Dalton Transactions



PAPER

View Article Online
View Journal | View Issue



Cite this: *Dalton Trans.*, 2015, **44**, 6268

Received 19th January 2015, Accepted 20th February 2015 DOI: 10.1039/c5dt00246j

www.rsc.org/dalton

Synthesis and (spectro)electrochemistry of mixed-valent diferrocenyl-dihydrothiopyran derivatives†

Konrad Kowalski,*^a Rafał Karpowicz,^a Grzegorz Mlostoń,^b Dominique Miesel,^c Alexander Hildebrandt,^c Heinrich Lang,^c Rafał Czerwieniec^d and Bruno Therrien^e

Three novel diferrocenyl complexes were prepared and characterised. 2,2-Diferrocenyl-4,5-dimethyl-3,6-dihydro-2H-thiopyran (**1**, sulphide) was accessible by the hetero-Diels-Alder reaction of diferrocenyl thioketone with 2,3-dimethyl-1,3-butadiene. Stepwise oxidation of **1** gave the respective oxides 2,2-diferrocenyl-4,5-dimethyl-3,6-dihydro-2H-thiopyran-1-oxide (**2**, sulfoxide) and 2,2-diferrocenyl-4,5-dimethyl-3,6-dihydro-2H-thiopyran-1,1-dioxide (**3**, sulfone), respectively. The molecular structures of **1** and **3** in the solid state were determined by single crystal X-ray crystallography. The oxidation of sulphide **1** to sulfone **3**, plays only a minor role on the overall structure of the two compounds. Electrochemical (cyclic voltammetry (= CV), square wave voltammetry (= SWV)) and spectroelectrochemical (*in situ* UV-Vis/NIR spectroscopy) studies were carried out. The CV and SWV measurements showed that an increase of the sulphur atom oxidation from -2 in **1** to +2 in **3** causes an anodic shift of the ferrocenyl-based oxidation potentials of about 100 mV. The electrochemical oxidation of **1–3** generates mixed-valent cations **1**⁺-**3**⁺. These monooxidised species display low-energy electronic absorption bands between 1000 and 3000 nm assigned to IVCT (= Inter-Valence Charge Transfer) electronic transitions. Accordingly, the mixed-valent cations **1**⁺-**3**⁺ are classified as weakly coupled class II systems according to Robin and Day.

Introduction

Recently, mixed-valent (= MV) species have attracted considerable attention in particular in the field of molecular electronics as they offer the possibility to act as model compounds for molecular wires, switches and other electronic building blocks. ^{1–15} Besides these foreseen technological applications MV compounds are used in electron transfer studies and are of key importance in biological systems. ^{16–20}

Ferrocenyl groups are often used in organometallic chemistry as redox-active terminal units, because ferrocene is thermally stable in its neutral and oxidised form.²¹ In addition,

ferrocenyl/ferrocenium groups in mixed-valent species show an excellent electrochemical reversibility of the Fe(II)/Fe(III) redox couple, *i.e.* 2,5-diferrocenyl five-membered heterocycles.^{8,22–32}

The electron transfer between the Fe(II)/Fe(III) centres, *i.e.* from the donor (Fe(II)) to the acceptor (Fe(III)) ion, manifests itself by the appearance of characteristic absorptions, *i.e.* IVCT (= Inter-Valence Charge Transfer) bands in the near infrared (NIR) spectral range.^{8,33}

Two distinct modes of the electronic communication between the donor and acceptor metal centres in the MV state exist: "through bond" and "through space". ^{5,6,34} The "through bond" mechanism is characteristic for molecules in which π-conjugated connectivities are linking the two redox-active metal termini, ^{5,6} while the "through space" mechanism requires a close proximity of the interacting centres. ³⁴ Depending on the degree of the electronic communication, three classes of MV complexes are distinguished according to the Robin and Day classification: non-coupled (class II) systems. ³⁵ Based on the linking group constitution, five structural types of dinuclear transition-metal compounds are known (types A–E), as schematically shown in Fig. 1.

Among the type **A** and type **B** molecules (Fig. 1), the respective diferrocenyl-functionalized systems have been extensively studied, $^{5,6,36,40-42}$ whereas compounds of structural type **D**

^aFaculty of Chemistry, Department of Organic Chemistry, University of Łódź, Tamka 12, PL-91403 Łódź, Poland. E-mail: kondor15@wp.pl

^bFaculty of Chemistry, Department of Organic and Applied Chemistry, University of Łódź, Tamka 12, PL-91403 Łódź, Poland

^cTechnische Universität Chemnitz, Faculty of Natural Sciences, Institute of Chemistry, Department of Inorganic Chemistry, D-09107 Chemnitz, Germany

^dUniversität Regensburg, Institut für Physikalische und Theoretische Chemie, Universitätsstraße 31, D-93040 Regensburg, Germany

^eInstitute of Chemistry, University of Neuchatel, Avenue de Bellevaux 51, CH-2000 Neuchatel, Switzerland

[†]Electronic supplementary information (ESI) available. CCDC 1031233 and 1031234. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/c5dt00246j

Dalton Transactions Paper

$$M \longrightarrow M$$

$$M \longrightarrow$$

Fig. 1 Classification of homobimetallic mixed-valence (= MV) metal complexes. M represents a metal containing redox centre. In type A molecules the centres are connected by a conjugated bridge consisting of C,C double or triple bonds, or aromatic moieties. 5,6 In type B molecules the M termini are directly connected to each other, 36 while in type D and E species the metal centres are separated by an aliphatic 37,38 or heteroatom fragment, respectively. 37-39 Type C molecules represent a special variety of type A compounds, in which the two M termini in the mixed-valent species interact most likely "through space". 34

remain almost unexplored. 37,43 The simplest representative of the Fc-based class D molecules is diferrocenylmethane $(Fc_2CH_2, Fc = Fe(\eta^5-C_5H_4)(\eta^5-C_5H_5))$, however, its monooxidised form (Fc2CH2+) does not show any detectable IVCT band. 44 On the contrary, such IVCT absorptions were observed for other derivatives, in which the methylene hydrogen atoms were replaced by methyl or ferrocenyl groups, 37,44 indicating that the appearance of IVCT transitions is related to the increased bulkiness of the linker unit due to the presence of sterically demanding groups in the latter compounds as compared to the Fc₂CH₂⁺ cation. In addition, the strength of the metal-metal interactions in type D compounds featuring CMe2, SiMe2 or GeMe2 bridges, depend on the effective distance between the metal centres.³⁷ It was found that the extent of the electronic coupling in the MV species decreases with an increase of the atomic radius of the bridging atom (e.g. in the order $CMe_2 > SiMe_2 > GeMe_2$).

In continuation of our works in the area of electron-transfer studies in (multi)ferrocenyl-functionalised organic and organometallic compounds, 8,11,22-32,45-49 we herein present the synthesis and characterisation of 2,2-diferrocenyl-4,5-dimethyl-3,6-dihydro-2*H*-thiopyran, 2,2-diferrocenyl-4,5-dimethyl-3,6dihydro-2H-thiopyran-1-oxide and 2,2-diferrocenyl-4,5dimethyl-3,6-dihydro-2H-thiopyran-1,1-dioxide. The influence of the electronic and steric effects on the electron transfer between the two ferrocenyl moieties in the respective mixedvalent species is reported. The 3,6-dihydro-2H-thiopyran

bridge was chosen, due to its bulkiness and the presence of the oxidisable sulphur atom in a position adjacent to the redox-active ferrocenyl groups. This S-atom reactivity allows us to synthesise three closely related derivatives 1-3 and investigate the influence of the different electron-withdrawing character of the aliphatic bridge on the IVCT properties of the mixed valence species 1⁺-3⁺.

Results and discussion

Synthesis

The synthetic methodologies for the preparation of 2,2-diferrocenyl-4,5-dimethyl-3,6-dihydro-2H-thiopyran (1, sulphide), 2,2diferrocenyl-4,5-dimethyl-3,6-dihydro-2*H*-thiopyran-1-oxide (2, sulfoxide) and 2,2-diferrocenyl-4,5-dimethyl-3,6-dihydro-2Hthiopyran-1,1-dioxide (3, sulfone) are shown in Scheme 1.

Sulphide 1 was prepared via the hetero-Diels-Alder cycloaddition of diferrocenylthioketone^{50,51} with 2,3-dimethyl-1,3-butadiene in a sealed glass-tube at 75 °C (Experimental section).⁵² After appropriate work-up, compound 1 was obtained as a red solid in 65% yield. Treatment of sulphide 1 with hydrogen peroxide (30%) and selenium dioxide as oxidising reagents⁵³ in methanol gave the respective sulfoxide 2, which was purified by column chromatography in 87% yield (Experimental section). Further oxidation of 2 with m-chloroperoxybenzoic acid (= MCPBA) in dichloromethane produced

$$(Fc)_{2}C=S + CH_{3} \xrightarrow{(i)} Fc \xrightarrow{S} CH_{3} \xrightarrow{(iii)} Fc \xrightarrow{CH_{3}} CH_{3} \xrightarrow{CH_{3}} CH_{3}$$

Scheme 1 Synthesis of 1-3 (MCPBA = m-chloroperoxybenzoic acid, Fc = ferrocenyl group). (i) 75 °C, 50 h; (ii) 30% H₂O₂, SeO₂, CH₃OH, 0 °C to r.t., 10 min; (iii) MCPBA, CH₂Cl₂, -20 °C, 2 h, then r.t. 24 h and 2nd portion MCPBA at r.t. 24 h.

Fig. 2 ORTEP diagrams of 1 and 3 (two independent molecules in the crystal, 3A and 3B) at 50% probability level

sulfone 3 at ambient temperature in 21% yield (Scheme 1, Experimental section). 54

Compounds 1–3 are red solids soluble in common organic solvents. All three compounds are stable towards air and moisture in the solid state and in solution.

Compounds 1–3 were characterised by IR and NMR (1 H, 13 C{ 1 H}; for more details see Fig. S1–S3, ESI†) spectroscopy and high-resolution mass spectrometry. The molecular structures of 1 and 3 in the solid state were determined by single crystal X-ray diffraction analysis. Electrochemical investigations were carried out by using cyclic voltammetry (= CV), square wave voltammetry (= SWV) and *in situ* UV-Vis/NIR spectroelectrochemistry.

X-ray structure determination

Single-crystals of 1 and 3 suitable for a single crystal X-ray diffraction analysis were obtained by slow diffusion of hexane into a dichloromethane solution of 1 and by slow diffusion of hexane into a diethyl ether solution of 3. ORTEP diagrams of 1 and 3 are shown in Fig. 2, while selected bond distances (Å) and angles (°) are listed in Table 1. Crystal and structure refinement data are presented in the ESI (Table S1†).

Compound 1 crystallizes in the orthorhombic space group Pcab, while complex 3 crystallizes in the triclinic space group $P\bar{1}$. In the crystal packing of 3, two independent molecules (A and B) are observed. The structure analysis of both molecules confirmed the expected structures in which the two ferrocenyl groups are bonded to a sulphide (1) or a sulfone (3) moiety (Fig. 2). All ferrocenyl units show an eclipsed conformation with nearly equivalent distances between Fe and the centroid of the cyclopentadienyl rings (Table 1).

The oxidation of sulphide 1 to sulfone 3, however, plays only a minor role on the overall structure of the two compounds. A similar behaviour was found for a series of sulphur-containing heterocycles in which the oxidation of the sulphur atom has almost no impact on the geometrical parameters of the heterocycles. 54,55 The only bond distances which are somewhat influenced by oxidation are S–C $_{Fc}$ and S–CH $_2$ (Table 1), whereby the corresponding bonds in 3, as compared to 1, are shortened by 0.02 Å, as previously observed by Petrov. 54,55 In both compounds, the thiopyran ring adopts a half-chair con-

Table 1 Selected bond distances (Å) and angles (°) of 1 and 3 (Cp- C_{Fc} = Cp ring connected to the bridging carbon atom (C_{Fc}) of the diferrocenyl unit)

3B

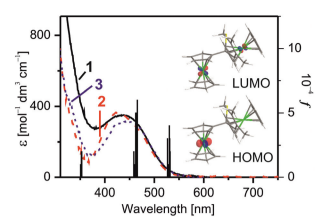
	1	3 A	3B
Bond distances			
Fe⋯Fe	5.6377(6)	5.6824(9)	5.6785(9)
$S-C_{Fc}$	1.850(2)	1.834(3)	1.832(3)
S-CH ₂	1.799(2)	1.767(3)	1.771(3)
C_{Fc} - CH_2	1.532(3)	1.530(4)	1.542(4)
C = C	1.339(3)	1.344(5)	1.331(5)
C_{Fc} – C_{Cp}	1.515(3)	1.527(4)	1.518(5)
C_{Fc} - C_{Cp}	1.517(3)	1.527(4)	1.526(4)
Fe-centroid (Cp-C _{Fc})	1.657	1.656	1.653
Fe-centroid (Cp-C _{Fc})	1.658	1.656	1.643
Fe-centroid (Cp)	1.660	1.659	1.646
Fe-centroid (Cp)	1.658	1.658	1.650
Bond angles			
S-C _{Fc} -CH ₂	107.91(13)	104.9(2)	105.3(2)
C _{Fe} -S-CH ₂	97.30(10)	100.65(16)	101.20(16
C_{Cp} - C_{Fc} - C_{Cp}	111.20(16)	112.1(2)	110.0(3)
0=S=O		117.81(15)	117.57(16
		, ,	,

formation to limit the steric repulsion between the different substituents (Fig. 2). In these carbon-bridged diferrocenyl complexes, the Fe···Fe distances are 5.6377(6) (1), 5.6824(9) (3A) and 5.6785(9) Å (3B), respectively. These values are comparable to those found in analogous carbon-bridged diferrocenyl compounds. Interestingly, the dihedral angles between the two planes of the covalently bonded cyclopentadienyl rings are quite acute in 1 (61.4°) and molecule 3A (63.1°), while in 3B this dihedral angle is normal at 76.6°.

UV-Vis spectroscopy, electrochemistry and spectroelectrochemistry

The electronic properties of the ferrocenyl-functionalised compounds 1–3 were studied by using UV-Vis, cyclic (= CV) and square wave voltammetry (= SWV) and *in situ* UV-Vis/NIR spectroelectrochemistry.

Compounds 1–3 show relatively weak absorptions in the visible region and stronger absorptions at higher energies (Fig. 3) as it is characteristic for ferrocene derivatives.^{57,58}



Dalton Transactions

Fig. 3 UV-Vis absorption spectra of 1–3 in dichloromethane at ambient temperature (1: black solid line; 2: red dashed line; 3: blue dotted line) and TD-DFT calculated electronic transitions for 1 (vertical bars; oscillator strengths are given on the y axis (right)). Contour plots of the frontier orbitals HOMO and LUMO for 1.

TD-DFT (= Time Dependent Density-Functional Theory) calculations (Fig. 3) performed for **1** predict eight lowest-energy transitions between 450 and 530 nm in which mainly Fecentred molecular orbitals of 3d-character are involved. The low energy spectral features are, thus, assigned to an unresolved series of broad overlapping bands resulting from ferrocene-centred d–d transitions. 40,59,60

The electrochemical measurements (CV and SWV) were performed under an argon atmosphere in dichloromethane solutions containing $[Bu_4N][B(C_6F_5)_4]$ (0.1 mol L^{-1}) as supporting electrolyte at a scan rate of 100 mV s⁻¹ at 25 °C. The data of the cyclic voltammetry experiments are summarised in Table 2. All potentials are referenced to the FcH/FcH⁺ redox couple. The voltammograms of 1–3 are shown in Fig. 5.

As it can be seen from Fig. 5, both ferrocenyl groups in 1–3 can be oxidised separately showing two well-resolved reversible one-electron oxidation steps. The chemical oxidation of the sulphur atom in the neighbouring position to the ferrocenyl units leads to an anodic shift of the both Fc-based oxidation processes, E_1° from 90 mV (1) over –15 mV (2) to 5 mV (3) and E_2° from 275 mV (1) over 355 mV (2) to 375 (3), respectively. This chemical oxidation process increases the group-electronegativity at the sulfur (for example, see group electronegativity: –SMe = 2.592; –SOMe = 2.841; –SO₂Me = 2.998)⁶² and hence,

Table 2 Cyclic voltammetry data of 1-3^a

Compd.	$E_1^{\circ'b}$ [mV] (ΔE_p^c [mV])	$E_2^{\circ'b}$ [mV] (ΔE_p^c [mV])	$\Delta E^{\circ\prime}{}^{d}$ [mV]
1	-90 (73)	275 (75)	365
2	-15 (75)	355 (76)	370
3	5 (73)	375 (83)	370

^a Potentials νs. FcH/FcH⁺, scan rate 100 mV s⁻¹ at glassy carbon electrode of a 1.0 mmol L⁻¹ solution in dry dichloromethane; 0.1 mol L⁻¹ [NⁿBu₄][B(C₆F₅)₄] as supporting electrolyte at 25 °C. ^b E° ' = Formal potential. ^c $\Delta E_{\rm p}$ = difference between the oxidation and the reduction potential. ^d ΔE° ' = potential difference between the two ferrocenyl-related redox processes $(E_2^{\circ}-E_1^{\circ})$.

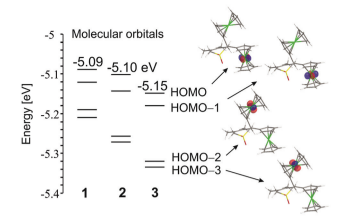


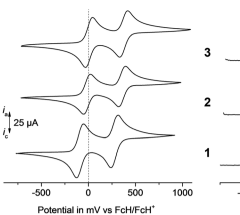
Fig. 4 DFT energies of the four lowest occupied orbitals of 1–3 and contour plots of the corresponding Kohn–Sham orbitals for 3.

reduces the electron density at the ferrocenyl groups due to the increased electron withdrawing effect. This trend is also reproduced in the results of the DFT calculations (see below). The Fe-centred occupied frontier orbitals undergo substantial stabilisation upon oxidation of the sulphur atom. For example, the HOMO energy in 3 is about 60 mV smaller than that in non-oxidised 1 (Fig. 4).

The redox separation between the 1st and the 2nd oxidation processes, however, is not affected by the different degree of the sulphur oxidation and is ca. 370 mV throughout the series. Due to the use of $[Bu_4N][B(C_6F_5)_4]^{25,63-69}$ as weakly coordination counter-ion within the electrolyte, the ion-pairing effects are minimised⁷⁰⁻⁷² and thus, the electrostatic stabilisation forces between the ferrocenyl groups are increased, when compared with diferrocenylmethane measured in $[Bu_4N][ClO_4]$ $(\Delta E^{o'}=120 \text{ mV}).^{73}$

The UV-Vis/NIR spectroelectrochemical measurements were performed in an OTTLE (= Optically Transparent Thin-Layer Electrochemistry) cell using an analyte concentration of 2.0 mmol $\rm L^{-1}$ and $\rm [Bu_4N][B(C_6F_5)_4]$ (0.1 mol $\rm L^{-1})$ as supporting electrolyte in dichloromethane (1–3) or acetonitrile (2). The UV-Vis/NIR spectra are depicted in Fig. 6 (1), 7 (2), and 8 (3). The spectrum measured in acetonitrile (2) is shown in Fig. S5.†

The appropriate compounds were oxidised by stepwise increasing the potentials (step width 25, 50 and 100 mV). Thus, the studied compounds 1–3 underwent oxidation to the mono-cationic 1^+-3^+ and di-cationic $1^{2+}-3^{2+}$ species, respectively. After complete oxidation, each sample was reduced at -200 mV to prove the reversibility of the redox processes. The resulting UV-Vis/NIR spectra were identical to those of the starting molecules. During the oxidation of 1-3 a broad band with a very low intensity ($\varepsilon_{\rm max} = 100$ L mol $^{-1}$ cm $^{-1}$) between 1000 and 3000 nm appeared (Fig. 6–8). A further increase of the potential resulted in the decrease of this band. Such a behaviour is typically observed for intervalence charge transfer (= IVCT) absorptions. 2,13 The experimental spectra can be deconvoluted into three Gaussian-shape bands assigned to an



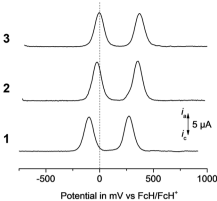


Fig. 5 Left: cyclic voltammograms of 1–3, scan rate: 100 mV s⁻¹. Right: square wave voltammograms of 1–3 in dichloromethane solutions (1.0 mmol L⁻¹) at 25 °C, supporting electrolyte 0.1 mol L⁻¹ [Bu₄N][B(C₆F₅)₄], working electrode: glassy carbon electrode (surface area 0.031 cm²).

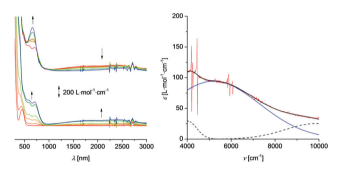


Fig. 6 Left: UV-Vis/NIR spectra of $\bf 1$ at 25 °C in dichloromethane (2.0 mmol L $^{-1}$) at rising potentials (bottom: -200 to 475 mV; top: 475 to 1200 mV vs. Ag/AgCl); supporting electrolyte [Bu₄N][B(C₆F₅)₄]. Right: deconvolution of the NIR absorptions of $\bf 1$ ⁺ using three Gaussian shaped bands determined by spectroelectrochemistry in an OTTLE cell.

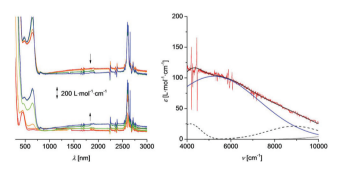


Fig. 7 Left: UV-Vis/NIR spectra of 2 at 25 °C in dichloromethane (2.0 mmol L^{-1}) at rising potentials (bottom: -200 to 375 mV; top: 375 to 1200 mV vs. Ag/AgCl); supporting electrolyte $[Bu_4N][B(C_6F_5)_4].$ Right: deconvolution of the NIR absorptions of 2^+ using three Gaussian shaped bands determined by spectroelectrochemistry in an OTTLE cell.

IVCT, a ligand field transition, and a band representing the edge to the higher energy absorptions. The sum of these three Gaussian-shaped bands fits almost exactly with the experimental spectra. The deconvolution reveals the intensity $\varepsilon_{\rm max}$, the full width-at-half-height $\Delta\nu_{1/2}$ and the $\nu_{\rm max}$ values for the IVCT component. The solvent polarity change from P=3.1 (dichloromethane) to P=5.8 (acetonitrile), 75 resulting in a shift

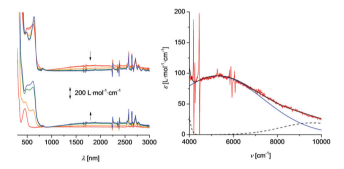


Fig. 8 Left: UV-Vis/NIR spectra of 3 at 25 °C in dichloromethane (2.0 mmol L^{-1}) at rising potentials (bottom: -200 to 375 mV; top: 375 to 1200 mV vs. Ag/AgCl); supporting electrolyte [Bu₄N][B(C₆F₅)₄]. Right: deconvolution of the NIR absorptions of 3⁺ using three Gaussian shaped bands determined by spectroelectrochemistry in an OTTLE cell.

Table 3 NIR absorption data of the IVCT band of 1^+ – 3^+ a

Compd.	$ u_{ m max} ({ m cm}^{-1}) $ $(\varepsilon_{ m max} ({ m L~mol}^{-1} { m cm}^{-1}))$	$\frac{\Delta \nu_{1/2}}{(\mathrm{cm}^{-1})}$	$(\Delta \nu_{1/2})_{\text{theo}}^b$ (cm^{-1})
1 (DCM)	5200 (100)	4950	3468
2^+ (DCM)	5250 (100)	4900	3478
2 ⁺ (ACN)	7525 (60)	7850	4169
3^{+} (DCM)	5300 (95)	4900	3512

 a Measured in dry dichloromethane (DCM) or acetonitrile (ACN) using [Bu_4N][B(C_6F_5)_4] (0.1 mol dm^-³) as supporting electrolyte at 25 °C. b Values calculated as $(\Delta\nu_{1/2})_{theo}=(2310~\nu_{max})^{1/2}$ according to the Hush relationships for weakly coupled systems. 76

of the $\nu_{\rm max}$ value from 5250 cm⁻¹ to 7525 cm⁻¹. It is remarked that strong solvatochromic shifts are expected for IVCT absorption bands being of distinct charge transfer character. Thus, the IVCT assignment of the observed NIR absorption features (Fig. 6–8 and Table 3) is further substantiated. The appearance of low energy ligand field transitions is characteristic for ferrocenyl containing compounds as for example demonstrated by UV/Vis-NIR measurements of mono ferrocenyl thiophenes.²⁷ The numerical data derived from the deconvolution procedure is summarised in Table 3. However, the data should be

Dalton Transactions

handled with care since the very small extinctions of these bands cause them to be susceptible to errors.

The $\Delta \nu_{1/2}$ values exceed the theoretical values according to the Hush model, which is a common behaviour for weakly coupled class II systems according to Robin and Day³⁵ and is caused by interactions with the respective solvent. The absorption energy is slightly blue shifted with increasing oxidation state of the sulphur atom (the ν_{max} value increases from 5200 cm⁻¹ for 1⁺ to 5300 cm⁻¹ determined for 3⁺). However, these differences are small and lie within the margin of the experimental error of our measurements. Thus, the sulphur oxidation state does not have any significant influence on the electron transfer properties of differrocenes 1⁺-3⁺. Remarkably, for the diferrocenylmethane cation (Fc₂CH₂⁺) no IVCT absorptions could be detected, while for the mixed-valent triferrocenylmethane (Fc₃CH⁺) weak charge transfer excitations (ε_{max} = 165 L mol⁻¹ cm⁻¹; $\Delta \nu_{1/2} = 3750 \text{ cm}^{-1}$; $\nu_{\text{max}} = 5900 \text{ cm}^{-1}$) were found, possessing similar characteristics as mixed-valent 1⁺-3⁺.44

Conclusions

Within this study it was shown that 2,2-diferrocenyl-4,5dimethyl-3,6-dihydro-2H-thiopyran (1) is readily available through a hetero-Diels-Alder cyclo-addition reaction of diferrocenylthioketone with 2,3-dimethyl-1,3-butadiene. Stepwise oxidation of the sulphur atom in 1 afforded the corresponding 2,2-diferrocenyl-4,5-dimethyl-3,6-dihydro-2H-thio-S-oxides: pyran-1-oxide (2, sulfoxide) and 2,2-diferrocenyl-4,5-dimethyl-3,6-dihydro-2*H*-thiopyran-1,1-dioxide (3, sulfone). Electrochemical measurements on 1-3 revealed well separated redox events related to the two ferrocenyl groups. The corresponding $E_1^{\circ\prime}$ and $E_2^{\circ\prime}$ potentials are shifted anodically in the order of 1 < 2< 3. This is accounted for the increasing electron-withdrawing effect of the adjacent S, SO, and SO₂ fragments on the ferrocenyl groups. Thus, the Fe-centred orbitals in 2 and 3 are more stabilised than in 1. A single crystal X-ray diffraction study of 1 and 3 revealed that the chemical oxidation of the sulphur atom has only a negligible influence on the overall molecule's geometry (bond lengths and angels). Nevertheless, the X-ray analyses have also shown that the Fe-Fe distances in 1 and 3 are shorter than that in Fc_2CH_2 (5.765 Å).⁷⁷ This effect is ascribed to the steric hindrance exerted by the 4,5-dimethyl-3,6-dihydro-2H-thiopyran ring. It has been also demonstrated that the cyclic 4,5-dimethyl-3,6-dihydro-2H-thiopyran group enables weak metal-metal electronic interactions in the monooxidised species 1^+-3^+ , as confirmed by the appearance of weak IVCT absorptions characteristic for mixed-valent systems (= MV). This allows to categorise cations 1^+-3^+ as weakly coupled class II MV systems according to Robin and Day. 35 Moreover, the extent of metal-metal electronic interactions in 1⁺-3⁺ does not change significantly with the oxidation state of the sulphur atom (sulphide $(1) \rightarrow$ sulfoxide $(2) \rightarrow$ sulfone (3)). In summary, our results demonstrate that sterically demanding 2H-thiopyran-derived bridges enable metal-metal electronic interactions between redox centres in mixed-valent molecular systems. In the studied MV species, most probably, the "through space" mechanism is dominantly operative.

Experimental section

General data and reagents

All reactions were carried out under an atmosphere of argon standard Schlenk techniques. Chromatographic separations were carried out using silica gel 60 (Merck, 230-400 mesh ASTM). Dichloromethane was purified by distillation from CaH2 prior to use and methanol was purified by distillation over magnesium. 2,3-Dimethyl-1,3-butadiene, m-chloroperoxybenzoic acid, 30% hydrogen peroxide and selenium dioxide were purchased from commercial suppliers and were used without further purification.

Instruments

¹H NMR (600 MHz) and ¹³C{H} NMR (150 MHz) spectra were recorded with a Bruker Avance III 600 spectrometer operating at 298 K in the Fourier transform mode. Chemical shifts are reported in δ units (ppm) using as residual CDCl₃ (¹H δ 7.26 ppm, 13 C δ 77.00 ppm) as the reference. Infrared spectra were recorded with a FTIR Nexus Nicolet apparatus. Mass spectra were recorded with a Varian 500-MS iT mass spectrometer (ESI) or with a Finnigan Mat95 mass spectrometer (EI). Microanalyses were determined by Analytical Services of the Polish Academy of the Sciences, Łódź. UV-Vis absorption spectra were recorded with a Varian Cary 300 double beam spectrometer.

DFT computations and spectroelectrochemical measurements

The geometry optimisations and electronic transition calculations were performed using density-functional theory (= DFT) and time dependent density-functional theory (= TD DFT) with Becke's three parameter functional⁷⁸ with the nonlocal Lee-Yang-Parr correlation functional (B3LYP)⁷⁹ and the standard 6-31G(d,p) basis set as implemented in the Gaussian 09 program package.80

Electrochemical measurements of 1.0 mmol L⁻¹ dichloromethane solutions of 1-3 were performed in a dried, argon purged cell at 25 °C with a Radiometer Voltalab PGZ 100 electrochemical workstation interfaced with a personal computer. Dichloromethane solutions (0.1 mol L⁻¹) containing [Bu₄N]- $[B(C_6F_5)_4]$ were used as supporting electrolyte. For the measurements a three electrode cell containing a Pt auxiliary electrode, a glassy carbon working electrode (surface area 0.031 cm^2) and an Ag/Ag⁺ ($0.01 \text{ mmol L}^{-1} \text{ [AgNO}_3\text{])}$ reference electrode fixed on a Luggin capillary was applied. The working electrode was pretreated by polishing on a Buehler microcloth first with a 1 micron and then with a 1/4 micron diamond paste. The reference electrode was constructed from a silver wire inserted into a 0.01 mmol L^{-1} [AgNO₃] and 0.1 mol L^{-1} [Bu₄N][B(C₆F₅)₄] acetonitrile solution in a Luggin capillary with a Vycor tip. This Luggin capillary was inserted into a second Luggin capillary containing a 0.1 mol L⁻¹ [Bu₄N][B(C₆F₅)₄]

dichloromethane solution and a Vycor tip. Experiments under the same conditions showed that all reduction and oxidation potentials were reproducible within 5 mV. Experimental potentials were referenced against an Ag/Ag⁺ reference electrode but the presented results are referenced against ferrocene as an internal standard as required by IUPAC. 61 To achieve this, each experiment was repeated in the presence of 1 mmol L^{-1} decamethylferrocene (= Fc^*). Data were processed on a Microsoft Excel worksheet to set the formal reduction potentials of the FcH/FcH⁺ couple to 0.0 V. Under our conditions the Fc^*/Fc^{*+} couple was at -619 mV vs. FcH/FcH^{+} ($\Delta E_{p} = 60 \text{ mV}$), while the FcH/FcH^{+} couple itself was at 220 mV vs. Ag/Ag^+ ($\Delta E_p = 61$ mV). Spectroelectrochemical UV-Vis/NIR measurements of 2.0 mmol L⁻¹ solutions of 1-3 in dichloromethane (1-3) or acetonitrile (2) containing 0.1 mol L^{-1} of $[Bu_4N][B(C_6F_5)_4]$ as the supporting electrolyte were performed in an OTTLE (OTTLE = Optically Transparent Thin-Layer Electrochemistry)⁷⁴ cell with a Varian Cary 5000 spectrophotometer at 25 °C. The values obtained by deconvolution could be reproduced within $\varepsilon_{\rm max}$, 100 L mol⁻¹ cm⁻¹; $\nu_{\rm max}$,

Synthesis of 2,2-diferrocenyl-4,5-dimethyl-3,6-dihydro-2*H*-thiopyran (1)

50 cm⁻¹ and $\Delta \nu_{1/2}$, 50 cm⁻¹.

A mixture of diferrocenyl thioketone (399 mg, 0.96 mmol) and 2,3-dimethyl-1,3-butadiene (8.5 mL) was stirred in a tightly closed glass-tube for 50 h at 75 °C. Afterwards, the reaction mixture was evaporated to dryness and the thus obtained solid was subjected to column chromatography on SiO_2 (chloroform–hexane, ratio 1/1 (v/v)). Crystallization from chloroform–hexane gave pure 1 as red crystals in a 65% yield (309 mg).

¹H NMR (600 MHz, CDCl₃): δ = 4.26 (bs, 2H, C₅H₄), 4.20 (bs, 2H, C₅H₄), 4.13 (bs, 2H, C₅H₄), 4.11 (bs, 2H, C₅H₄), 4.07 (s, 10H, C₅H₅), 3.00 (s, 2H, CH₂), 2.64 (s, 2H, CH₂), 1.85 (s, 3H, CH₃), 1.74 (s, 3H, CH₃). ¹³C NMR (150 MHz, CDCl₃): δ = 126.6, 124.3, 97.1, 69.0, 66.9, 66.8, 66.7, 66.4, 44.5, 43.3, 31.3, 20.4, 19.4. FTIR (KBr): 3088, 2989, 2911, 2872, 1628, 1443, 1409, 1264, 1106, 1032, 1000, 824, 483 cm⁻¹. MS (ESI): m/z = 496 (M⁺). MS (EI, 70 eV): m/z = 496 (M⁺). HRMS: m/z = 496.0614 (Calc. for C₂₇H₂₈SFe₂: 496.0610). Anal. Calcd for: C₂₇H₂₈SFe₂: C, 65.35; H, 5.69; S, 6.46%. Found: C, 65.10; H, 5.57; S, 6.72%.

Synthesis of 2,2-diferrocenyl-4,5-dimethyl-3,6-dihydro-2*H*-thiopyran-1-oxide (2)

Hydrogen peroxide (30%, 71 mg) and selenium dioxide (72 mg, 0.65 mmol) in water (1 mL) were subsequently added to a stirred solution of 1 (312 mg, 0.63 mmol) in methanol (10 mL) at 0 °C. The resulting mixture was stirred at 0 °C for 5 min and then for an additional 5 min at ambient temperature. The reaction was quenched with water (\sim 7 mL) and extracted with dichloromethane. The organic layer was washed with brine, separated and dried over anhydrous MgSO₄ and then all volatile materials were evaporated. The residue was subjected to chromatography on SiO₂ (diethyl ether–hexane, ratio 3/1 (v/v)) to give pure sulfoxide 2 as red crystals in 87% yield (281 mg).

¹H NMR (600 MHz, CDCl₃): δ = 4.90–4.89 (m, 1H, C₅H₄), 4.36–4.35 (m, 1H, C₅H₄), 4.32–4.31 (m, 1H, C₅H₄), 4.30–4.29 (m, 1H, C₅H₄), 4.28 (s, 5H, C₅H₅), 4.27–4.26 (m, 1H, C₅H₄), 4.22–4.21 (m, 1H, C₅H₄), 4.05–4.04 (m, 1H, C₅H₄), 4.00–3.99 (m, 1H, C₅H₄), 3.88 (s, 5H, C₅H₅), 3.18 (d, $J_{\rm H,H}$ = 18.6 Hz, 1H, CH₂), 3.01 (d, $J_{\rm H,H}$ = 18.0 Hz, 1H, CH₂), 2.97 (d, $J_{\rm H,H}$ = 15.6 Hz, 1H, CH₂), 2.50 (d, $J_{\rm H,H}$ = 15.6 Hz, 1H, CH₂), 1.84 (s, 3H, CH₃), 1.68 (s, 3H, CH₃). ¹³C NMR (150 MHz, CDCl₃): δ = 127.0, 118.6, 94.3, 87.1, 70.1, 69.8, 69.2, 68.9, 68.7, 68.4, 66.9, 66.7, 66.2, 66.1, 59.4, 51.1, 41.6, 20.0, 19.4. FTIR (KBr): 3089, 2917, 2891, 2857, 1629, 1406, 1104, 1053 (s, S=O), 1031, 1000, 824, 487 cm⁻¹. MS (ESI): m/z = 535 (MNa⁺), 513 (MH⁺). MS (EI, 70 eV): m/z = 512 (M⁺). HRMS: m/z = 512.0558 (Calc. for C₂₇H₂₈OSFe₂: 512.0560). Anal. Calcd for: C₂₇H₂₈OSFe₂: C, 63.30; H, 5.51; S, 6.26%. Found: C, 63.23; H, 5.72; S, 6.27%.

Synthesis of 2,2-diferrocenyl-4,5-dimethyl-3,6-dihydro-2*H*-thiopyran-1,1-dioxide (3)

m-Chloroperoxybenzoic acid (= MCPBA) (45 mg, 0.26 mmol) was added to a stirred solution of sulphoxide 2 (100 mg, 0.19 mmol) in dichloromethane (20 mL) at -20 °C. The resulting reaction mixture was stirred at -20 °C for 2 h and then for an additional 24 h at ambient temperature. Afterwards, the $2^{\rm nd}$ portion of MCPBA (45 mg, 0.26 mmol) was added and the mixture was stirred at ambient temperature for another 24 h. Subsequently, a saturated NaHCO3 solution was added in a single portion and the resulting mixture was extracted with dichloromethane. The organic layer was washed with brine, separated and dried over anhydrous MgSO4. Then all volatile materials were evaporated. The residue was subjected to preparative TLC on SiO2 (diethyl ether–hexane, ratio 1/1 (v/v) as eluent). Sulfone 3 was obtained as a red solid in 21% (21 mg) yield.

¹H NMR (600 MHz, CDCl₃): δ = 4.47 (bs, 2H, C₅H₄), 4.24 (s, 4H, C₅H₄), 4.21–4.20 (pq, $J_{\rm H,H}$ = 1.98 Hz, 1.74 Hz, 2H, C₅H₄), 4.13 (s, 10H, C₅H₅), 3.32 (s, 2H, CH₂), 3.06 (s, 2H, CH₂), 1.91 (s, 3H, CH₃), 1.69 (s, 3H, CH₃). ¹³C NMR (150 MHz, CDCl₃): δ = 127.0, 119.9, 89.0, 69.6, 68.7, 67.6, 67.5, 67.2, 62.5, 51.6, 44.5, 19.9, 19.6. FTIR (KBr): 3103, 2918, 2857, 1629, 1305 (s, SO₂), 1121(s, SO₂), 820, 480 cm⁻¹. MS (ESI): m/z = 551 (MNa⁺), 528 (M⁺). MS (EI, 70 eV): m/z = 528 (M⁺), 464 (M⁺ – SO₂), 462 (M⁺ – H₂SO₂). HRMS: m/z = 528.0510 (Calc. for C₂₇H₂₈O₂SFe₂: 528.0509). Anal. Calcd for: C₂₇H₂₈O₂SFe₂: C, 61.39; H, 5.34; S, 6.07%. Found: C, 61.20; H, 5.48; S, 5.91%.

Single-crystal X-ray structure analysis of 1 and 3

Red crystals of 1 were obtained by slow evaporation of a chloro-form–hexane solution containing 1, while red crystals of 3 were grown by the slow evaporation of a diethyl ether–hexane solution containing 3 at ambient temperature. Data were collected with a Stoe Image Plate Diffraction system equipped with a ϕ circle goniometer using Mo K_{α} graphite monochromatic radiation (λ = 0.71073 Å) with ϕ range 0–200°. The structures were solved by direct methods applying the program SHELXS-97, while the refinement and all further calculations were carried out with SHELXL-97. ^{82,83} The hydrogen atoms

Dalton Transactions Paper

were included in calculated positions and treated as riding atoms using the SHELXL default parameters, except for the N-H hydrogen atoms which were located on the Fourier difference map and refined. The non-hydrogen atoms were refined anisotropically using weighted full-matrix least-square on F^2 . Crystallographic details for 1 and 3 are summarized in Table S1 (see ESI†).

CCDC 1031233 (1) and 1031234 (3) contain the supplementary crystallographic data for this paper.

Acknowledgements

Authors (K. K. and G. M.) thank the National Science Centre (Poland) for financial support (Project Maestro-3; Dec-2012/06/A/ST5/00219) and R. C. thanks the German Federal Ministry of Education and Research (BMBF) for support. The support from the German Academic Exchange Service (DAAD) in the framework of the exchange program "Ostpartnerschaften" is highly appreciated.

Notes and references

- 1 S. D. Glover, J. C. Goeltz, B. J. Lear and C. P. Kubiak, *Eur. J. Inorg. Chem.*, 2009, 5, 585–594.
- 2 D. M. D'Alessandro and F. R. Keene, *Chem. Rev.*, 2006, **106**, 2270–2298.
- 3 C. Lapinte, J. Organomet. Chem., 2008, 693, 793-801.
- 4 A. K. Diallo, C. Absalon, J. Ruiz and D. Astruc, J. Am. Chem. Soc., 2011, 133, 629–641.
- 5 A. Ceccon, S. Santi, L. Orian and A. Bisello, *Coord. Chem. Rev.*, 2004, 248, 683–724.
- 6 P. Aguirre-Etcheverry and D. O'Hare, Chem. Rev., 2010, 110, 4839–4864.
- 7 R. F. Winter, Organometallics, 2014, 33, 4517-4536.
- 8 A. Hildebrandt and H. Lang, *Organometallics*, 2013, 32, 5640–5653.
- 9 K. Costuas and S. Rigaut, *Dalton Trans.*, 2011, **40**, 5643–5658.
- 10 T. Kienz, C. Förster and K. Heinze, *Organometallics*, 2014, 33, 4803–4812.
- 11 K. Kowalski, M. Linseis, R. F. Winter, M. Zabel, S. Záliš, H. Kelm, H.-J. Krüger, B. Sarkar and W. Kaim, *Organometallics*, 2009, **28**, 4196–4209.
- 12 M. Linseis, S. Záliš, M. Zabel and R. F. Winter, *J. Am. Chem. Soc.*, 2012, **134**, 16671–16692.
- 13 F. Paul and C. Lapinte, Coord. Chem. Rev., 1998, 178–180, 431–509.
- 14 M. D. Ward, Chem. Soc. Rev., 1995, 121-134.
- 15 M. Ratner and J. Jortner, *Molecular Electronics*, Malden, MA, 1997.
- 16 Electron Transfer in Chemistry and Biology, ed. V. Balzani, Wiley-VCH, Weinheim, 2001.
- 17 Electron and Proton Transfer in Chemistry and Biology, ed. A. Müller, H. Ratajczak, W. Junge and E. Dieman, Elsevier, New York, 1992.

18 H. B. Gray and J. R. Winkler, Annu. Rev. Biochem., 1996, 65, 537–561.

- 19 A. K. Diallo, C. Absalon, J. Ruiz and D. Astruc, *J. Am. Chem. Soc.*, 2011, **133**, 629–641.
- 20 M. Pichlmaier, R. F. Winter, M. Zabel and S. Záliš, J. Am. Chem. Soc., 2009, 131, 4892–4903.
- 21 W. E. Geiger, Organometallics, 2007, 26, 5738-5765.
- 22 A. Hildebrandt, D. Schaarschmidt, R. Claus and H. Lang, *Inorg. Chem.*, 2011, **50**, 10623–10632.
- 23 J. M. Speck, M. Korb, T. Rüffer, A. Hildebrandt and H. Lang, *Organometallics*, 2014, 33, 4813–4823.
- 24 S. W. Lehrich, A. Hildebrandt, T. Rüffer, M. Korb, P. J. Low and H. Lang, *Organometallics*, 2014, 33, 4836–4845.
- 25 D. Miesel, A. Hildebrandt, M. Korb, P. J. Low and H. Lang, *Organometallics*, 2013, 32, 2993–3002.
- 26 U. Pfaff, A. Hildebrandt, M. Korb and H. Lang, *Polyhedron*, 2015, **86**, 2–9.
- 27 J. M. Speck, R. Claus, A. Hildebrandt, T. Rüffer, E. Erasmus, L. van As, J. C. Swarts and H. Lang, *Organo-metallics*, 2012, 31, 6373–6380.
- 28 K. Kaleta, F. Strehler, A. Hildebrandt, T. Beweries, P. Arndt, T. Rüffer, A. Spannenberg, H. Lang and U. Rosenthal, *Chem. Eur. J.*, 2012, **18**, 12672–12680.
- 29 A. Hildebrandt, U. Pfaff and H. Lang, *Rev. Inorg. Chem.*, 2011, 31, 111–141.
- 30 A. Hildebrandt and H. Lang, *Dalton Trans.*, 2011, **40**, 11831–11837.
- 31 K. Kaleta, A. Hildebrandt, F. Strehler, P. Arndt, H. Jiao, A. Spannenberg, H. Lang and U. Rosenthal, *Angew. Chem., Int. Ed.*, 2011, **50**, 11248–11252.
- 32 A. Hildebrandt, D. Schaarschmidt and H. Lang, *Organometallics*, 2011, **30**, 556–563.
- 33 C. G. Allen and S. N. Hush, *Prog. Inorg. Chem.*, 1967, 8, 357–339.
- 34 P. Mücke, M. Zabel, R. Edge, D. Collison, S. Clément, S. Záliš and R. F. Winter, *J. Organomet. Chem.*, 2011, **696**, 3186–3197.
- 35 M. B. Robin and P. Day, *Adv. Inorg. Chem. Radiochem.*, 1967, **10**, 247–422.
- 36 T.-Y. Dong, C.-H. Huang, C.-K. Chang, H.-C. Hsieh, S.-M. Peng and G.-H. Lee, *Organometallics*, 1995, **14**, 1776– 1785.
- 37 S. C. Jones, S. Barlow and D. O'Hare, *Chem. Eur. J.*, 2005, **11**, 4473–4481.
- 38 J. C. Kotz, C. L. Nivert, J. M. Lieber and R. C. Reed, *J. Organomet. Chem.*, 1975, **91**, 87–95.
- 39 D. C. O'Connor Salazar and D. O. Cowan, J. Organomet. Chem., 1991, 408, 227–231.
- 40 A.-C. Ribou, J.-P. Launay, M. L. Sachtleben, H. Li and C. W. Spangler, *Inorg. Chem.*, 1996, 35, 3735–3740.
- 41 T.-Y. Dong, T.-Y. Lee, S.-H. Lee, G.-H. Lee and S.-M. Peng, Organometallics, 1994, 13, 2337–2348.
- 42 R. J. Weeb, S. J. Geib, D. L. Staley, A. L. Rheingold and D. N. Hendrickson, J. Am. Chem. Soc., 1990, 112, 5031.
- 43 R. Rulkens, A. J. Lough, I. Manners, S. R. Lovelace, C. Grant and W. Geiger, *J. Am. Chem. Soc.*, 1996, **118**, 12683–12695.

44 F. Delgado-Pena, D. R. Talham and D. O. Cowan, J. Organo-

- met. Chem., 1983, 253, C43-C46.
- 45 U. Pfaff, G. Filipczyk, A. Hildebrandt, M. Korb and H. Lang, *Dalton Trans.*, 2014, **43**, 16310–16321.
- 46 G. Filipczyk, A. Hildebrandt, U. Pfaff, M. Korb, T. Rüffer and H. Lang, *Eur. J. Inorg. Chem.*, 2014, 4258–4262.
- 47 U. Pfaff, A. Hildebrandt, D. Schaarschmidt, T. Hahn, S. Liebing, J. Kortus and H. Lang, *Organometallics*, 2012, 31, 6761–6771.
- 48 M. Lohan, F. Justaud, H. Lang and C. Lapinte, *Organometallics*, 2012, **31**, 3565–3574.
- 49 M. Lohan, F. Justaud, T. Roisnel, P. Ecorchard, H. Lang and C. Lapinte, *Organometallics*, 2010, **29**, 4804–4817.
- 50 P. Denifl and B. Bildstein, *J. Organomet. Chem.*, 1993, **453**, 53–59.
- 51 D. Enders, E. A. Jonas and T. Klumpen, *Eur. J. Org. Chem.*, 2009, 2149–2162.
- 52 S. D. Larsen, P. V. Fisher, B. E. Libby, R. M. Jensen, S. A. Mizsak and W. Watt, *J. Org. Chem.*, 1996, 61, 4725– 4738.
- 53 J. Drabowicz and M. Mikołajczyk, Synthesis, 1978, 758-759.
- 54 V. A. Petrov, S. Lustig and W. Marshall, J. Fluorine Chem., 2007, 128, 1227–1234.
- 55 V. A. Petrov and W. Marshall, *J. Fluorine Chem.*, 2007, **128**, 729–735.
- 56 R.-J. Xie, L.-M. Han, Q.-L. Suo, H.-L. Hong and M.-H. Luo, J. Coord. Chem., 2010, 63, 1700–1710.
- 57 Y. Yamaguchi, W. Ding, C. T. Sanderson, M. L. Borden, M. J. Morgan and C. Kutal, *Coord. Chem. Rev.*, 2007, 251, 515–524.
- 58 S. Barlow and D. O'Hare, Chem. Rev., 1997, 97, 637-670.
- 59 D. M. Duggan and D. N. Hendrickson, *Inorg. Chem.*, 1975, 14, 955–970.
- 60 K. Kowalski, Ł. Szczupak, J. Skiba, O. S. Abdel-Rahman, R. F. Winter, R. Czerwieniec and B. Therrien, *Organometallics*, 2014, 33, 4697–4705.
- 61 G. Gritzner and J. Kuta, *Pure Appl. Chem.*, 1984, **56**, 461–466.
- 62 N. Inamoto and S. Masuda, *Chem. Lett.*, 1982, 11, 1003-1006.
- 63 H. J. Gericke, N. I. Barnard, E. Erasmus, J. C. Swarts, M. J. Cook and M. A. S. Aquino, *Inorg. Chim. Acta*, 2010, 363, 2222–2232.
- 64 E. Fourie, J. C. Swarts, D. Lorcy and N. Bellec, *Inorg. Chem.*, 2010, 49, 952–959.
- 65 J. C. Swarts, A. Nafady, J. H. Roudebush, S. Trupia and W. E. Geiger, *Inorg. Chem.*, 2009, 48, 2156–2165.
- 66 V. N. Nemykin, G. T. Rohde, C. D. Barrett, R. G. Hadt, J. R. Sabin, G. Reina, P. Galloni and B. Floris, *Inorg. Chem.*, 2010, 49, 7497–7509.

- 67 V. N. Nemykin, G. T. Rohde, C. D. Barrett, R. G. Hadt, C. Bizzarri, P. Galloni, B. Floris, I. Nowik, R. H. Herber, A. G. Marrani, R. Zanoni and N. M. Loim, *J. Am. Chem. Soc.*, 2009, 131, 14969–14978.
- 68 E. A. Poppitz, A. Hildebrandt, M. Korb and H. Lang, J. Organomet. Chem., 2014, 752, 133–140.
- 69 U. Pfaff, A. Hildebrandt, D. Schaarschmidt, T. Rüffer, P. J. Low and H. Lang, *Organometallics*, 2013, 32, 6106–6117.
- 70 W. E. Geiger and F. Barrière, Acc. Chem. Res., 2010, 43, 1030–1039.
- 71 F. Barrière and W. E. Geiger, J. Am. Chem. Soc., 2006, 128, 3980–3989.
- 72 F. Barrière, N. Camire, W. E. Geiger, U. T. Mueller-Westerhoff and R. Sanders, *J. Am. Chem. Soc.*, 2002, **124**, 7262–7263.
- 73 G. Ferguson, C. Glidewell, G. Opromolla, C. M. Zakaria and P. Zanello, J. Organomet. Chem., 1996, 517, 183–190.
- 74 M. Krejcik, M. Danek and F. Hartl, *J. Electroanal. Chem.*, 1991, 317, 179–187.
- 75 L. R. Snyder, J. Chromatogr. Sci., 1978, 16, 223-234.
- 76 N. S. Hush, Electrochim. Acta, 1968, 13, 1005-1023.
- 77 R.-J. Xie, L.-M. Han, N. Zhu, H.-L. Hong, Q.-L. Suo and C.-L. Ke, *Asian J. Chem.*, 2013, 25, 197–201.
- 78 A. D. Becke, J. Chem. Phys., 1993, 98, 5648-5652.
- 79 C. Lee, W. Yang and R. G. Parr, *Phys. Rev. B: Condens. Matter*, 1988, 37, 785–789.
- 80 M. J. Frisch, G. W. Trucks, H. B. Schlegel, G. E. Scuseria, M. A. Robb, J. R. Cheeseman, G. Scalmani, V. Barone, B. Mennucci, G. A. Petersson, H. Nakatsuji, M. Caricato, X. Li, H. P. Hratchian, A. F. Izmaylov, J. Bloino, G. Zheng, J. L. Sonnenberg, M. Hada, M. Ehara, K. Toyota, R. Fukuda, J. Hasegawa, M. Ishida, T. Nakajima, Y. Honda, O. Kitao, H. Nakai, T. Vreven, J. A. Montgomery, J. E. Peralta Jr., F. Ogliaro, M. Bearpark, J. J. Heyd, E. Brothers, K. N. Kudin, V. N. Staroverov, R. Kobayashi, J. Normand, K. Raghavachari, A. Rendell, J. C. Burant, S. S. Iyengar, J. Tomasi, M. Cossi, N. Rega, J. M. Millam, M. Klene, J. E. Knox, J. B. Cross, V. Bakken, C. Adamo, J. Jaramillo, R. Gomperts, R. E. Stratmann, O. Yazyev, A. J. Austin, R. Cammi, C. Pomelli, J. W. Ochterski, R. L. Martin, K. Morokuma, V. G. Zakrzewski, G. A. Voth, P. Salvador, J. J. Dannenberg, S. Dapprich, A. D. Daniels, Ö. Farkas, J. B. Foresman, J. V. Ortiz, J. Cioslowski and D. J. Fox, Gaus-
- 81 A. Nafady and W. E. Geiger, *Organometallics*, 2008, 27, 5624–5631.

sian 09W, Version 8.0, Gaussian Inc., Wallingford, CT, 2009.

- 82 G. M. Sheldrick, Acta Crystallogr., Sect. A: Fundam. Crystallogr., 2008, 64, 112–122.
- 83 L. J. Farrugia, J. Appl. Crystallogr., 1997, 30, 565.