applications.⁵ Intramolecular metal \rightleftharpoons dioxolene electron transfer can lead to valence tautomerism equilibria,¹ which may be accompanied by a metal ion spin-state change.⁶ Alternatively, some homoleptic metal/dioxolene complexes can be obtained in mixed-valent cat/

^bDepartment of Chemistry, University of Sheffield, Brook Hill, Sheffield S3 7HF, UK ^cSchool of Chemistry and Photon Science Institute, University of Manchester, Oxford Road, Manchester M13 9PL, UK

Platinum(II) complexes of mixed-valent radicals derived from cyclotricatechylene, a macrocyclic tris-dioxolene†

Jonathan J. Loughrey,^a Nathan J. Patmore,^{‡b} Amgalanbaatar Baldansuren,^c Alistair J. Fielding,^c Eric J. L. McInnes,^c Michaele J. Hardie,^a Stephen Sproules^{*d} and Malcolm A. Halcrow^{*a}

Three complexes of cyclotricatechylene (H₆ctc), [{PtL}₃(μ_3 -ctc)], have been synthesised: (L = 1,2-bis(diphenylphosphino)benzene {dppb}, **1**; L = 1,2-bis(diphenylphosphino)ethane {dppe}, **2**; L = 4,4'-bis(*tert*-butyl)-2,2'-bipyridyl {^tBu₂bipy}, **3**). The complexes show three low-potential, chemically reversible voltammetric oxidations separated by *ca*. 180 mV, corresponding to stepwise oxidation of the [ctc]⁶⁻ catecholato rings to the semiquinonate level. The redox series [**1**]^{0/1+/2+/3+} and [**3**]^{0/1+/2+/3+} have been characterised by UV/vis/NIR spectroelectrochemistry. The mono- and di-cations have class II mixed valent character, with reduced radical delocalisation compared to an analogous bis-dioxolene system. The SOMO composition of [**1**']⁺ and [**3**']⁺ has been delineated by cw EPR, ENDOR and HYSCORE spectroscopies, with the aid of two monometallic model compounds [PtL(DBsq')]⁺ (DBsqH = 3,5-bis(*tert*-butyl)-1,2-benzosemiquinone; L = dppe or ^tBu₂bipy). DF and time-dependent DF calculations confirm these interpretations, and demonstrate changes to spin-delocalisation in the ctc macrocycle as it is sequentially oxidised.

sq or sq/q ligand oxidation states, showing strong coupling and delocalisation between the ligand redox sites leading to strong NIR absorptions as before.⁷ Finally, these considerations can lead to dioxolene-related ligands and substrates acting as electron reservoirs during synthetic and biological catalysis.⁸

Complexes of dinucleating and polynucleating dioxolenes have potential for even more complicated electronic structures and redox, involving electron transfer between multiple metal and ligand sites. Most studied are 2,5-hydroxy-1,4-benzoquinonate complexes, where two metal ions chelate to one ligand redox centre.4 More unusual, are complexes of ligands bearing two or more dioxolene rings separated by a spacer,^{2,3} which have potential for ligand-based mixed valency.9 Several bis- and trisdioxolenes have been investigated in this regard, examples of which are shown in Scheme 1.10-26 However, since early interest in these compounds was focussed on the magnetic properties of molecular poly-sq radicals,3 only a handful of these complexed poly-dioxolenes have been spectroscopically characterised in their mixed-valent cat/sq forms.10-15,23 These range from being fully delocalised ([biscat⁻]^{3-2,10-13} and [triscat⁻]⁵⁻/[triscat⁻]⁴⁻²³) to being valence-localised, with electron hopping between the oxidised and unoxidised dioxolene rings occurring near the EPR timescale ([spiro']³⁻).^{2,14}

Seventeen years ago, Bohle and Stasko communicated two compounds of type [{PtL}₃(μ_3 -ctc)], where H₆ctc is cyclotricatechylene (Scheme 1) and L is a diphosphine.²⁷ Both compounds show three low-potential oxidations by cyclic



View Article Online

View Journal | View Issue

[&]quot;School of Chemistry, University of Leeds, Woodhouse Lane, Leeds LS2 9JT, UK. E-mail: m.a.halcrow@leeds.ac.uk

^dWestCHEM, School of Chemistry, University of Glasgow, Glasgow G12 8QQ, UK. E-mail: stephen.sproules@glasgow.ac.uk

[†] Electronic supplementary information (ESI) available: Experimental procedures for the physical and computational characterisation of the compounds in this work; crystallographic figures, and tables of crystallographic parameters; spectroelectrochemical data for [4[•]]⁺ and [5[•]]⁺; cw and pulsed EPR spectra; additional MO manifolds, orbital and spin density plots. CCDC 1037214 and 1037215. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/c5sc02776d

[‡] Current address: Department of Chemical and Biological Sciences, University of Huddersfield, Huddersfield HD1 3DH, UK.



Scheme 1 The complexes studied in this work, and other bis- and trisdioxolene complexes referred to in the article.

voltammetry, indicating that the $[ctc]^{6-}$ catecholate rings are oxidised sequentially from the cat to sq levels. No characterisation of those unusual radical species was undertaken, however. More recently cage complexes²⁸ and supramolecular assemblies²⁹ based on $[ctc]^{6-}$ have been crystallographically characterised, although their redox properties also remain to be explored.

Apart from the potential of ctc complexes to act as redoxactive supramolecular hosts, $[ctc']^{5-}$ and $[ctc'']^{4-}$ radicals also represent a unique example of multi-center mixed valency^{9,30} in a cyclophane-type scaffold.³¹ Since some of us have a longstanding interest in the host:guest chemistry of receptors derived from ctc,³² we decided to re-investigate the redox activity of the [{PtL}₃(μ_3 -ctc)] system. To shed further light on the trimetallic compounds, we have also quantified the electronic structures of mononuclear platinum/semiquinonate complexes by EPR and DFT methods.

Results and discussion

Reaction of H_6 ctc with metal ion reagents is complicated by its decomposition in the presence of base, and the moisture- and air-sensitivity of its complex products. Following extensive experiments with different transition ions, Pt(n) complexes of $[ctc]^{6-}$ were found to be the most tractable. Thus $[{PtL}_3(\mu_3-ctc)]$ (L = 1,2-bis(diphenylphosphino)benzene {dppb}, 1; L = 1,2-bis(diphenylphosphino)ethane {dppe}, 2; L = 4,4'-bis(tert-

butyl)-2,2'-bipyridyl { ${}^{t}Bu_{2}bipy$ }, 3) were obtained by reaction of H₆ctc with the appropriate [PtCl₂L] precursor in the presence of K₂CO₃, in a *N*,*N*-dimethylacetamide/methanol solvent mixture. Compound **1** is from Bohle and Stasko's initial study,²⁷ but **2** and **3** have not been reported before. Compound **2** is less soluble than **1** in most organic solvents, which hampered its solution characterisation. The known mononuclear complexes [Pt(dppe)(DBcat)] (**4**) and [Pt(${}^{t}Bu_{2}bipy$)(DBcat)] (**5**, Scheme **1**; DBcatH₂ = 3,5-di-*tert*-butylcatechol) were also prepared,¹³ as an aid to interpretating the spectroscopic data from **1–3**.

Electrospray mass spectra of 1–3 exhibit strong mono-, diand tri-cationic peaks associated with intact $[{PtL}_3(\mu_3-ctc)]^{z+}$ $(L = dppb, dppe \text{ or } {}^t\text{Bu}_2\text{bipy}; z = 1, 2 \text{ or } 3)$ ions. Only moderate fragmentation through loss of L or ${PtL}^{2+}$ fragments was observed (ESI†). The ¹H NMR spectra of 1–3 show the anticipated distribution of peaks, with a characteristic pair of resonances from the ditopic CH₂ groups in the $[ctc]^{6-}$ macrocycle. The spectra are sharp when obtained under an inert atmosphere, but are broadened when run in air with reduced integrals for the $[ctc]^{6-}$ ligand peaks. X-band EPR of the air-exposed solutions showed a weak resonance at g ≈ 2.03 at 150 K, implying the NMR peak-broadening reflects partial aerobic oxidation of the coordinated $[ctc]^{6-}$.

Crystallisation of 2 from N,N-dimethylacetamide (dma)/ methanol yielded a mixture of two solvates. The complex molecule in $2 \cdot H_2O \cdot 8$ dma has crystallographic C_3 symmetry, and is disordered about a mirror plane. Its molecular structure resembles that of 1 (Fig. 1).27 The bond lengths and angles within the metal coordination sphere are typical for a platinum(II)/catecholate/phosphine complex,33 while the dimensions of the unique dioxolene ring provide no evidence for oxidation of the [ctc]⁶⁻ ligand (ESI[†]).³⁴ Like all ctc derivatives,³² 2 has a bowl-shaped cavity at its centre, of approximate dimensions 6.8 Å rim diameter imes 9.5 Å depth. Precedent implies this cavity should be occupied by solvent;25,35,36 although this was not apparent in the Fourier map, a SQUEEZE analysis³⁷ implied the presence of two unresolved molecules of dma in this region of the lattice. The dihedral angle between the catecholate groups in the molecule is $71.2(5)^{\circ}$, which is again similar to the published solvate of 1 (70.6-73.2°).27

The complex molecule in the other solvate, $2 \cdot 2H_2$ -O·1.3dma·0.5MeOH, lies on a general crystallographic site. The molecular structure of 2 in this crystal is very similar to the first solvate, apart from differences in the orientations of some phenyl substituents. In particular, the dimensions of the ctc dioxolene units are again consistent with the catecholate oxidation level (ESI[†]).³⁴ The depth of the molecular cavity in this solvate is ca. 9.2 Å, but its rim dimensions cannot be uniquely defined because of disorder in the dppe phenyl groups. The cavity contains a disordered dma molecule, which is incompletely resolved in the model. The intramolecular dihedral angles between neighbouring catecholate groups in this solvate span a wider range at 62.9(3)-76.2(3)°, implying some flexibility in the $[ctc]^{6-}$ macrocycle. The average of these angles, $69.1(5)^{\circ}$, is similar to the value from the first crystal however.



Fig. 1 Two views of the complex molecule in $2 \cdot H_2O \cdot 8$ dma. Displacement ellipsoids are at the 50% probability level, except for the dppe C atoms in the lower view which are de-emphasised, and H atoms are omitted for clarity. Colour code: C, white; O, red; P, green; Pt, cyan. Symmetry codes: (i) 1 - y, 1 + x - y, z; (ii) -x + y, 1 - x, z.

Electrochemistry and spectroelectrochemistry

The cyclic voltammogram (CV) and differential pulse voltammogram (DPV) of **1** in CH₂Cl₂/0.5 M [^{*n*}Bu₄N]BF₄ at 298 K confirmed Bohle and Stasko's original report, in showing the [ctc]^{6-/5-/4-/3-} redox series (cat/cat/cat \rightleftharpoons cat/cat/sq \rightleftharpoons cat/sq/ sq \rightleftharpoons sq/sq/sq, Scheme 2; Table 1, Fig. 2).²⁷ These oxidations in 2 occur at similar potentials as in **1**, although **2** also exhibits a fourth process near +0.2 V that is not shown by **1**. The CV and DPV of 3 resemble those of **2**, albeit with slightly more positive potentials for the first two oxidations. The third and fourth oxidation processes in **2** and **3** lie at almost identical potentials and have *ca*. half the intensity of the first two processes by DPV (Fig. 2). Since its relative intensity is reduced at higher scan

rates, we tentatively attribute the fourth oxidation to a daughter arising from deposition of $[2^{\circ}]^{2+}$ and $[3^{\circ}]^{2+}$ at the electrode surface. That would explain why such a peak is not observed for 1, which is more soluble than 2 in the base electrolyte medium. The separation between the first three oxidation potentials in 1 and 2, $\Delta E_{1/2} = 0.18-0.22 \text{ V} (K_c = 1-5 \times 10^3)$,³⁰ is larger than in 3 (0.12 V, $K_c = 1 \times 10^2$) and implies moderate-to-weak communication between the dioxolene rings in these complexes. In comparison, complexes of $[\text{thea}]^{4-}$ and $[\text{spiro}]^{4-}$, whose dioxolene rings are also linked by methylene spacers (Scheme 1), show $\Delta E_{1/2} (K_c) = 0.25 \text{ V} (2 \times 10^4)$ and *ca.* 0.15 V (3 × 10²) respectively.^{2,14,15}

In addition to these low-potential oxidations, **1**–3 exhibit an envelope of irreversible oxidations between +0.6 and +1.1 V, attributable to the $[ctc]^{3-/2-/1-/0}$ (sq/sq/sq \Rightarrow sq/sq/q \Rightarrow sq/q/q $q \Rightarrow q/q/q$) sequence, while 3 also exhibits a chemically reversible three-electron reduction at -1.8 V (Fig. 2). This arises from reduction of the ${}^{t}Bu_{2}$ bipy co-ligands, ${}^{13,38-43}$ and is split into at least two components in the DPV trace.

The oxidation processes of **1** and **3** were monitored spectroelectrochemically in CH₂Cl₂/0.5 M [N^{*n*}Bu₄]BF₄ at 253 K (Fig. 3). Three of the oxidations have no potential isosbestic points, but the other oxidations exhibit minor decomposition of the compounds during the experiments. Their small potential separation (ΔE) also made it difficult to poise the samples cleanly at the intermediate oxidation levels. Despite these issues, however, the following trends can be noted.

Oxidation of **1** to $[\mathbf{1}^{\cdot}]^{+}$ leads to the appearance of a broad near-IR absorption centred at 7.9 × 10³ cm⁻¹, assignable to an intervalence charge transfer (ICVT) band (Fig. 3). Generation of the $[\mathbf{1}^{\cdot}]^{+} \rightarrow [\mathbf{1}^{\cdot \cdot}]^{2+}$ oxidation causes the IVCT band to shift to $\nu_{\text{max}} = 7.1 \times 10^3 \text{ cm}^{-1}$, while increasing slightly in intensity. The $\mathbf{3} \rightarrow [\mathbf{3}^{\cdot}]^{+} \rightarrow [\mathbf{3}^{\cdot \cdot}]^{2+}$ sequence also generates an IVCT absorption, which also red-shifts by 800 cm⁻¹ during the second oxidation but is stronger and lower energy than for $[\mathbf{1}^{\cdot}]^{+}$ - $[\mathbf{1}^{\cdot \cdot}]^{2+}$ (Fig. 3; Table 2). Generation of the third oxidation of both

Table 1 Electrochemical data for the complexes ^a					
	sq/cat	$(E_{1/2}/V)$		q/sq (irr ^b , Ep_a/V)	$[^{t}Bu_{2}bipy]^{0/-}(E_{1/2}/V)$
1 ^c 2 3	$-0.30 \\ -0.35 \\ -0.17$	$-0.12 \\ -0.17 \\ -0.05$	+0.10 +0.04, +0.23 +0.07, +0.20	+0.62 +0.61 ^{d} , +0.79 +1.12	- -1.78, -1.86 ^d

^{*a*} Potentials referenced to Fc^{+/0}. ^{*b*} irr = irreversible. ^{*c*} cat/sq $E_{1/2}$ = +0.04, 0.24 and 0.49 V ν s. NHE from ref. 27. ^{*d*} Shoulder.



Scheme 2 Ligand-based redox series exhibited by $[{PtL}_3(\mu_3-ctc)]$ (L = neutral bidentate ligand)



Fig. 2 Cyclic and differential pulse voltammograms of 1 (top) and 3 (bottom) at 298 K, in $CH_2Cl_2/0.5$ M [NⁿBu₄]BF₄ and scan rate 100 mV s⁻¹. Only the cat/sq oxidation window is shown in the CVs, while the DPVs scan the full range of the solvent window.

compounds almost completely quenches the near-IR absorption. Assignment of the other spectral changes was aided by comparison with the mono-metallic complexes (see below). A weak absorption at 11.1 × 10³ cm⁻¹ in $[3^{\cdot}]^{2+}$ and $[3^{\cdot}]^{3+}$ may be a Pt(d σ) \rightarrow sq MLCT transition, and a strong peak at 20–24 × 10³ cm⁻¹ in all the radical spectra is assigned to a Pt(d π) \rightarrow sq MLCT band.⁴⁴ A transition near 18 × 10³ cm⁻¹, which is shown by 3–[3[•]]²⁺ but collapses upon the third oxidation, is a cat \rightarrow bipy inter-ligand CT band,³⁹ while a peak at 27.2 × 10³ cm⁻¹ in 3 that blue-shifts upon oxidation is a Pt \rightarrow bipy MLCT.^{41,42} More detailed assignment of these peaks is described below.

Despite the non-isosbesticity of the electrochemical oxidations, re-reduction of the fully oxidised solutions at -0.4 V regenerated the neutral precursor spectra with transient appearance of the IVCT peaks described above. That implies 1 and 3 spontaneously reassemble from their regenerated constituents.

Allowing for overlapping high-energy absorptions,⁴⁵ the width at half height $(\Delta \nu_{1/2})$ of the IVCT bands in [3']⁺ and [3'']²⁺ can be estimated at 4950 and 3550 cm⁻¹, respectively. In comparison, the values predicted for a class II mixed-valent system⁴⁶ with these IVCT energies are $\Delta \nu_{1/2} \approx 3800 \text{ cm}^{-1}$ for [3']⁺ and 3500 cm⁻¹ for [3'']²⁺ (eqn (1)).^{9,30}

$$\Delta \nu_{1/2} = (2310E)^{1/2} \tag{1}$$

 $\Delta v_{1/2}$ for $[1^{\cdot}]^+$ and $[1^{\cdot}]^{2+}$ cannot be measured directly but is greater than 5000 cm⁻¹, which is also larger than predicted by

eqn (1) (4250 cm⁻¹ for $[1']^+$ and 4050 cm⁻¹ for $[1'']^{2+}$). Assuming they lie in the class II limit, the IVCT v_{max} for the mixed-valent radicals corresponds to their inter-valence reorganisation energy (λ); that is, $5.4 \le \lambda \le 7.9 \times 10^3 \text{ cm}^{-1}$ (Table 2).³⁰ Those are at the low end of the typical range of λ values for class II organic radicals.9 UV/vis/NIR spectra in different solvents to define λ in more detail were not undertaken,^{9,30} because of the limited stability of the oxidised complexes. Nonethe-less, all these mixed-valent radicals adopt electronic structures whose dioxolene rings are partly valence-localised, although [3^{••}]²⁺ may exhibit a reduced barrier to electronhopping since its $\Delta v_{1/2}$ value is closer to the classical class II description.9,30,46 That contrasts with the cat/sq radicals $[{PtL}_2(\text{thea'})]^+$ (L = dppb or dppe; Scheme 1) which exhibit narrower IVCT bands that are more characteristic of the class III formalism.15

Mononuclear 4 and 5 undergo a reversible cat \rightarrow sq oxidation near $-0.2 \text{ V} \nu s. [\text{Fc}]^{+/0}$, which was monitored by chemical redox titration with [Fc]PF₆. The changes in the UV/vis spectra of 4 and 5 upon oxidation resemble the first oxidations of the trinuclear compounds (Fig. S9;† Table 2), in depletion of cat \rightarrow L LLCT absorptions (L = dppe or ${}^{t}\text{Bu}_2$ bipy) and the growth of new Pt(d σ) \rightarrow sq and Pt(d π) \rightarrow sq MLCT peaks near 12 and 23 × 10³ cm⁻¹, respectively.^{13,44} The energy difference between the Pt(d π) \rightarrow sq band in [4⁻]⁺ compared to [5⁻]⁺ ($\Delta E_{\text{MLCT}} = 2.4 \times 10^3 \text{ cm}^{-1}$) is identical to the difference between the same transition in [1⁻]⁺ and [3⁻]⁺ (2.4 × 10³ cm⁻¹), and [1⁻]²⁺ and **Edge Article**



Fig. 3 The first three oxidations of 1 (left) and 3 (right) at 253 K, in $CH_2Cl_2/0.1$ M NBu_4BF_4 , monitored by UV/vis/NIR spectroscopy using an optically transparent electrode. The spectra of the pure starting material and product spectra are highlighted in black while the intermediate spectra are paler. Discontinuities near 12.4 and 28.7×10^3 cm⁻¹ are artifacts from grating changes in the spectrometer.

 $[3^{\cdot}]^{2+}$ (2.3 × 10³ cm⁻¹; Table 2). That implies the Pt/sq interactions in the ctc-based radicals and the mono-nuclear semiquinonate complexes are similar in character.

EPR spectroscopy of monometallic complexes

Fluid solution S-band (3.9 GHz) and X-band (9.4 GHz) EPR spectra of [4[•]]⁺ and [5[•]]⁺ (Scheme 1) in dichloromethane/THF solution at 210–230 K exhibit one central line from naturally occurring metal isotopes with no nuclear spin, flanked by ¹⁹⁵Pt

(I = 1/2, 33.8% abundant) hyperfine satellites. The isotropic ¹⁹⁵Pt coupling is much larger in [5[•]]⁺ than in [4[•]]⁺ (96.4 vs. 36.6 MHz, respectively). In [4[•]]⁺, there is further hyperfine to one ¹H and two equivalent ³¹P nuclei ($a^{H} = 9.5, a^{P} = 7.8$ MHz; Fig. S10,[†] Table 3). For [5[•]]⁺, there is splitting to a single ¹H nucleus, with $a^{H} = 9.4$ MHz (Fig. 4, Table 3). The $g \approx 2.002$ value and hyperfine analysis (see below) for both radicals identify the singly occupied molecular orbital (SOMO) as largely localised in the dioxolene ligand π -system, but with some contribution from metal-based orbitals. The strongly coupled

8

This article is licensed under a Creative Commons Attribution 3.0 Unported Licence.

Open Access Article. Published on 20 August 2015. Downloaded on 27/02/2018 08:37:02.

Table 2 Electronic absorption data for the complexes

	$\nu_{\rm max}/10^3 {\rm ~cm^{-1}} \left(\varepsilon_{\rm max}/10^3 {\rm ~M^{-1}} {\rm ~cm^{-1}} \right)$
1	13.9 (0.1), 15.7 (0.1), 17.7 (sh), 26.3 (2.8)
[1 [.]] ⁺	7.9 (0.5), 14.4 (0.4), 15.9 (sh), 24.7 (sh)
[1] ²⁺	7.1 (0.7), 14.3 (1.1), 15.7 (sh), 18.6 (sh), 23.5 (3.3)
[1 ^{···}] ³⁺	10.6 (sh), 14.3 (1.4), 18.6 (sh), 23.6 (3.9)
2	14.6 (0.5), 26.9 (sh)
3	14.1 (sh), 17.6 (11.6), 27.2 (6.4)
[3 [.]] ⁺	6.2 (2.4), 14.2 (9.0), 18.1 (11.9), 20.8 (11.1), 27.2 (sh)
[3 ⁻¹] ²⁺	5.4 (5.7), 11.0 (3.2), 14.2 (10.0), 18.2 (12.8), 21.1 (14.0)
[3 ^{···}] ³⁺	11.4 (5.0), 14.1 (11.7), 16.3 (sh), 21.3 (14.7)
4	28.2 (sh), 33.2 (6.6)
[4 '] ⁺	12.3 (0.1), 14.3 (0.2), 16.0 (0.2), 18.7 (0.5), 24.0 (3.4)
5	16.8 (4.0), 26.0 (1.7)
[5 [.]] ⁺	10.4 (0.1), 16.0 (sh), 17.0 (2.0), 20.5 (sh),
	21.6 (5.3), 25.3 (sh), 28.5 (sh), 29.7 (sh)

Table 3 Spin-Hamiltonian parameters (MHz) and Pt contribution to the SOMO for the radical complexes

		[1 [•]] ⁺	[3 [•]] ⁺	[4 [•]] ⁺	[5 [•]] ⁺
	g _{iso}	2.0019	2.0017	2.0022	2.0025
	g_{y}	2.0112	2.0406	2.0095	2.0403
	g_x	2.0088	2.0032	2.0062	2.0036
	g_z	1.9838	1.9507	1.9789	1.9509
¹⁹⁵ Pt	A _{iso}	$\approx -12^{a}$	-31	-37	-97
	A_{v}	-70	-100	-66	-111
	A_x	-71	-144	-78	-150
	A_z	$+33^{b}$	-35^{b}	$+33^{b}$	-32^{b}
${}^{1}\text{H}^{c}$	a_{iso}			9.5	9.4
	a_x			d	15
	a_{v}			d	2.6
	a_z			d	10.6
	$\langle a \rangle$				9.4
	γ^{e} (deg)				60
${}^{1}\text{H}^{f}$	$a_{x,y}$	1.4	2.0	d	0.9
	a_z	2.6	3.2		3.6
	$\langle a \rangle$	1.8	2.4		1.8
${}^{1}\mathrm{H}^{g}$	$a_{y,z}$			d	5
	a_r				7
³¹ P	a_{iso}	2.7^{b}		7.8	
	$a_{x,y}$	8.5		8.2	
	a ₂	8.0		7.6	
	$\langle a \rangle$	8.3		8.0	
^{14}N	a_{iso}		1.0		1.2
	K		0.36		0.38
	a^{2h}	0.001	0.033	0.009	0.028
	b^{2h}	0.071	0.044	0.069	0.055
	Total Pt	0.078	0.074	0.078	0.083
	DFT	0.043	0.050	0.040	0.074

^{*a*} Value in fluid solution is *ca.* 1/3 as large as the average value in frozen solution. See text for details. ^{*b*} From $A_{iso,x,y}$. ^{*c*} H4 of 3,5-DBsq. ^{*d*} Similar to those in [5']⁺. ^{*e*} Euler angle about *z* relating $\mathbf{a}_{\mathbf{H}}$ and \mathbf{g} frames. ^{*f*} H6 of DBsq or H3,6 of ctc (modelled as axial from HYSCORE, but orientation not well defined). ^{*g*} Unassigned. ^{*h*} Pt 5d_{yz} and 6p_z admixture to SOMO.

proton is assigned to *H*4 of the [DBsq⁻]⁻ ligand (HDBsq = 3,5di(*tert*-butyl)-1,2-benzosemiquinone),⁴⁷ since the *C*4,5 positions carry much greater spin density than *C*3,6 (see below).

EPR spectra of frozen solutions were recorded at S-, X- and Q-band (3.9, 9.4 and 33.7 GHz, respectively) to determine the **g**



Fig. 4 Fluid solution S-band EPR spectra of $[5^{-}]^+$ and $[3^{-}]^+$ in CH₂Cl₂/ THF at 230 K. Simulations (red) used the parameters in Table 3.

and ¹⁹⁵Pt **A** matrices. The spectra are near-axial for [4']⁺ but clearly rhombic for [5']⁺ (Fig. 5, S10 and S11[†]) and are dominated by ¹⁹⁵Pt hyperfine coupling, as is common for Pt(II) complexes of π -radical ligands.^{41,42,48-50} Following the literature analyses,^{41,42,48,51} the molecular *z*-axis is defined perpendicular to the plane of the molecule, and the *y*-axis as bisecting the dioxolene ligand bite angle (Fig. S12[†]) with the assignment of the principal *g*-values in Table 3. The ¹⁹⁵Pt A_x and A_y components are resolved, but the high-field A_z component is not. This was therefore derived from $A_z = 3A_{iso} - (A_x + A_y)$. This component has an estimated uncertainty of 12 MHz, but clearly has a different sign to $A_{iso,x,y}$ for [4']⁺. In contrast, all the hyperfine components in [5']⁺ have the same sign.

Approximating the molecular symmetry to C_{2v} , the SOMO has b_2 symmetry (being dominated by the [DBsq[•]]⁻ ligand; see below). In this case only the metal $5d_{yz}$ and $6p_z$ valence orbitals have the correct symmetry to admix.^{41,42,48,50,52,53} These contributions can be calculated from the ¹⁹⁵Pt hyperfine matrix *via* eqn (2)–(4),^{41,42} where: a^2 and b^2 are the $5d_{yz}$ and $6p_z$ admixtures



Fig. 5 Frozen solution X-band EPR spectra of $[1^{-}]^+$ and $[3^{-}]^+$ at 150 K, and $[4^{-}]^+$ and $[5^{-}]^+$ at 30 K in CH₂Cl₂/THF. Simulation parameters are given in Table 3.

to the SOMO, respectively; $P_{\rm d}$ and $P_{\rm p}$ are the electron nuclear dipolar coupling parameters for Pt 5d and 6p electrons; and $A_{\rm s}$ is the isotropic Fermi contact term.

$$A_x = A_s - 4/7P_d a^2 - 2/5P_p b^2$$
(2)

$$A_y = A_s + 2/7P_d a^2 - 2/5P_p b^2$$
(3)

$$A_z = A_s + 2/7P_d a^2 + 4/5P_p b^2 \tag{4}$$

Hence, A_x is expected to be the largest hyperfine component and A_z the smallest, as per the assignment in Table 3. Using $P_d = +1.65 \times 10^3$ and $P_p = +1.21 \times 10^3$ MHz,^{41,48} we get total Pt contribution to the SOMO ($a^2 + b^2$) of 8%, dominated by the 6p_z contribution.⁴¹ The larger isotropic hyperfine for [5⁻]⁺ *cf.* [4⁻]⁺ reflects an increase in the 5d_{yz} contribution from *ca.* 1 to 3%. Thus, the SOMO in both species has *ca.* 90% dioxolene character, which is consistent with DFT calculations (see below; Table 3).

Since there is no resolution of ligand hyperfine in the frozen solution spectra, orientation-selective Davies electron nuclear double resonance (ENDOR) and hyperfine sublevel correlation (HYSCORE) spectroscopies were undertaken to probe the spin distribution further. Detailed Q- and W-band ENDOR spectra have been reported for *para*-semiquinones,⁵⁴ but we are unaware of any comparable study for the *ortho* equivalents. Q-band Davies ENDOR spectra for [4[•]]⁺ and [5[•]]⁺ and have nearly identical profiles in the ¹H region (Fig. 6 and S15;[†] v_n (¹H) = 51.33 MHz at 1200 mT), so we only discuss the latter in detail here.



Fig. 6 Q-band Davies ENDOR spectra of [5']⁺ in CH₂Cl₂/THF solution at 20 K (black), and simulations (red) with the *g* and *a*^H parameters listed in Table 3. Static magnetic fields as shown, corresponding to the echo-detected field-swept spectrum in Fig. S14.† Experimental conditions: $\nu_{mw} = 33.710$ GHz, $t_{RF} = 16$ µs, $t_{inv} = 200$ ns, $t_{\pi/2} = 22$ ns, $\tau = 440$ ns.

The hyperfine couplings for the α -protons of the [DBsq[•]]⁻ ligands (ring positions H4 and H6; Fig. S12[†]) arise from polarisation of C 2pz spin density. The principal ¹H hyperfine components are expected to be parallel to the C $2p_z(a_z)$, along the C-H bond (a_y) , and orthogonal to these two axes $(a_x, with a_x)$ $> a_z > a_y$).⁵⁴ However, spin density on neighbouring carbons can distort this pattern through rotation of the latter two axes about z. The C $2p_7$ orbitals are parallel to g_7 : this corresponds to the high-field extreme of the EPR spectrum. Hence, spectra at this field (1235.3 mT; Fig. 6, top) are single crystal-like: the largest splitting observed at this field is $a_z = 10.6$ MHz. This must be due to H4, because the C4,5 positions carry much greater spin density than C3,6.47 The maximum splitting observed is ca. 15 MHz at 1192 mT, that is, in the molecular xy plane. This corresponds to a_x of H4. Simulations (Fig. 5) give $a_{x,y,z}H4 = 15$, 2.6, 10.6 MHz with $\gamma = 60^{\circ}$, where γ is an Euler rotation angle about z relating the hyperfine (a_{xyz}) and the molecular (g_{xyz}) frames. The average $\langle a \rangle$ agrees well with the observed A_{iso} value from fluid solution.

A further ¹H coupling of 5–7 MHz is observed at each orientation. This is too large to arise from the other α-proton H6 (very little spin density is carried by C6), while its near isotropic nature is also inconsistent with an α -proton. An alternative assignment might reflect hyperconjugation to the ^tBu group in the C5 position, oriented such that a proton folds back towards the ring π -system.⁵⁵ These couplings are consistent with X-band HYSCORE spectra (Fig. S16[†]). Linear regression analysis of HYSCORE spectra gives evidence of a further, weak coupling with isotropic and anisotropic tensor components of 1.8 and 0.9 MHz, respectively, which we assign as H6. The ENDOR simulations are improved by making this rhombic (Table 3), but there is some uncertainty over these parameters. The isotropic H4 and H6 values of $\langle a \rangle = 9.4$ and 1.8 MHz are a close match for those of [DBsq[•]]⁻ itself in fluid solution (9.5 and 1.8 MHz).⁴⁷ These parameters imply the spin density at C4,5 is ca. five times that at C3,6.

X-band HYSCORE measurements on [5⁻]⁺ (Fig. 7b) also reveal coupling to the ¹⁴N atoms of the bipy co-ligand. The ¹⁴N coupling is near-isotropic, with analysis of the double quantum transitions giving $a^{\rm N} \approx 1.0$ MHz and quadrupole constant K =0.36 MHz ($\eta \approx 0$) (Fig. S17†). A near-isotropic ³¹P coupling is observed for [4⁻]⁺ in both HYSCORE (Fig. 7a) and Q-band ENDOR (Fig. S15†), giving $\langle a^{\rm P} \rangle = 8.0$ MHz, in agreement with the observed $a_{\rm iso}$. The very small anisotropic components (*ca.* 0.2 MHz) to the ¹⁴N and ³¹P hyperfine matrices put a limit of *ca.* 0.1% spin density at these positions.

These results were used to calibrate EPR data from the more elaborate tris-dioxolene systems derived from 1–3.

EPR spectroscopy of $[{PtL}_3(\mu_3-ctc')]^+$

The trinuclear radicals $[1^{\cdot}]^{+}$ and $[3^{\cdot}]^{+}$ were generated by *in situ* oxidation of 1 and 3 by 1 equiv. [Fc]PF₆ at 220 K, inside a precooled EPR tube. S-band spectra of chilled (210 K) fluid solutions of $[1^{\cdot}]^{+}$ and $[3^{\cdot}]^{+}$ show partial hyperfine resolution (Fig. S18 and S19†). The spectrum of $[3^{\cdot}]^{+}$ (Fig. 4) shows a dominant hyperfine coupling to more than one ¹⁹⁵Pt nucleus: simulation



Fig. 7 X-band HYSCORE spectra showing ³¹P coupling in (a) [4']⁺ and (c) [1']⁺, and ¹⁴N coupling in (b) [5']⁺ and (d) [3']⁺, measured at the maximum of the ESE-detected EPR spectra at 20 K. The anti-diagonals are in red, marking single and double quantum (for ¹⁴N) transitions frequencies.

assuming coupling to three equivalent Pt nuclei with the appropriate statistical distribution gives A^{Pt}_{iso} of 31 MHz. This is very close to one third of the value for the related monometallic complex [5⁻]⁺ (Table 3) implying that, at this temperature, the radical is delocalised over the entire complex. The spectra of [1⁻]⁺ are consistent with this delocalisation: although the Pt satellites are not clearly resolved, simulations give an upper bound of A_{iso}^{Pt} < 15 MHz, much smaller than for the equivalent monomer [4[•]]⁺. There is partial resolution of superhyperfine coupling with $a_{iso} \approx 2.7$ MHz, which could not be unambiguously assigned. However, there is only one unique dioxolene α -proton environment in $[1^{\cdot}]^+$ and $[3^{\cdot}]^+$, whose C-atoms (equivalent to the C3,6 sites in $[4^{-}]^{+}$ and $[5^{-}]^{+}$) should carry little spin density. Hence, the superhyperfine in [1']⁺ should be dominated by the six equivalent ³¹P atoms, and the observed a_{iso} is close to one-third the value found for $[4^{\cdot}]^+$ (the spectra can be simulated with this model {Fig. S18[†]}; including ¹⁹⁵Pt coupling in these simulations gives a best fit with A_{iso}^{Pt} = 12 MHz, but this is not well defined). In summary, fluid solution spectra of [1']⁺ and [3']⁺ are consistent with a delocalised radical model (Table 3).

In contrast, frozen solution X-band spectra of $[1^{\cdot}]^{+}$ and $[3^{\cdot}]^{+}$ at 150 K are very similar to their monometallic counterparts, with similar **g** and **A**^{Pt} matrices to $[4^{\cdot}]^{+}$ and $[5^{\cdot}]^{+}$, respectively (Fig. 5 and Table 3). Hence, the radicals are localised in the frozen solutions of $[1^{\cdot}]^{+}$ and $[3^{\cdot}]^{+}$, giving very similar electronic structures to the monomers $[4^{\cdot}]^{+}$ and $[5^{\cdot}]^{+}$. This is confirmed by measurements of ligand hyperfines. ¹H HYSCORE spectra for $[1^{\cdot}]^{+}$ and $[3^{\cdot}]^{+}$ at 20 K are similar, and show only a weak coupling with $\langle a \rangle \approx 2$ MHz, similar to that of the weaker coupled of the two α -protons (*H*6) in $[4^{\cdot}]^{+}$ and $[5^{\cdot}]^{+}$ (Fig. S20 and S21;[†] Table 3).

Hence we assign this to the two α -protons (*H*3,6) of a single dioxolene ring of [ctc[•]]^{5–}. Convincingly, ³¹P and ¹⁴N HYSCORE give essentially identical couplings to those for [4[•]]⁺ and [5[•]]⁺ (Fig. 7). Therefore, electron hopping between the dioxolene groups of [ctc[•]]^{5–} is suspended upon freezing of the matrix, localizing the electron spin on a single Pt-dioxolene fragment at these temperatures. A similar quenching of delocalisation on freezing was also reported in the dimetallic [{PtL}₂(thea[•])]⁺ radical (Scheme 1).¹⁵

Calculations

Geometry optimised structures and electronic properties of the free $[DBsq']^-$ radical, $[1]^{0/1+/2+/3+}$, $[3']^+$, $[4']^+$ and $[5']^+$ were calculated by spin-unrestricted broken symmetry (BS) DFT calculations at the B3LYP-ZORA level. The ^{*t*}Bu substituents on the bipy in $[3']^+$ and $[5']^+$, and the phosphine Ph groups in $[1]^{0/1+/2+/3+}$ and $[4']^+$, were replaced with H atoms.

The optimised structure of $[DBsq^{\cdot}]^{-}$ shows the alternating long-short aromatic C–C bond lengths expected for a semiquinone (Fig. S22†), with the average C–C distance intermediate between those typical for closed-shell catecholates and quinones.^{56,57} Calculated structures of the complexed ligands in [4[•]]⁺ and [5[•]]⁺ are similar (Table S4†), with slightly longer average C–O (*ca.* 1.30 Å) and shorter average C–C(ring) (1.41 Å) distances than free [DBsq[•]]⁻ (1.26 and 1.43 Å, respectively).

Calculated Mulliken spin densities (Fig. 8a–c) confirm the ligand radical nature of the complexes. There is slightly greater localisation of the spin on the [DBsq[•]][–] ring, and less on the O atoms, in the complexes than in the free radical. The total ($5d_{yz}$ plus $6p_z$) Pt composition of the SOMO is calculated to be 7.4% in [5[•]]⁺ and 4.0% in [4[•]]⁺. This agrees well with the 8% calculated from the EPR parameters, with the higher value for [5[•]]⁺ matching the increase in $5d_{yz}$ character from EPR (Table 3). Hence, the co-ligand has a small influence on the nature of the SOMO and we can speculate that the π -donor orbitals of the bipy in [5[•]]⁺ destabilise the 5d manifold such that they are energetically closer to the [DBsq[•]][–] π orbitals.

The optimised structure of **1** is close to its crystallographic geometry (Table S7[†]).²⁷ The bond lengths within the μ_3 -ctc ligand are accurately reproduced but the Pt-P and Pt-O distances are over- and under-estimated by 0.017 and 0.015 Å, respectively. The bowl structure of the cavitand is slightly more open in the computed structure as measured by the longer Pt… Pt separation (10.42 Å, cf. crystallographic 9.85 Å) and a larger dihedral angle between the dioxolene plane and the molecular three-fold axis. This stems from the absence of a solvent molecule trapped within the cavitand.27 The two highest occupied molecular orbitals (HOMOs) are the 1A1 and 2E combinations (C_{3v} molecular symmetry) of the three dioxolene π orbitals (which are b_2 in C_{2v} local symmetry; Fig. 10). The next filled MOs are the $1A_2$ (HOMO-2) and 1E (HOMO-3) combinations from the dioxolene π orbital with a_2 symmetry. The lowest unoccupied orbitals (LUMOs) are combinations of dppb π^* orbitals in groupings of 2A₁ and 3E, and 2A₂ and 4E. The 5E and 3A2 unoccupied orbitals are the lowest with large Pt

character (\sim 34%), corresponding to combinations of the inplane 5d_{xy} orbital, and are at considerably higher energy.

Removal of an electron from the ctc ligand to form [1⁻]⁺ does not affect the topology of the molecule, but lowers its symmetry to $C_{\rm s}$ (Table S7[†]). This is evident in the ctc ligand geometry, where one dioxolene unit has significantly shorter C-O bond lengths (1.337 Å) compared with the other two (1.347 Å). All degenerate E-type MOs in 1, including the 2E HOMO, are split into A' and A'' components upon oxidation; this yields a 3A' SOMO and 3A'' HOMO-1 in $[1']^+$ (Fig. 9). The overall decrease in symmetry is relatively small, however, when considering the energy separation between previously degenerate pairs. The reduction in symmetry is more pronounced when the optimisation is performed in a highly polar solvent continuum. This effect is depicted in the Mulliken spin density plot for $[1']^+$, where the majority of the spin (+0.74) is found on one arm of the complex (Fig. 8d). The corresponding plot for [3[•]]⁺ was derived from a gas-phase optimisation, and here the spin density is more evenly distributed across the three arms of the complex (Fig. S29[†]).

Interestingly, the formation of a semiquinone and the subsequent attenuation of the π -donor strength substantially stabilises the dppb π^* (4A' and 4A'') and Pt 5d_{xy} (5A'') orbitals (LUMO, LUMO+1 and LUMO+2, respectively; Fig. S26†) of one arm of $[1^{\cdot}]^+$ relative to the other two, which remain near degenerate. The Pt content of the 4A' (10.3%) and 5A'' (40%) MOs is larger than for the corresponding orbitals in 1. The MO manifold for $[3^{\cdot}]^+$ is essentially identical, albeit with a smaller HOMO–LUMO energy gap reflecting the low-lying π^* orbital of bipy (Fig. S31†).

The Mulliken spin density analysis for the S = 1/2 monocations of 1, 3, 4 and 5 can be used to validate the ¹H hyperfine couplings measured by cw and pulsed EPR spectroscopy. The DFT-derived spin density on protons on the dioxolene unit are listed in Table 4, and show excellent agreement with the experimentally determined values according to the relationship $\rho_{\rm H} = a_{\rm iso}/1419.^{58}$ The accuracy of these DFT calculations was calibrated by comparison with experimental literature data from [DBsq[•]]^{-.47}

The dicationic complex, $[1^{\cdot}]^{2^+}$, was investigated using a BS(1,1) method, and by spin unrestricted S = 0 (singlet) and S = 1 (triplet) calculations. The spin unrestricted singlet state is defined as removal of an electron from the SOMO of $[1^{\cdot}]^+$; the spin unrestricted triplet state (S = 1) is identical to the triplet state of the BS(1,1) calculation. The calculated spin triplet solution is 2 kcal mol⁻¹ more stable than the BS singlet, and 11 kcal mol⁻¹ more stable than the spin unrestricted singlet state. The exchange coupling constant, determined from the high-spin and BS energies together with the corresponding spin-expectation values $\langle S^2 \rangle$,^{59,60} is calculated to be J = +731 cm⁻¹.

The tris-dioxolene ligand in $[1^{\cdot}]^{2+}$ is in its tetra-anionic form, $[ctc^{+}]^{4-}$, involving an electron being removed from the 3A" MO of [1']⁺ (Fig. 9). All three dioxolene units are metrically identical (Table S7†). The mean C-O (1.327 Å) and aromatic C-C (1.413 Å) distances are consistent with the average obtained for two semiquinones and one catecholate. The inter-metal separation is greater for $[1^{\cdot}]^{2+}$ than $[1^{\cdot}]^{+}$, and stems from a subtle tilt of the PtP₂O₂ square plane away from the mean plane of the dioxolene unit. Examination of the frontier orbitals shows two near degenerate SOMOs whose composition resembles the corresponding MOs in 1. The HOMO-1 is the totally symmetric 1A₁ MO. Oxidation of the ligand has stabilised the Pt-L σ^* orbitals (3A₂ and 5E shown in blue, Fig. 9) such that they are energetically matched with the dppb π^* (2A₂ and E, in red) and mix in $[1^{\cdot}]^{2+}$ (shown in purple). This dilutes their Pt content to \sim 28%, down from \sim 40% for [1']⁺. The subtle shift in energy of



Fig. 8 Mulliken spin populations for: (a) [DBsq[•]]⁻; (b) [4[•]]⁺; (c) [5[•]]⁺; (d) [1[•]]²⁺; (f) [1^{••}]³⁺ (red: α-spin; yellow: β-spin).



Fig. 9 Qualitative MO scheme depicting the ordering of frontier orbitals for the $[1]^{0/1+/2+/3+}$ electron transfer series, derived from B3LYP-ZORA DFT calculations. The idealised orbitals shown left are derived from the calculation of 1 annotated with C_{3v} symmetry labels. The purple levels represent mixing of 2A₂ and 4E (red) with 3A₂ and 5E (blue). Detailed schemes and MO plots are in Fig. S25–S28†.

the Pt 5d orbitals leads to an increase in the metal content to 7% in HOMO-2 and HOMO-3 (1A₂ and 1E, Fig. 9). The even distribution unpaired spins with +0.66 spins per arm of the cavitand underscores the three-fold symmetry of $[1^{\cdot}]^{2+}$ (Fig. 6e).

Broken symmetry calculation of the electronic structure of $[1^{3^+}]^{3^+}$ give the spin doublet to be only 0.4 kcal mol⁻¹ more stable than the quartet: the S = 1/2 solution is shown in Fig. 7. The complex is again C_{3v} symmetric as demonstrated by the equivalent bond distances in the three {Pt(dioxolene)} units. The average C–O bond length is shortened to 1.308 Å, and the average aromatic C-C distance of 1.418 Å match those for the semiquinone ligands in the optimised structures of $[4^{\cdot}]^{+}$ and $[5']^+$ (Table S4[†]). Thus, $[1'']^{3+}$ possesses a tris-semiquinone $[ctc^{*}]^{3-}$ ligand. The accumulation of unpaired spins on each of the dioxolene arms results in a subtle opening of the cavitand. The HOMO and HOMO-1 are the ctc-based 1A₂ and 1E MOs. The 5d orbitals are further stabilised by the reduced ligand field strength of the oxidised ctc ligand where the LUMO+1 and LUMO+2 are now exclusively the Pt-L σ^* possessing \sim 42% metal character, and match the corresponding

Table 4	Electronic	absorption	data for	the comp	olexes
---------	------------	------------	----------	----------	--------

	Proton ^a	$ ho_{ m calcd}$	$ ho_{\mathrm{exptl}}$
[DBsg [•]] ^{-b}	H4	-0.0060	-0.0068
1]	H6	-0.0017	-0.0013
[4 '] ⁺	H4	-0.0062	-0.0067
	<i>H</i> 6	-0.0010	-0.0013
[5 [•]] ⁺	H4	-0.0054	-0.0068
	H6	-0.0012	-0.0013
[1 [.]] ⁺	$H3/H6^{c}$	-0.0008	-0.0013
[3 [•]] ⁺	$H3/H6^{c}$	-0.0009	-0.0017
$[DBsq']^{-b}$	H4	-0.0060	-0.0068

^a See ESI for atom numbering. ^b Experimental data taken from ref. 46.
^c Values averaged for these protons.

orbitals in **1** and $[1^{\cdot}]^{+}$ (*vide supra*). Three SOMOs originating from the 1A₁ and 2E MOs are generated: two α -spin (spin-up) and one β -spin (spin-down), where the latter is antiferromagnetically coupled to the corresponding α -spin SOMO of matching symmetry (Fig. 9 and S28†). The magnitude of the coupling is expressed by the overlap integral (*S*), which ranges from S = 0 for fully uncoupled (or perfectly orthogonal magnetic orbitals) to S = 1 for two electrons in a single MO. The value computed here of S = 0.19 is shows the antiferromagnetic coupling is relatively weak, consistent with the small exchange constant. This energy gap of 134 cm⁻¹ (2*J*) represents the energy required to flip the β -spin to generate a spin quartet with three α -spin SOMOs, which would be populated at room temperature. The Mulliken spin density analysis shows one unpaired electron per dioxolene unit with 2% of the spin located on the Pt ions (Fig. 8f).

Notably, the calculated value of $2J = 134 \text{ cm}^{-1}$ for $[1 \cdot \cdot \cdot]^{3^*}$ is comparable to measured values from other poly-dioxolene complexes at the all-sq level. These exhibit $1 \leq |J| \leq 209 \text{ cm}^{-1}$, depending on the organic linker between the dioxolene rings and the dihedral angle between them.^{14,20,21} No measured J values from a mixed-valent cat/sq system analogous to $[1^* \cdot \cdot]^{2^+}$ are available. However, an *ab initio* study of a different metal/ organic tris-dioxolene system predicted that magnetic coupling in the mixed-valent cat/sq/sq diradical should be $10-100 \times$ stronger than the corresponding sq/sq/sq triradical.⁶¹ That agrees with the much larger superexchange constant calculated for $[1^* \cdot \cdot]^{2^+}$ (+731 cm⁻¹) than for $[1^* \cdot \cdot]^{3^+}$.

Time-dependent (TD) DF calculations were carried out for the four-membered series $[1]^{0/1+/2+/3+}$ in a dichloromethane solvent continuum. This method has reliably reproduced experimental data in analogous systems,^{13,23} the use of a solvent continuum being essential as corresponding gas phase calculations greatly underestimate the energy of IVCT bands.⁶² The position of the computed transitions matches the spectral profile, but their intensities are an order of magnitude larger than experiment (Fig. 10). Contributors to that discrepancy may include partial decomposition during electro-generation of the radical complexes (Fig. 3); the influence of rigidly encapsulated solvent within the molecular cavity;27 and the absence of vibrationally induced transition moments in the TD-DF treatment. The hallmark feature reproduced by TD-DF is the IVCT band in $[1^{\cdot}]^{+}$ and $[1^{\cdot}]^{2+}$, whose computed energies at 6310 cm^{-1} (cf. 7900 cm $^{-1}$ in [1']⁺) and 6204 cm $^{-1}$ (cf. 7100 cm $^{-1}$ in $[1^{\cdot}]^{2^+}$) are well within the generally accepted error.⁶³ This is defined as the $1A_1 \rightarrow 2E$ excitation in Fig. 9, or more explicitly the $2A'' \rightarrow 3A''$ transition for $[1']^+$ with its C_s molecular symmetry (Fig. S26[†]). The transition is computed to be twice as intense in the dication.

The weak band at 13 700 cm⁻¹ in the calculated spectrum of $[1^{\cdot}]^{2+}$ most likely corresponds to the 14 300 cm⁻¹ peak observed experimentally (Fig. 3). It is defined as the 1E \rightarrow 2E transition, where the low intensity stems from the poor overlap between these combinations of dioxolene π orbital (Fig. 9). The analogous transition in $[1^{\cdot}]^+$ is not evident in the calculation, despite its very weak appearance in the experimental spectrum (Fig. 3). The broad transition envelope centered at ~22 000 cm⁻¹ in the mixed-valent species comprises excitations from the HOMO-8 and HOMO-10, and HOMO-4 and HOMO-5 (Fig. S32 and S33†). The former are a combination of d_{yz} (~20%), dppb π and some tris-dioxolene character of the same phase as the singly occupied acceptor orbitals – 3A" in $[1^{\cdot}]^+$ and 2E in $[1^{\cdot}]^{2+}$ – while the latter are excitations to the



Fig. 10 Calculated electronic absorption spectra for 1 (black), $[1^{\cdot}]^+$ (blue), $[1^{\cdot}]^{2+}$ (red), and $[1^{\cdot} \cdot \cdot]^{3+}$ (green).

SOMOs from MOs with *ca.* 57% d_{z^2} character. These transitions dominate the electronic spectrum of $[1^{\cdot \cdot \cdot}]^{3+}$ but in this case, the donor orbitals are have ~45% mixed d_{yz} and d_{z^2} character (Fig. S34[†]) reflecting the weaker field strength of the tris-semiquinone ligand. The spectrum of $[1^{\cdot \cdot \cdot}]^{3+}$ lacks the intense IVCT band shown by $[1^{\cdot}]^+$ and $[1^{\cdot \cdot}]^{2+}$, which is the hallmark for three dioxolene groups at the same oxidation level in the $[Cr(dioxolene)_3]^z$ (z = 3+, 2+, 1+, 0, 1-, 2-, 3-) series.^{7,64} A weak band at 10 080 cm⁻¹ is assigned as the SOMO \rightarrow LUMO excitation whose intensity is proportional to the poor overlap between this collection of near orthogonal orbitals (Fig. S28[†]).

Conclusions

The $[ctc]^{6-/5-/4-/3-}$ redox series, which was first noted seventeen years ago in a series of platinum(II) complexes (Scheme 1),²⁷ has been characterised by spectroscopic (UV/vis/NIR, cw and pulsed EPR) and computational methods. The ctc radical complexes are poorly stable above 270 K, which has thus far precluded their isolation as pure compounds. Their instability may stem from the poor steric protection afforded by the exposed CH₂ groups in the ctc macrocycle.⁶⁵ None-the-less, with the help of data from monometallic model compounds, the mixed valent character of coordinated $[ctc']^{5-}$ radicals has been determined in detail.

The ligand radicals [1[•]]⁺ and [3[•]]⁺ have class II character, with more localised SOMOs than radicals derived from fully conjugated triscat,²³ or the comparable bis-dioxolene thea (Scheme 1).¹⁵ In the latter case, this will reflect differing degrees

View Article Online Edge Article

of through-space overlap between neighbouring dioxolene π -systems, which are linked by one methylene bridge in ctc and two methylene bridges in thea. Electron hopping around the [ctc⁻]⁵⁻ macrocycle is rapid in fluid solution, but slows below the EPR timescale when the solution is frozen.¹⁵ While the metal/ligand character of the SOMO in [1⁻]⁺ and [3⁻]⁺ is essentially the same (Table 3), they exhibit differing temperature dependence of unpaired spin delocalisation. That might explain the different trends in the oxidation potentials and IVCT linewidths of [1]ⁿ⁺ and [3]ⁿ⁺ (n = 1–3), as their dioxolene rings are sequentially oxidised.

Intramolecular mixed-valency between three or more redox centers is well-known,⁹ with examples based on redox sites that are well-separated (by a 1,3,5-triphenylene scaffold, for example); linked by fully conjugated spacers (like triscat (Scheme 1) or hexaazatriphenylene);^{9,23,30} or, bound by a centrally coordinated metal ion.^{7,43,66} The mixed-valent macrocycle [ctc⁻]^{5–} is distinct from these scenarios, in having three dioxolene redox sites that interaction *via* through-space π – π overlap.³¹ As such, it is the first mixed-valent radical derived from a tri-cyclophane-type precursor to be spectroscopically characterised.³¹ Our current work aims to incorporate additional steric protection onto the ctc framework,⁶⁷ to produce mixed-valent radicals that are stable enough for use in frameworks and supramolecular architectures.^{24,32}

Experimental

Cyclotricatechylene (H_6ctc) ,³⁵ [PtCl₂(L)] (L = dppb, dppe and ^tBu₂bipy),⁶⁸ [{Pt(dppb)}₃(μ_3 -ctc)] (1),²⁷ [Pt(dppe)(DBcat)] (4) and [Pt(^tBu₂bipy)(DBcat)] (5)¹³ were all prepared by the literature procedures. Other reagents were used as supplied. Experimental data and procedures for the crystal structure determinations, electrochemical and spectroelectrochemical measurements, cw and pulsed EPR experiments and the computational study are given in the ESI.†

Synthesis of $[{Pt(dppe)}_3(\mu_3-ctc)]$ (2)

A solution of H_6 ctc (0.037 g, 0.1 mmol) and [PtCl₂(dppe)] (0.24 g, 0.37 mmol) in deoxygenated dimethylacetamide (25 cm³) was stirred for 30 min at room temperature. A suspension of K₂CO₃ (0.102 g, 0.73 mmol) in dry, deoxygenated methanol (10 cm^3) was then added, and the mixture was stirred at 70 $^\circ$ C for 18 h resulting in the formation of a deep yellow colouration. After cooling, excess dry, deoxygenated diethyl ether was added to precipitate the crude yellow product which was isolated under a nitrogen atmosphere. Recrystallisation from dry, deoxygenated dichloromethane/acetone yielded the product as a deep yellow powder. Yield: 0.035 g (16%). Found C, 53.2; H, 4.00%. Calcd for C₉₉H₉₂O₁₀P₆Pt₃ C, 53.7; H, 4.19%. ES-MS *m*/*z* 629.1 $(53, [Pt(dppe)(OH_2)_2]^+), 689.1 (23, [Pt(dppe)(OH_2)(dmso)]^+),$ 706.1 (27, [NaPt(dppe)(O₂CH)₂]⁺), 1093.1 (16, [Na₂Pt₃(dppe)₃-(ctc)]²⁺), 1101.1 (26, [NaKPt₃(dppe)₃(ctc)]²⁺), 1109.1 (38, [K₂Pt₃- $(dppe)_3(ctc)]^{2+}$, 1131.7 (23, $[Na_2Pt_3(dppe)_3(ctc)(dmso)]^{2+}$), 1140.2 (33, [NaKPt₃(dppe)₃(ctc)(dmso)]²⁺), 1146.2 (27, [K₂Pt₃-(dppe)₃(ctc)(dmso)]²⁺), 2163.3 (15, [NaPt₃(dppe)₃(ctc)]⁺), 2179.4

(100, $[\text{KPt}_3(\text{dppe})_3(\text{ctc})]^+$). ¹H NMR ({CD₃}₂SO) δ 3.60 (m, 12H, dppe C₂H₄), 4.54 (br, 3H, ctc CH₂), 6.53 (s, 6H, ctc CH), 7.51 and 8.06 (both m, 60H, dppe C₆H₅) – an additional diastereotopic ctc CH₂ resonance expected near 3.3 ppm was obscured under the dmso water peak. ³¹P NMR ({CD₃}₂SO) δ 30.8 (J_{P-Pt} 3350 Hz).

Synthesis of $[{Pt(^{t}Bu_{2}bipy)}_{3}(\mu_{3}-ctc)](3)$

Method as for 2, using $[PtCl_2(^tBu_2bipy)]$ (0.20 g, 0.37 mmol). After addition of K₂CO₃ the mixture was heated to 40 °C for 18 h. After cooling, the blue solution was diluted with dry, deoxygenated 1:1 diethyl ether/pentane and the blue precipitate collected. The product could be purified by either recrystallisation from dichloromethane/acetone at -20 °C or silica column chromatography (98.5:1.5 CH₂Cl₂/MeOH eluent) to yield the product as a blue powder. Yield: 0.049 g (28%). Found C, 50.1; H, 5.15; N, 4.3%. Calcd for C₇₅H₉₀N₆O₉Pt₃ C, 49.9; H, 5.03; N, 4.7%. ES-MS (MeCN) m/z 269.2 (48, $[H(^tBu_2bipy)]^+)$, 522.1 (26, $[Pt(^{t}Bu_{2}bipy)(NCCH_{3})(OH_{2})]^{+}$), 549.1 (26, $[Pt(^{t}Bu_{2}-t)]^{+}$), 549.1 (26, bipy)(NCCH₃)(O₂CH)]⁺), 572.2 (39, [NaPt(^tBu₂bipy)(NCCH₃)(O₂- $(CH)^{+}$, 584.2 (100, $[H_3Pt_3-(^{t}Bu_2bipy)_3(ctc)]^{3+}$), 597.2 (20, $[H_2KPt_3(^tBu_2bipy)_3(ctc)]^{3+}), 609.7 (26, [HK_2Pt_3(^tBu_2bipy)_3-$ (ctc)]³⁺), 617.3 (19, [NaK₂Pt₃(^tBu₂bipy)₃(ctc)]³⁺), 645.4 (24, [H₂- $Pt_2(^{t}Bu_2bipy)_2(ctcH_2)]^{2+}$, 663.4 (23, $[HKPt_2(^{t}Bu_2bipy)_2(ctcH_2)]^{2+}$), 875.8 (30, $[H_2Pt_3(^tBu_2bipy)_3(ctc)]^{2+}$), 887.3 (23, $[HNaPt_3(^tBu_2$ bipy)₃-(ctc)]²⁺), 895.3 (23, [HKPt₃(^tBu₂bipy)₃(ctc)]²⁺), 1167.4 (27, $[NaPt_2(^tBu_2bipy)(NCMe)_2(ctcH)(O_2CH)]^+)$, 1288.4 (9, $[Pt_2(^tBu_2$ bipy)₂(ctcH₂)]⁺), 1751.5 (49, [HPt₃(^tBu₂bipy)₃(ctc)]⁺), 1772.5 (28, [NaPt₃(^tBu₂bipy)₃(ctc)]⁺), 1789.5 (28, [KPt₃(^tBu₂bipy)₃(ctc)]⁺). ¹H NMR (CDCl₃) δ 1.28 (s, 54H, C{CH₃}₃), 3.30 and 4.51 (both d, 14.0 Hz, 3H, ctc CH₂), 6.67 (s, 6H, ctc CH), 7.35 (dd, 1.9 and 6.4 Hz, 6H, Py H⁵), 7.84 (d, 1.9 Hz, 6H, Py H³), 8.77 (d, 6.4 Hz, 6H, Py H⁶).

Acknowledgements

This work was supported by the EPSRC (EP/F040547/1). EPR spectra were recorded at the EPSRC National EPR Facility at Manchester. We also thank Diamond Light Source for access to beamline I19 (MT7847) that contributed to the results presented here, and Professor Mike Ward (University of Sheffield) for use of the spectroelectrochemistry facilities.

Notes and references

- C. G. Pierpont and R. M. Buchanan, *Coord. Chem. Rev.*, 1981, 38, 45; C. G. Pierpont and C. W. Lange, *Prog. Inorg. Chem.*, 1993, 41, 331; C. G. Pierpont, *Coord. Chem. Rev.*, 2001, 216– 217, 99.
- 2 A. Dei, D. Gatteschi, C. Sangregorio and L. Sorace, *Acc. Chem. Res.*, 2004, **37**, 827.
- 3 D. A. Shultz, Comments Inorg. Chem., 2002, 23, 1.
- 4 J. S. Miller and K. S. Min, Angew. Chem., Int. Ed., 2009, 48, 262.
- 5 S. Archer and J. A. Weinstein, *Coord. Chem. Rev.*, 2012, 256, 2530.
- 6 T. Tezgerevska, K. G. Alley and C. Boskovic, *Coord. Chem. Rev.*, 2014, **268**, 23.

- 7 C. G. Pierpont, *Coord. Chem. Rev.*, 2001, 219–221, 415;
 C. G. Pierpont, *Inorg. Chem.*, 2011, 50, 9766.
- 8 W. Kaim and B. Schwederski, *Coord. Chem. Rev.*, 2010, 254, 1580; O. R. Luca and R. H. Crabtree, *Chem. Soc. Rev.*, 2013, 42, 1440.
- 9 J. Hankache and O. S. Wenger, *Chem. Rev.*, 2011, 111, 5138;
 A. Heckmann and C. Lambert, *Angew. Chem., Int. Ed.*, 2012, 51, 326.
- 10 L. F. Joulié, E. Schatz, M. D. Ward, F. Weber and L. J. Yellowlees, *J. Chem. Soc., Dalton Trans.*, 1994, 800;
 A. M. Barthram, Z. R. Reeves, J. C. Jeffery and M. D. Ward, *J. Chem. Soc., Dalton Trans.*, 2000, 3162.
- 11 G. A. Abakumov, V. K. Cherkasov, V. I. Nevodchikov, N. O. Druzhkov and V. A. Kuropatov, *Russ. Chem. Bull.*, 1998, 47, 875.
- 12 A. Bencini, C. A. Daul, A. Dei, F. Mariotti, H. Lee, D. A. Shultz and L. Sorace, *Inorg. Chem.*, 2001, **40**, 1582.
- 13 J. Best, I. V. Sazanovich, H. Adams, R. D. Bennett, E. S. Davies, A. J. H. M. Meijer, M. Towrie, S. A. Tikhomirov, O. V. Bouganov, M. D. Ward and J. A. Weinstein, *Inorg. Chem.*, 2010, **49**, 10041.
- K. G. Alley, G. Poneti, P. S. D. Robinson, A. Nafady,
 B. Moubaraki, J. B. Aitken, S. C. Drew, C. Ritchie,
 B. F. Abrahams, R. K. Hocking, K. S. Murray, A. M. Bond,
 H. H. Harris, L. Sorace and C. Boskovic, *J. Am. Chem. Soc.*, 2013, 135, 8304.
- 15 J. J. Loughrey, S. Sproules, E. J. L. McInnes, M. J. Hardie and M. A. Halcrow, *Chem.–Eur. J.*, 2014, **20**, 6272.
- 16 G. A. Abakumov, V. K. Cherkasov, V. I. Nevodchikov and V. A. Kuropatov, *Inorg. Chem.*, 1998, 37, 6117;
 G. A. Abakumov, V. K. Cherkasov, V. I. Nevodchikov and V. A. Kuropatov, *Inorg. Chem.*, 2001, 40, 2434;
 A. I. Poddel'sky, A. V. Piskunov, N. O. Druzhkov,
 G. K. Fukin, V. K. Cherkasov and G. A. Abakumov, *Z. Anorg. Allg. Chem.*, 2009, 635, 2563;
 A. I. Poddel'sky, N. O. Druzhkov, V. K. Cherkasov and
 G. A. Abakumov, *J. Organomet.Chem.*, 2011, 696, 517.
- 17 Y. Suenaga, Y. Hirano, Y. Umehata and T. Minematsu, *Inorg. Chim. Acta*, 2011, **365**, 505.
- 18 A. Dei and L. Sorace, Dalton Trans., 2003, 3382.
- M. Affronte, A. Beni, A. Dei and L. Sorace, *Dalton Trans.*, 2007, 5253; K. G. Alley, G. Poneti, J. B. Aitken, R. K. Hocking, B. Moubaraki, K. S. Murray, B. F. Abrahams, H. H. Harris, L. Sorace and C. Boskovic, *Inorg. Chem.*, 2012, **51**, 3944.
- 20 D. A. Shultz, R. M. Fico Jr, S. H. Bodnar, R. K. Kumar, K. E. Vostrikova, J. W. Kampf and P. D. Boyle, *J. Am. Chem. Soc.*, 2003, **125**, 11761.
- 21 A. Caneschi, A. Dei, H. Lee, D. A. Shultz and L. Sorace, *Inorg. Chem.*, 2001, 40, 408; D. A. Shultz, S. H. Bodnar, H. Lee, J. W. Kampf, C. D. Incarvito and A. L. Rheingold, *J. Am. Chem. Soc.*, 2002, 124, 10054; A. Beni, A. Dei, D. A. Shultz and L. Sorace, *Chem. Phys. Lett.*, 2006, 428, 400; J. C. Sloop, D. A. Shultz, T. Coote, B. Shepler, U. Sullivan, J. W. Kampf and P. D. Boyle, *J. Phys. Org. Chem.*, 2012, 25, 314.
- Y. Suenaga and C. G. Pierpont, *Inorg. Chem.*, 2005, 44, 6183;
 Y. Suenaga, Y. Umehata, Y. Hirano, T. Minematsu and C. G. Pierpont, *Inorg. Chim. Acta*, 2008, 361, 2941.

- 23 C. S. Grange, A. J. H. M. Meijer and M. D. Ward, *Dalton Trans.*, 2010, **39**, 200.
- 24 M. Hmadeh, Z. Lu, Z. Liu, F. Gándara, H. Furukawa, S. Wan, V. Augustyn, R. Chang, L. Liao, F. Zhou, E. Perre, V. Ozolins, K. Suenaga, X. Duan, B. Dunn, Y. Yamamto, O. Terasaki and O. M. Yaghi, *Chem. Mater.*, 2012, 24, 3511.
- 25 A. Caneschi, A. Dei, C. P. Mussari, D. A. Shultz, L. Sorace and K. E. Vostrikova, *Inorg. Chem.*, 2002, **41**, 1086.
- 26 D. A. Shultz and R. K. Kumar, J. Am. Chem. Soc., 2001, 123, 6431.
- 27 D. S. Bohle and D. Stasko, Chem. Commun., 1998, 567.
- 28 B. F. Abrahams, N. J. FitzGerald, T. A. Hudson, R. Robson and T. Waters, Angew. Chem., Int. Ed., 2009, 48, 3129;
 B. F. Abrahams, N. J. FitzGerald and R. Robson, Angew. Chem., Int. Ed., 2010, 49, 2896;
 B. F. Abrahams, B. A. Boughton, N. J. FitzGerald, J. L. Holmes and R. Robson, Chem. Commun., 2011, 47, 7404.
- 29 J. J. Loughrey, C. A. Kilner, M. J. Hardie and M. A. Halcrow, Supramol. Chem., 2012, 24, 2; P. Satha, G. Illa and C. S. Purohit, Cryst. Growth Des., 2013, 13, 2636.
- 30 K. D. Demadis, C. M. Hartshorn and T. J. Meyer, *Chem. Rev.*, 2001, **101**, 2655; D. M. D'Alessandro and F. R. Keene, *Chem. Soc. Rev.*, 2006, **35**, 424.
- 31 Intramolecular delocalisation of unpaired spins between pairs of cofacial aromatic rings is well known, in radicals related to [2,2]cyclophanes.⁹ See: e.g. D. Sun, S. V. Rosokha and J. K. Kochi, J. Am. Chem. Soc., 2004, 126, 1388; P. H. Dinolfo, V. Coropceanu, J. L. Brédas and J. T. Hupp, J. Am. Chem. Soc., 2006, 128, 12592; A. S. Jalilov, S. F. Nelsen, I. A. Guzei and Q. Wu, Angew. Chem., Int. Ed., 2011, 50, 6860; Y. Wu, M. Frasconi, D. M. Gardner, P. R. McGonigal, S. T. Schneebeli, M. R. Wasielewski and J. F. Stoddart, Angew. Chem., Int. Ed., 2014, 53, 9476; M. Castellano, R. Ruiz-García, J. Cano, J. Ferrando-Soria, E. Pardo, F. R. Fortea-Pérez, S.-E. Stiriba, W. P. Barros, H. O. Stumpf, L. Cañadillas-Delgado, J. Pasán, C. Ruiz-Pérez, G. de Munno, D. Armentano, Y. Journaux, F. Lloret and M. Julve, Coord. Chem. Rev., 2015, 303, 110.
- 32 M. J. Hardie, *Chem. Soc. Rev.*, 2010, **39**, 516; J. J. Henkelis and
 M. J. Hardie, *Chem. Commun.*, 2015, **51**, 11929.
- J. M. Clemente, C. Y. Wong, P. Bhattacharyya, A. M. Z. Slawin,
 D. J. Williams and J. D. Woollins, *Polyhedron*, 1994, 13, 261;
 J. D. Kinder and W. J. Youngs, *Organometallics*, 1996, 15, 460;
 M. J. G. Lesley, W. Clegg, T. B. Marder, N. C. Norman,
 A. G. Orpen, A. J. Scott and J. Starbuck, *Acta Crystallogr.*, *Sect. C: Struct. Chem.*, 1999, 55, 1272.
- 34 O. Carugo, C. B. Castellani, K. Djinović and M. Rizzi, *J. Chem. Soc., Dalton Trans.*, 1992, 837; S. Brown, *Inorg. Chem.*, 2012, 51, 1251.
- 35 A. Chakrabarti, H. M. Chawla, G. Hundal and N. Pant, *Tetrahedron*, 2005, **61**, 12323.
- 36 See e.g. J. A. Hyatt, E. N. Duesler, D. Y. Curtin and I. C. Paul, *J. Org. Chem.*, 1980, 45, 5074; J. W. Steed, H. Zhang and J. A. Atwood, *Supramol. Chem.*, 1997, 7, 37; C. J. Sumby and M. J. Hardie, *Acta Crystallogr., Sect. E: Struct. Rep. Online*, 2007, 63, 01537.
- 37 A. L. Spek, J. Appl. Crystallogr., 2003, 36, 7.

- 38 J. A. Weinstein, M. T. Tierney, E. S. Davies, K. Base, A. A. Robeiro and M. W. Grinstaff, *Inorg. Chem.*, 2006, 45, 4544.
- 39 N. M. Shavaleev, E. S. Davies, H. Adams, J. Best and J. A. Weinstein, *Inorg. Chem.*, 2008, 47, 1532.
- 40 H.-C. Chang, T. Shiozaki, A. Kamata, K. Kishida, T. Ohmori, D. Kiriya, T. Yamauchi, H. Furukawa and S. Kitagawa, J. Mater. Chem., 2007, 17, 4136.
- 41 E. J. L. McInnes, R. D. Farley, S. A. Macgregor, K. J. Taylor, L. J. Yellowlees and C. C. Rowlands, *J. Chem. Soc., Faraday Trans.*, 1998, 94, 2985.
- 42 E. J. L. McInnes, R. D. Farley, C. C. Rowlands, A. J. Welch, L. Rovatti and L. J. Yellowlees, *J. Chem. Soc., Dalton Trans.*, 1999, 4203.
- 43 J. England, C. C. Scarborough, T. Weyhermüller, S. Sproules and K. Wieghardt, *Eur. J. Inorg. Chem.*, 2012, 4605;
 A. C. Bowman, J. England, S. Sproules, T. Weyhermüller and K. Wieghardt, *Inorg. Chem.*, 2013, 52, 2242; M. Wang, T. Weyhermüller, J. England and K. Wieghardt, *Inorg. Chem.*, 2013, 52, 12763; M. Wang, J. England, T. Weyhermüller and K. Wieghardt, *Inorg. Chem.*, 2014, 53, 2276; M. Wang, T. Weyhermüller and K. Wieghardt, *Chem.-Eur. J.*, 2014, 20, 9037.
- 44 F. Hartl and A. Vlček jr, Inorg. Chem., 1996, 35, 1257.
- 45 T. Weyland, K. Costuas, L. Toupet, J.-F. Halet and C. Lapinte, Organometallics, 2000, **19**, 4228; D. M. D'Alessandro and F. R. Keene, *Chem.-Eur. J.*, 2005, **11**, 3679.
- 46 M. B. Robin and P. Day, *Adv. Inorg. Chem. Radiochem.*, 1967, 10, 247.
- 47 O. Ryba, J. Pillař and J. Petránek, *Collect. Czech. Chem. Commun.*, 1968, **33**, 26.
- 48 D. Collison, F. E. Mabbs, E. J. L. McInnes, K. J. Taylor, A. J. Welch and L. J. Yellowlees, *J. Chem. Soc., Dalton Trans.*, 1996, 329.
- 49 E. A. M. Geary, K. L. McCall, A. Turner, P. R. Murray, E. J. L. McInnes, L. A. Jack, L. J. Yellowlees and N. Robertson, *Dalton Trans.*, 2008, 3701.
- 50 J. S. Pap, F. L. Benedito, E. Bothe, E. Bill, S. DeBeer George, T. Weyhermüller and K. Wieghardt, *Inorg. Chem.*, 2007, 46, 4187.
- 51 P. H. Rieger, J. Magn. Reson., 1997, 124, 140.
- 52 W. Kaim, A. Dogan, M. Wanner, A. Klein, I. Tiritiris, T. Schleid, D. J. Sturfkens, T. L. Snoeck, E. J. L. McInnes, J. Fiedler and S. Záliš, *Inorg. Chem.*, 2002, **41**, 4139.

- 53 A. Klein, E. J. L. McInnes, T. Scheiring and S. Záliš, *J. Chem. Soc., Faraday Trans.*, 1998, **94**, 2979.
- 54 O. Burghaus, M. Plato, M. Rohrer, K. Mobius, F. MacMillan and W. Lubitz, J. Phys. Chem., 1993, 97, 7639.
- 55 F. Gerson and S. Coskun, J. Chem. Soc., Perkin Trans. 2, 1997, 1127.
- 56 K. Shiren and K. Tanaka, Inorg. Chem., 2002, 41, 5912.
- 57 D.-N. Horng, J.-P. Chyn, K.-J. Shieh, J.-L. Chou and Y.-S. Wen, Acta Crystallogr., Sect. C: Struct. Chem., 1999, 55, 652.
- 58 A. Carrington and A. D. McLachlan, *Introduction to Magnetic Resonance*, Harper & Row, New York, USA, 1967.
- 59 T. Soda, Y. Kitagawa, T. Onishi, Y. Takano, Y. Shigetu, H. Nagao, Y. Yoshioka and K. Yamaguchi, *Chem. Phys. Lett.*, 2000, **319**, 223.
- 60 K. Yamaguchi, Y. Takahara and T. Fueno, in *Applied Quantum Chemistry*, ed. V. H. Smith, Reidel, Dordrecht, The Netherlands, 1986, p. 155.
- 61 S. Franzen and D. A. Shultz, J. Phys. Chem. A, 2003, 107, 4292.
- 62 A. A. Vlček and S. Záliš, Coord. Chem. Rev., 2007, 251, 258.
- 63 F. Neese, JBIC, J. Biol. Inorg. Chem., 2006, 11, 702.
- 64 H.-C. Chang, H. Myasaka and S. Kitagawa, *Inorg. Chem.*, 2001, 40, 146; H.-C. Chang and S. Kitagawa, *Angew. Chem.*, *Int. Ed.*, 2002, 41, 130; R. R. Kapre, E. Bothe, T. Weyhermüller, S. DeBeer George, N. Muresan and K. Wieghardt, *Inorg. Chem.*, 2007, 46, 7827.
- 65 Steric protection *para* to the oxy substituent(s) in phenoxyltype radicals greatly enhances their stability. P. Chaudhuri and K. Wieghardt, *Prog. Inorg. Chem.*, 2001, **50**, 151.
- 66 Another series of macrocyclic mixed valent compounds based on a metallacyclic architecture has also been reported. N. Shan, S. J. Vickers, H. Adams, M. D. Ward and J. A. Thomas, *Angew. Chem., Int. Ed.*, 2004, 43, 3938; N. Shan, J. D. Ingram, T. L. Easun, S. J. Vickers, H. Adams, M. D. Ward and J. A. Thomas, *Dalton Trans.*, 2006, 2900; A. Zubi, A. Wragg, S. Turega, H. Adams, P. J. Costa, V. Félix and J. A. Thomas, *Chem. Sci.*, 2015, 6, 1334.
- 67 M. Harig, B. Neumann, H.-G. Stammler and D. Kuck, *Eur. J. Org. Chem.*, 2004, 2381.
- 68 W. Levason and C. A. McAuliffe, *Inorg. Chim. Acta*, 1976, 16, 167; K. Yasufuku, H. Noda and H. Yamazaki, *Inorg. Synth.*, 1989, 26, 369; S. Huertas, M. Hissler, J. E. McGarrah, R. J. Lachicotte and R. Eisenberg, *Inorg. Chem.*, 2001, 40, 1183.