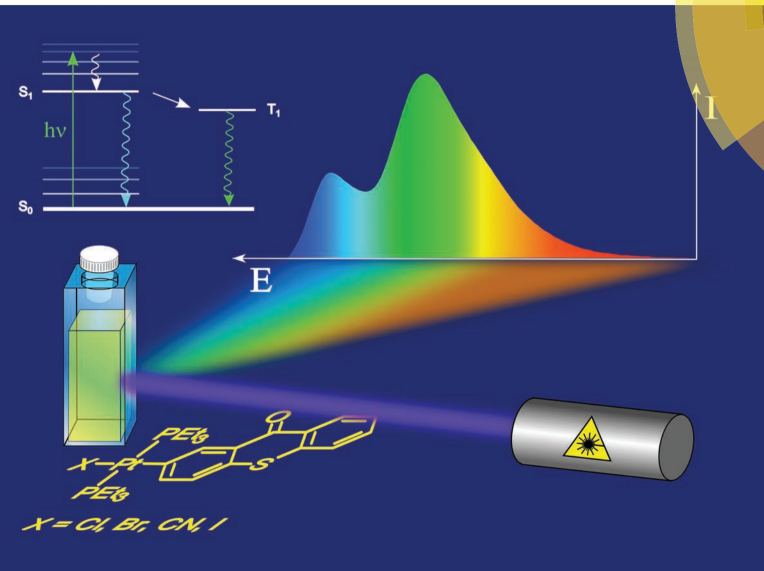
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Dual ligand-based fluorescence and phosphorescence emission at room temperature from platinum thioxanthonyl complexes†

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New square-planar platinum(II) complexes of the type trans-[Pt(PEt₃)₂(Tx)(X)] (X = Br, Cl, I or CN) bearing a σ -bonded thioxanthon-2-yl (Tx) ligand have been prepared and characterised by X-ray crystallography, cyclic voltammetry, and by NMR and electronic absorption and luminescence spectroscopy. The ligand X hardly influences the electronic transitions, which indicates that the relevant molecular orbitals are largely confined to the Pt-Tx chromophore. In agreement with TD-DFT calculations the energetically lowest electronic transition is assigned as the Tx-centred $\pi \rightarrow \pi^*$ HOMO \rightarrow LUMO excitation. All four complexes display dual emission from the σ -bonded Tx ligand at ca. 450 nm and at ca. 510 nm, which are assigned as fluorescence originating from the $^{1}\pi^*$ -state and as phosphorescence originating from the $^{3}\pi^*$ -state, respectively. The phosphorescence quantum yield increases with increasing σ -donor strength of the ligand X and reaches a uniquely high value of 18.8% for the chlorido complex **Pt-Cl**. Switching-on of Tx phosphorescence emission by the Pt(PEt₃)₂(X) fragment goes along with a reduction of the lifetime of the Tx triplet state from several ms in purely organic derivatives to ca. 2 μ s in the complexes.

Introduction

Within the last two decades, complexes of mainly the heavy transition metals have been the focus of intense research activities devoted to the fields of light emitting devices, ¹⁻⁶ bioimaging, ^{5,7-9} sensing ^{5,10-14} or sensitisation. ¹⁵⁻¹⁷ One major advantage of metal complexes when compared to purely organic emitters is their ability to trigger long-lived phosphorescence at ambient temperature. Platinum, in particular, is a widely applied metal in this field because its heavy atom effect, which emanates from its large spin-orbit coupling constant, promotes high intersystem crossing (ISC) efficiencies. ¹⁸ The utilisation of triplet emitters in organic light emitting devices can theoretically lead to a fourfold increase in efficiency compared to the 25% efficiency limit of organic fluorescent compounds. ^{19,20}

Fachbereich Chemie der Universität Konstanz, Universitätsstraße 10, D-78457 Konstanz, Germany. E-mail: rainer.winter@uni-konstanz.de † Electronic supplementary information (ESI) available: Depictions of the ¹H, ¹³C { ¹H}, ³¹P{ ¹H} and ¹⁹⁵Pt NMR spectra of the complexes; crystal and refinement data as well as ORTEPs and packing diagrams of complexes Pt-Cl and Pt-CN, graphical representations of exemplary cyclic voltammograms and electronic absorption and emission spectra, Tables with structure parameters for the optimised geometries of the complexes, energies and graphical representations of the crucial MOs of the complexes and TD-DFT calculated transitions. CCDC 1017997 (Pt-Cl) and 1017999 (Pt-CN). For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/c4dt02410a

In practice, the theoretical limit of unitary quantum efficiency is, however, rarely approached. One possible deactivation pathway for the excited states of square-planar platinum(II) complexes involves distortion towards a more tetrahedral geometry by a twisting of the plane of two donors with respect to that of the other two (the so-called D_{2d} distortion). This pathway is particularly relevant for complexes with only mono- or bidentate ligands. The main reason for low quantum yields and hence increased deactivation rates of emissive states is, however, the thermal activation of non-emissive excited metal d-states which are energetically above the emissive triplet states. A ligand field-induced destabilisation of these excited metal d-states will thus lead to increased quantum yields. 18,21 Much effort has therefore been directed to introducing strong-field ligands into the coordination sphere of such complexes. Prominent examples are cyclometalating ligands such as the ubiquitous N,C- or N,N,C-chelating arylpyridines or -bipyridines, 18,22-39 simple or cyclometalating carbenes and bis(carbenes), 3,6,36,37,40-45 or σ-bonded alkynyl ligands.46-51 Particularly abundant are trans-bis(alkynyl)bis-(trialkylphosphine)platinum(II) complexes $(C \equiv CR)_2(PR_3)_2$. Such complexes have, for example been used as constituents of luminescent coordination oligomers and polymers featuring platinum atoms in the conjugated main chain⁵⁴⁻⁶³ or for two-photon absorption (TPA).⁶⁴ Other representatives have organic dyes as the substituents at the alkynyl ligands. These complexes exhibit intense ligand-based

100 - 140 ms

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 $\pi \rightarrow \pi^*$ -transitions and usually emit from ligand-centred π^* -states with moderate fluorescence or phosphorescence quantum yields in fluid solution temperature. 49,53,65-69 Again, ISC to a ligand-based excited triplet state is triggered by the large heavy-atom effect of the Pt(II) coordination centre. In rare cases, where the excited states are mainly localised at those ligands and spin-orbit coupling is attenuated by small Pt (5d) contribution to the relevant frontier molecular orbitals, dual emission from the 1L and ³L or ³MLCT excited states has been observed. ^{70–72}

σ-Bonded aryl ligands exert an even stronger ligand-field splitting than alkynyl ones. Despite the large and rapidly growing number of complexes featuring the above-mentioned cyclometalating arylpyridine²⁴ or arylphosphine⁷³ ligands, there are relatively few examples of luminescent Pt(II) complexes possessing simple, non-chelating σ-bonded aryl ligands.74-76 Most of these complexes show exclusively aryl ligand-centred emission from the $^1\pi^*$ -state with lifetimes in the ns regime⁷⁷ or emit with low quantum yields from the $^{3}\pi^{*}$ -state. Notable exceptions are trans-Pt(PEt₃)₂Br-complexes featuring the anthracen-9-yl ligand or some of its brominated derivatives of Sharp et al., which are formed by oxidative addition of the respective 9-bromoanthracenes to Pt(PEt₃)₄. These complexes were found to undergo P-type delayed fluorescence with quantum yields of close to 90% and lifetimes of a few microseconds.79 This finding clearly indicates that a triplet state is involved in the radiative deactivation process of the initially generated excited state and that the Pt(PEt₃)₂Br fragment is capable of strongly accelerating the ISC rate from the ${}^{1}\pi^{*}$ - to the ${}^{3}\pi^{*}$ -state. In contrast, attaching the Pt(PEt₃)₂Brfragment to the 3-position of perylene or the 9-position of perylene-3,4-dicarboximide (PMI) does not lead to the expected fast ISC, but promotes ligand-based fluorescence with quantum yields of 70 or 30%, respectively. Further modification of these complexes by substituting the bromido ligand by either cyanido, thiocyanato or neutral ligands such as 4-methylpyridine or butylisocyanide hardly affects the quantum yields in the case of the perylene complexes (70-80%) whereas those for the PMI analogues were found to vary from 30% for the neutral complexes to 78% for the cationic isocyanide derivative.80

The favourable photophysical properties of thioxanthones have prompted us to prepare and to investigate the complexes trans-Pt(PEt₃)₂(X)(thioxanthonyl) (X = Cl, Br, I or CN) bearing the σ-bonded thioxanthon-2-yl ligand. The ready formation of the $3\pi\pi^*$ and $3\pi\pi^*$ -states of aromatic carbonyl compounds after photoexcitation into the $^1\pi \rightarrow \pi^*$ - and $^1n \rightarrow \pi^*$ -bands $^{81-84}$ renders thioxanthone derivatives useful sensitisers 85,86 or initiators for radical photopolymerisations of olefins.^{87–93} The excited states of thioxanthones involve two singlet states S₁ $(^{1}\pi\pi^{*})$ and S_{2} $(^{1}n\pi^{*})$ and two corresponding triplet states T_{1} $(^3\pi\pi^*)$ and T_2 $(^3n\pi^*)$, which readily equilibrate (Fig. 1). 94,95 Apolar solvents decrease the energy gap between the two singlet states due to either a stabilisation of the 1 n π^* -state or a destabilisation of the ${}^{1}\pi\pi^*$ -state. The energetical proximity of these two states leads to strong vibronic coupling between

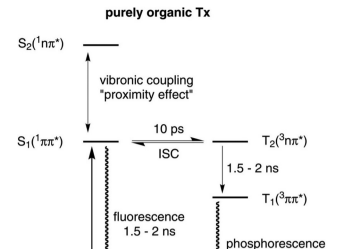


Fig. 1 Qualitative Jablonski diagram illustrating the photophysical behaviour of thioxanthones.

them. This so-called proximity effect is responsible for increasing the non-radiative deactivation rate as well as increasing the fluorescence energy. Furthermore, this effect also lowers the quantum yield for triplet formation.⁹⁶ Two principal pathways for ISC in thioxanthones were found: (a) a very fast direct mechanism leading from the ${}^{1}n\pi^{*}$ - to the ${}^{3}\pi\pi^{*}$ -state and (b) a slow indirect mechanism leading from the $^{1}\pi\pi^{*}$ - to the $^{3}\pi\pi^{*}$ state with the ³nπ*-state as the intermediate. ^{95,97} Although ISC of thioxanthones is rather efficient, they mainly emit through fluorescence because the T1 state with a lifetime of 100 to 140 ms is so long-lived that radiationless deactivation pathways become dominant. Phosphorescence is usually observed in glassy matrices at 77 K but only rarely in fluid solution with much lower quantum yields. 89,90,97 It was therefore our hope that attachment of the {Pt(PEt₃)₂X}-fragment to the thioxanthonyl core should further increase the ISC rate and hence open a pathway for radiative decay of the T1 to the S0 state. We here report our findings on the complexes Pt-Cl to Pt-CN of Scheme 1, including the observation of dual fluorescence and phosphorescence emissions from the thioxanthone chromophore.

Results and discussion

Synthesis

Sn

Oxidative addition of 2-bromo-9H-thioxanthen-9-one (Br-Tx) to Pt(PEt₃)₄ gave the trans complex Pt-Br (Scheme 1). Monitoring the progress of the reaction via ¹H and ³¹P NMR spectroscopy provided no evidence for initial formation of the kinetically favoured cis-isomer at room temperature. Subsequent abstraction of the bromido ligand of Pt-Br with AgBF₄ in methanol

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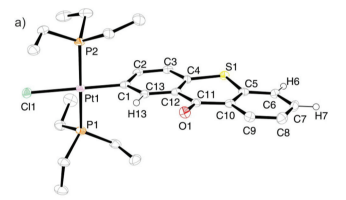
Scheme 1 Synthesis of complexes Pt-Br, Pt-CN, Pt-Cl, and Pt-I.

and addition of NaCl or NaI gave the halogenido complexes **Pt–Cl** and **Pt–I**, respectively. The analogous cyanido complex **Pt–CN** was similarly obtained by treatment of **Pt–Br** with AgCN. All complexes were purified by recrystallisation from toluene or acetone and were obtained in yields of 63 to 89%. We note that all *trans*-XPt(PEt₃)₂(Tx) complexes (X = Br, Cl, I or CN) are much more soluble in polar organic solvents than **Br–Tx**.

Attempts to purify **Pt-Br** by chromatography on basic alumina with CH₂Cl₂ as the eluent resulted in halide exchange, putatively *via* oxidative addition of CH₂Cl₂ and reductive elimination of CH₂BrCl to form complex **Pt-Cl.**^{98,99} A second possible reaction sequence is the bromide abstraction by alumina resulting in a cationic complex that subsequently reacts with CH₂Cl₂ to form **Pt-Cl.** When basic alumina was added to CD₂Cl₂ solutions of **Pt-Br** that were kept in the dark, exchange of the bromido by the chlorido ligand was much slower and occurred only over a period of two weeks as probed by ³¹P NMR spectroscopy.

Single-crystal X-ray diffraction studies of Pt-Cl and Pt-CN

Slow evaporation of concentrated dichloromethane solutions of **Pt-I**, **Pt-CN**, **Pt-Cl** and **Pt-Br** gave single crystals in each case. While data sets of **Pt-I** and **Pt-Br** were unsatisfactory, needle shaped single-crystals of **Pt-Cl** and **Pt-CN** provided data sets of sufficient quality to allow for successful refinement. The ORTEPs of the structures are shown in Fig. 2a for **Pt-Cl** and in Fig. 2b for **Pt-CN** while details of the crystallographic measurement, data collection and refinement are provided as Tables S1 and S3 of the ESI.† Complex **Pt-Cl** crystallises in the monoclinic space group $P2_1/c$ and **Pt-CN** in the triclinic space group $P\overline{1}$. The Pt atoms of both complexes are in almost ideal square-planar coordination environments with bond angles of 86.67(4) to $93.52(15)^{\circ}$ and angle sums of



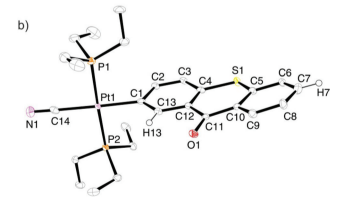


Fig. 2 ORTEPs of Pt-Cl (a) and Pt-CN (b); ellipsoids are drawn at a 40% probability level. H atoms are omitted for clarity except for H6, H7 and H13 of Pt-Cl and H7 and H13 of Pt-CN. Selected bond lengths [Å] and angles [°] of Pt-Cl: Cl1-Pt1 2.4001(13), P1-Pt1 2.3084(11), P2-Pt1 2.3058(11), C1-Pt1 2.010(5), P1-Pt1-P2 175.07(4), C1-Pt1-Cl1 178.98(13), P1-Pt1-Cl1 86.67(4), C1-Pt1-P1 92.63(10), C1-Pt1-P2 87.42(10), P2-Pt1-Cl1 93.21(4); Pt-CN: C14-Pt1 2.048(6), P1-Pt1 2.3098(14), P2-Pt1 2.3049(13), C1-Pt1 2.052(5), P1-Pt1-P2 175.39(5), C1-Pt1-C14 178.6(2), P1-Pt1-C14 88.16(15), C1-Pt1-P1 90.39(14), C1-Pt1-P2 87.92(14), P2-Pt1-C14 93.52(15).

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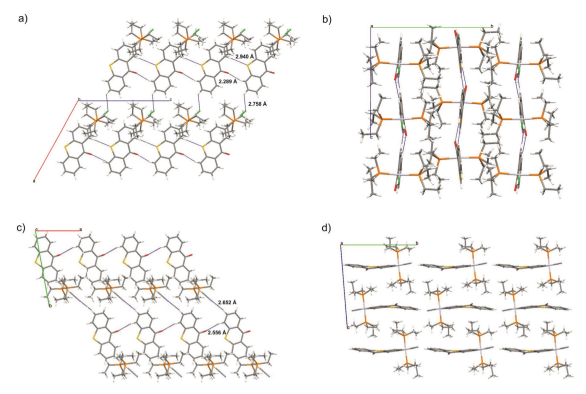


Fig. 3 Molecule packing of Pt-Cl and Pt-CN in the crystal. View along the crystallographic b axis (a) and the a axis (b) of Pt-Cl and along the crystallographic b tallographic c axis (c) and the a axis (d) of Pt-CN.

359.9 and 360.0, respectively (Fig. 2 and 3). While the X-Pt1-C1 (X = Cl1 of Pt-Cl or C14 of Pt-CN) bond angles of 178.98(13) and 178.6(2)° match the expected 180° of a squareplanar coordination sphere almost perfectly, the P1-Pt1-P2 bond angles of 175.07(4)° for Pt-Cl and 175.39(5)° for Pt-CN differ by about 5° from the ideal value. This is the result of a slight bending of the PEt₃ ligands toward the side of the intracyclic S atom of the thioxanthonyl ligand. The thioxanthonyl ligand is not exactly planar with some puckering about the S1-C11-O1 vector, which amounts to 5.0° for Pt-Cl and to 11.8° for Pt-CN. The best plane through the Tx ligand of the complexes adopts a nearly orthogonal orientation to the platinum coordination plane as is seen by the angles of 85.0° for Pt-Cl and 86.5° for Pt-CN formed between the metal bonded aryl ring of that ligand and the best plane through atoms Pt1, P1, P2, C1 and Cl1 or C14 of the cyanido ligand. The Pt-CN axis of complex Pt-CN is tilted by 1.2° with respect to the plane formed by atoms Pt1, C14, P1, P2 and C1. Pt-C1 bond lengths of 2.010(5) Å for Pt-Cl and of 2.052(5) Å for Pt-CN reflect the trans-effect of the (pseudo)halogenido ligand.

In the crystal, individual molecules of Pt-Cl associate to 1D chains along the crystallographic c axis by a total of four hydrogen bonds O1-H6a and S1-H13a with O···H and S···H distances of 2.289 and 2.940 Å for Pt-Cl, respectively, which is 43 and 19 pm shorter than the sum of the van der Waals radii (see Fig. 3a). In contrast, individual molecules of Pt-CN form two hydrogen bonds O1-H13a along the crystallographic a axis with a distance of 2.556 Å, 16 pm shorter than the sum of the

van der Waals radii, Fig. 3c. Within each chain of Pt-Cl, neighbouring molecules are tilted with respect to each other by 13.3° with respect to their thioxanthone planes in a slight zigzag pattern (Fig. 3b). Individual chains of both complexes are further connected through hydrogen bonds between H7a and the chlorido or cyanido ligand with H...X distances of 2.758 or 2.652 Å, respectively. For Pt-CN, these chains are offset along the bc plane and arrange in a brick wall-type fashion (see Fig. 3d).

Individual molecules of these complexes thus form twodimensional sheet structures within the ac or ab planes, respectively. Individual chains of Pt-Cl are further offset along the ac plane whereas those of Pt-CN are offset along the bc plane. Each thioxanthonyl ligand of the complexes is surrounded by two phosphine ligands belonging to neighbouring chains, one above and one below the thioxanthone plane. Thus no Pt···Pt or π -stacking interactions are observed in the solid state. This may contribute to the much higher solubility of the complexes when compared to the free ligand.

NMR spectroscopic characterisation

The ³¹P{¹H} NMR spectra of **Pt-Cl**, **Pt-Br**, **Pt-CN** and **Pt-I** all reveal just one sharp singlet flanked by 195Pt satellites (natural abundance 33.8%; Fig. 4a) which testifies to the trans arrangement of the phosphine ligands. The chemical shifts (Table 1) increase in the order I < CN < Br < Cl. For the halogenido ligands the ordering parallels decreasing σ-basicity against platinum, while the cyanido ligand one sticks out because of

a) b) c) Pt-CN Pt-CI Pt-CI Pt-Br Pt-Br Pt-I Pt-CI 18 10 6 151 149 147 145 143 141 139 137 8.55 8.45 7.75 7.65

Fig. 4 (a) ³¹P{¹H}NMR spectra of Pt-Cl, Pt-Br, Pt-CN and Pt-I. (b) ¹³C{¹H} NMR spectra of Pt-Cl, Pt-Br, Pt-CN and Pt-I. The asterisks indicate the signal of the platinum bonded carbon atom C1. (c) ¹H NMR spectra Pt-Cl, Pt-Br, Pt-CN and Pt-I in the spectral region of the ortho protons of the Tx ligand. The asterisks mark the resonance signal atom H13 and the circles that of H2.

 δ^{13} C (ppm)

Table 1 δ^{195} Pt, δ^{31} P, δ^{13} C and δ^{1} H chemical shifts of Pt–Br, Pt–Cl, Pt–I and Pt–CN with selected coupling constants; n.d. = not determined

	δ 195 Pt	$\delta^{~31}$ P [ppm]/ $^1J_{\mathrm{Pt-P}}$ [Hz]	δ 13 C, C1 [ppm]/ $^{1}J_{\mathrm{Pt-C}}$ [Hz]/ $^{2}J_{\mathrm{P-C}}$ [Hz]	δ ¹ H, H13 [ppm]/ $^3J_{\mathrm{Pt-H}}$ [Hz]	δ ¹ H, H2 [ppm]/ $^3J_{\mathrm{Pt-H}}$ [Hz]
Pt-Cl	-4257	13.93/2725	137.8/960/9.2	8.55/69.2	7.71/61.9
Pt-Br	-4382	11.91/2701	139.1/972/9.0	8.54/71.3	7.70/63.4
Pt-CN	-4674	10.73/2524	151.3/n.d./10.5	8.50/45.1	7.61/40.9
Pt-I	-4602	8.39/2668	143.7/n.d./8.8	8.52/71.9	7.66/64.0

its strong π -accepting capability. Observed coupling constants range from 2524 Hz for Pt-CN to 2725 Hz for Pt-Cl and are unexceptional when compared to other complexes trans- $Pt(\sigma-aryl)(PEt_3)_2(X)$. 79,80,100 The $^1J_{Pt-P}$ coupling constants reflect the s character of the Pt-P and Pt-X σ -bond 101,102 and follow the order CN < I < Br < Cl. The ¹⁹⁵Pt signal of every complex is split into a triplet by coupling to the two equivalent phosphorus nuclei and the shift follows a similar trend to the ³¹P chemical shift with the exception of complex **Pt-CN**, whose ¹⁹⁵Pt resonance appears at higher field than that of **Pt-I**.

 $\delta^{31}P$ (ppm)

The ¹³C NMR signal of the metal-bonded C1 atom of every complex is also split into a triplet due to coupling to the two cis positioned phosphorus nuclei with ²J_{P-C} coupling constants ranging from 8.8 Hz for Pt-I to 10.5 Hz for Pt-CN. The chemical shift of the C1 atom clearly depends on the (pseudo)halogenido ligand (Fig. 4b, Table 1 and Experimental section). From the data in Table 1 and Fig. 4b one can infer that the resonance signal of carbon atom C1 dramatically shifts to lower field as the back bonding ability of the trans disposed ligand increases. 195Pt satellites of the C1 13C resonance signal of 960 and 972 Hz, respectively, were detected for complexes Pt-Cl and Pt-Br but not for the other two complexes because of the lower signal to noise ratio of the experimental spectra. For the two complexes Pt-Cl and Pt-Br the ${}^{1}J_{\text{Pt-C1}}$ coupling constants have an inverse ordering to the ${}^{1}J_{Pt-P}$ coupling constants, which indicates that these two kinds of ligands compete for the 6s electron density at the Pt atom. The chemical shifts of

all other carbon atoms remain almost unchanged along this series. The coupling constants ${}^{3}J_{\text{Pt-H2}}$ and ${}^{3}J_{\text{Pt-H13}}$ to the Tx protons in the ortho positions to the platinum atom amount to ca. 70 Hz and 63 Hz, respectively, for the halogenido complexes and to ca. 45 or 41 Hz for the cyanido complex and follow the ordering of the trans-influence of the ligand $X (CN \ll Cl < Br < I)$.

δ¹H (ppm)

Electronic absorption spectroscopy

Fig. 5a depicts the room temperature electronic absorption spectra of the four complexes and of the free ligand Br-Tx in CH2Cl2 while relevant data are collected in Table 2. Like for the Br-Tx precursor, the spectra of the platinum complexes feature prominent UV absorption with extinction coefficients ε of the order of 4×10^4 and a much weaker one ($\varepsilon \approx 5 \times 10^3$) at about 400 nm which is responsible for the yellow colour of Br-Tx and the complexes. Attachment of the {(X)Pt(PEt₃)₂}-fragment (X = Br, Cl, I or CN) to the thioxanthone core increases the intensity of the UV bands and red shifts the low energy band by 644 cm^{-1} for the cyanido complex and by ca. 895 cm⁻¹ for the halogenido complexes. Donor substitution of thioxanthones at the 2-position generally red shifts the low energy band owing to a destabilisation of the highest occupied molecular orbital (HOMO). Shifts are in the range of 1740 cm⁻¹ (in acetonitrile) for alkoxy⁸⁸ and of 410 cm⁻¹ (in benzene) for alkyl87 substituents. Thus, in a first approximation, the $\{(X)Pt(PEt_3)_2\}$ -fragment seems to act as a moderate

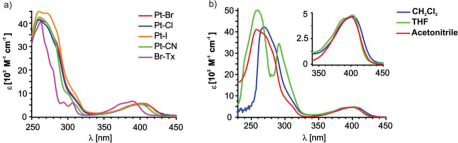


Fig. 5 (a) UV-Vis spectra in CH₂Cl₂ of Pt-Br, Pt-Cl, Pt-I, Pt-CN and the ligand bromothioxanthone (Br-Tx). (b) UV-Vis spectra of Pt-Br in acetonitrile (red), CH₂Cl₂ (blue) and THF (green).

Table 2 Absorption data of the ligand Br-Tx and the four complexes Pt-Cl, Pt-Br, Pt-CN and Pt-I (ca. 10^{-5} M in CH₂Cl₂, T = 298 K) and TD-DFT results of Pt-Br and Pt-CN in CH₂Cl₂; n.c. = not calculated

Compound	Absorption data	TD-DFT data			
	$\lambda_{\text{max}} [\text{nm}] (\varepsilon [10^3 \text{ M}^{-1} \text{ cm}^{-1}])$	λ [nm]	Main configuration	f^a	Assignment
Pt-Cl	260 (41.3), 272 (38.2), 286 (28.3),	240	HOMO → L+5 (83%)	0.19	MMLLCT
	310 (7.7), 403 (5.2)	248	$H-7 \rightarrow LUMO (61\%)$ $H-8 \rightarrow LUMO (16\%)$	0.29	ILCT/MMLL'CT
		261	$H-5 \rightarrow LUMO (54\%)$ $HOMO \rightarrow L+1 (19\%)$	0.63	$MMLL'CT/n {\rightarrow} \pi^*$
		274	$H-5 \rightarrow LUMO (11\%)$ $H-4 \rightarrow LUMO (14\%)$ $H-3 \rightarrow LUMO (19\%)$ $HOMO \rightarrow L+1 (36\%)$	0.23	$MMLL'CT/MLCT/\pi{\rightarrow}\pi^*$
		366	$HOMO \rightarrow LUMO (97\%)$	0.073	$\pi \rightarrow \pi^*/MMLL'CT$
Pt-Br	260 (41.5), 272 (39.5), 285 (31.2),	240	$HOMO \rightarrow L+5 (66\%)$	0.12	ILCT/MMLL'CT
	310 (7.7), 403 (5.7)	248	$H-8 \rightarrow LUMO (19\%)$ $H-7 \rightarrow LUMO (57\%)$	0.32	ILCT/MMLL'CT
		261	HOMO → L+1 (21%) H-6 → LUMO (57%)	0.73	$MLCT/n{\rightarrow}\pi^*$
		275	$H-1 \rightarrow L+2 (68\%)$	0.21	$\sigma \rightarrow \sigma^*$
		365	$HOMO \rightarrow LUMO (97\%)$	0.072	$\pi \rightarrow \pi^*/MMLL'CT$
Pt-CN	260 (42.5), 272 (39.3), 282 (32.5), 301 (9.6), 313 (4.3), 399 (5.3)	237	HOMO → L+4 (64%) H-1 → L+2 (22%)	0.21	$\pi \rightarrow \pi^*/\text{MLCT}$
		246	$H-7 \rightarrow LUMO (80\%)$	0.24	ILCT
		261	$H-5 \rightarrow LUMO (48\%)$ $HOMO \rightarrow L+1 (27\%)$ $H-2 \rightarrow LUMO (15\%)$	0.88	$MMLL'CT/ILCT/\pi{\rightarrow}\pi^*$
		362	$HOMO \rightarrow LUMO (97\%)$	0.075	$\pi \rightarrow \pi^*/MLCT$
Pt-I	260 (45.0), 273 (44.0), 285 (36.0), 310 (10.0), 403 (5.5)	248	$H-9 \rightarrow LUMO (15\%)$ $H-8 \rightarrow LUMO (58\%)$ $HOMO \rightarrow L+4 (12\%)$	0.32	MMLL'CT/ $\pi \rightarrow \pi^*$ /ILCT
		261	$H-6 \rightarrow LUMO(57\%)$ $HOMO \rightarrow L+1(12\%)$	0.86	$MLCT/ILCT/n{\rightarrow}\pi^*/MMLL'CT$
		279	$H-3 \rightarrow L+2 (74\%)$	0.13	$\sigma \rightarrow \sigma^*$
		362	$HOMO \rightarrow LUMO (97\%)$	0.072	$\pi \rightarrow \pi^*/MMLL'CT$
Tx-Br	262 (40.3), 270 (29.0), 279 (20.0), 293 (6.1), 306 (5.8), 389 (6.6)	n.c.	,		
^a Oscillator stre	ength.				

[&]quot;Oscillator strength.

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donor towards the thioxanthonyl core, thus lowering the HOMO–LUMO gap. 100 A more precise explanation of the redshift is that the highest occupied platinum d orbital $5d_{xz}$ is more strongly coupled to the π than to the π^* orbital of the thioxanthonyl ligand since the π and the $5d_{xz}$ orbitals are closer in energy. This leads to a destabilisation of the π orbital while the π^* orbital is stabilised by the interaction with the unoccupied Pt $6p_z$ orbital. Thus the energy gap of the $\pi \rightarrow \pi^*$ -transition is lowered. 78

As follows from our quantum chemical calculations (*vide infra*) the (relatively) weak low-energy band corresponds to the HOMO \rightarrow LUMO transition while several individual transitions contribute to the stronger, structured UV absorption. The energy of the low-energy band remains nearly unchanged upon substitution of the Br⁻ ligand by I⁻ or Cl⁻ while it is shifted by 370 cm⁻¹ to higher energy for the cyanido complex. This blue-shift is a result of the π -accepting ability of the cyanido ligand in *trans*-position to the Tx ligand, which

stabilises the HOMO. This observation is also in accord with the 13 C NMR chemical shifts where the strong back bonding ability of the cyanido ligand induces a deshielding of the *ipso* carbon. Nevertheless, the effect of the ligand X on the HOMO–LUMO gap is rather small. The electrochemical data of the four complexes derived from cyclic voltammetric experiments (0.1 M Bu₄NPF₆/CH₂Cl₂ ν = 400 mV s⁻¹) also show a small effect of the X-ligand since the potentials of the first irreversible oxidation are all in a similar range of 0.94 to 1.09 V (see Fig. S18–S21 of the ESI;† note, however, that the peak potentials of chemically irreversible processes do not strictly correspond to the thermodynamic E^0 values but are also affected by the equilibrium and forward rate constants of the chemical follow process).

Fig. 5b shows the electronic absorption spectra of Pt-Br at room temperature in different solvents. The position of the twin band at 400 nm reveals no solvent dependence whereas the onset of the UV band shifts to higher energy with increasing solvent polarity. Of note is also a distinct splitting of this latter band in THF whereas the lower energy components are only a shoulder superimposed on the more intense, higher energy one in the other solvents. From our observation that the split Vis band with individual peaks at ca. 400 nm and 390 nm exhibits no positional solvent dependence and closely resembles the Vis absorption of the free Tx ligand itself, we can infer that both relevant MOs are essentially confined to the σ -bonded Tx ligand and have obviously very similar spatial characteristics. In particular, the underlying transition(s) have no appreciable charge transfer character. Experimental and quantum chemical studies on thioxanthones point to mainly $\pi \rightarrow \pi^*$ -character of the underlying transition(s). That assumption is further supported by our present TD-DFT calculations on Pt-Br, Pt-Cl, Pt-I and Pt-CN (Table 2, Fig. 6, and Fig. 25, 27 and 28 of the ESI†). These calculations indicate that the HOMO, the LUMO and the LUMO+1 are almost exclusively located on the thioxanthonyl moiety with major orbital coefficients on the intracyclic sulfur atom for the HOMO, at the thioxanthone carbonyl function for the LUMO and a nodal plane along the S···CO axis for the LUMO+1.

In contrast, the UV bands in the range of 310 to 260 nm possess negative solvatochromism which indicates charge

Table 3 Luminescence data of Pt-Br, Pt-Cl, Pt-I and Pt-CN in nitrogen-saturated and of Br-Tx in aerated CH_2Cl_2 solutions at a concentration of 10^{-5} mol L^{-1} ; n.m. = not measured

	$\lambda_{\mathrm{max,Fl}}$ [nm] (St. shift [cm ⁻¹])	$\lambda_{\max, Ph}$ [nm] (St. shift [cm ⁻¹])	$\phi_{ m Fl}$	ϕ Ph	τ [μs]
Br-Tx	417 (1726)	_	n.m.	_	_
Pt-Cl	451 (2640)	520 (5583)	0.006	0.188	2.17
Pt-Br	$450/421^a$ (2592)	$517/491^a$ (5472)	0.006	0.120/	n.m.
				0.031^{a}	
Pt-I	441 (2138)	511 (5244)	0.004	0.017	n.m.
Pt-CN	439 (2283)	507 (5338)	0.015	0.069	n.m.

 $^{^{}a}$ Measured in benzene at ambient temperature in nitrogen saturated solutions.

transfer contributions to at least one of the major transitions within the band envelope. According to our TD-DFT calculations, the overall UV absorption comprises four major components: an absorption at a (calculated) energy of 275 nm, which is essentially the HOMO-1 \rightarrow LUMO+2 transition, one very strong absorption at 261 nm mainly corresponding to the HOMO-6 → LUMO (57%) and HOMO → LUMO+1 (21%) transitions, a band at 248 nm, which is dominated by the HOMO $-7 \rightarrow \text{LUMO}$ (57%) and the HOMO-8 \rightarrow LUMO (19%) transitions, and one rather weak band at 240 nm of mainly HOMO → LUMO+5 origin. As follows from Fig. 6, the band calculated at 275 nm is a $\sigma \rightarrow \sigma^*$ transition of the Br-Pt bond accompanied by an electron density increase at the Br ligand and the occurrence of a nodal plane between the Pt and the Br atom. The band calculated at 261 nm is a mixture of an intraligand charge-transfer (ILCT) and $n\rightarrow\pi^*$ transition, both centred at the thioxanthonyl ligand with some shift of electron density from the S and O atoms to the carbon atoms of the thioxanthonyl moiety and a shift of electron density from the Pt bound aryl to the outermost phenyl ring. The 248 nm band comprises a thioxanthone-based intraligand charge transfer (ILCT) and a mixed metal/ligand-to-ligand' charge-transfer (MLL'CT) from the {(PEt₃)₂Pt}-unit to the thioxanthonyl ligand owing to appreciable Pt-aryl δ character of the HOMO-8. Of note is also an energy splitting of 0.28 to 0.29 eV between the similar MOs HOMO-8 and HOMO-6, which represent the δ-bonding and -antibonding interactions between the Pt atom and the Ptbonded aryl ring of the Tx ligand. The band at 240 nm has also some ILCT- and MLL'CT component. Similar results have been obtained for complexes Pt-CN, Pt-Cl and Pt-I (see Fig. S25, S27 and S28 of the ESI†). However, the $\sigma \rightarrow \sigma^*$ transition of Pt-Br and Pt-I is absent in Pt-CN and Pt-Cl as a result of the energetically low-lying MOs for the Pt-Cl and Pt-CN σ-bonds.

Luminescence spectroscopy

Fig. 7a displays the room temperature emission spectra of Br-Tx in 10⁻⁵ M aerated and of Pt-CN, Pt-I, Pt-Br and Pt-Cl in 10⁻⁵ M nitrogen-saturated CH₂Cl₂ solutions. Br-Tx emits at 417 nm upon irradiation at 370 nm. Excitation at 400 nm into the HOMO → LUMO band of the four complexes results in dual emission at around 450 nm and at ca. 510 nm. Stokes shifts are in the range of 2140 to 2640 cm⁻¹ for the higher energy emission and of 5244 to 5583 cm1 for the one at the lower energy. We note here that the Stokes shifts of the higher energy emission band of the platinum complexes is only mildly larger as that of 1760 cm⁻¹ for the fluorescence emission of Br-Tx itself and it is clearly different from that of Br-Tx in each case. For every complex, the emission band at 450 nm is insensitive against oxygen while that at 510 nm is almost completely quenched in aerated solutions (Fig. 7c). This observation in concert with the large Stokes shifts let us assign the high energy emission as fluorescence and the one at lower energy as phosphorescence, both arising from the σ -bonded Tx ligand. Excitation spectra of Pt-Cl (see Fig. S32 of the ESI†) support this assignment. Emission detection at wavelengths of

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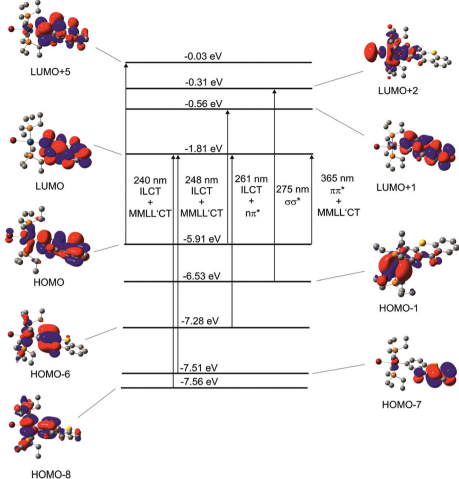


Fig. 6 Graphical representations of selected MOs and their TD-DFT energies for complex Pt-Br. Arrows symbolise the main contributors to the respective transitions.

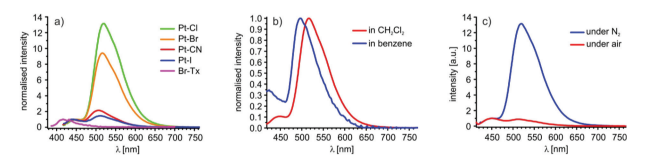


Fig. 7 (a) Emission spectra of Pt–Cl (green), Pt–Br (orange), Pt–CN (red), and Pt–I (blue) in nitrogen-saturated and Br–Tx in aerated CH₂Cl₂solutions. (b) Emission spectra of Pt–Br in CH₂Cl₂ (red), and benzene (blue). (c) Emission spectra of Pt–Cl under nitrogen atmosphere (blue) and under air (red).

450 and 501 nm provide essentially identical spectra with a prominent band 400 nm in exact match with the low-energy $\pi \rightarrow \pi^*$ absorption of the Pt-bonded Tx substituent. In further agreement with that hypothesis dual emission from the complexes with attenuated intensities was also observed when **Pt-Br** was irradiated at 280, 350 or 400 nm (see Fig. S33 of the ESI†). This clearly indicates that both emissions emanate from the same compound.

The influence of the ligand X on the emission energies is small, but both bands shift to slightly higher energy as Pt–X back bonding increases (Cl < Br \ll I < CN). The wavelength of the fluorescence emission of the (pseudo)halogenido complexes thus follows an inverse order as that of the electronic absorption. One possible explanation is that the first excited singlet state undergoes some structural reorientation before emission occurs which leads to a stronger contribution

of the (pseudo)halogenido ligand. Such structural reorienta-

tion upon irradiation of thioxanthones is not without precedence in the literature96 and also underlies the solvent dependency of Tx emission, very similar to the one that we observe for the investigated complexes (Fig. 7b). Computed structure changes between the T₁ and S₀ states are, however, rather minor and basically confined to the platinum bonded Tx ligand with a contraction of the Pt-C(Tx) and the C4-S bond lengths of ca. 3 pm and elongation of the Tx C=O and the C1-C2 bonds of about 4 pm and 2 pm, respectively (Tables S5 to S8 of the ESI†). The same holds for calculated spin densities as is exemplarily shown for Pt-Cl in Fig. S26 of the ESI.†

For the three halogenido complexes the phosphorescence quantum yields increase with increasing σ-basicity of the ligand X against platinum in the order I < CN < Br < Cl from smaller than 2% for Pt-I to close to 19% for Pt-Cl (Table 3). With increasing σ-basicity of the ligand X, the ligand-field splitting also increases which in turn destabilises the nonemissive Pt 5d states. Therefore, thermal excitation from the emissive thioxanthone-centred ${}^{3}\pi\pi^{*}$ -states to the Pt 5d states undergoing radiationless decay is diminished. 103 Ligand-field arguments would, however, predict that the cyanido complex is the strongest triplet emitter along this series of complexes, which is clearly not the case. Radiationless decay of the excited triplet state may also be triggered by excitation of vibrational quanta, and this decay channel may be particularly efficient in **Pt-CN** due to the high energy of the C≡N stretch. ¹⁰⁴

The phosphorescence quantum yield further increases with increasing polarity of the solvent as is exemplified by complex Pt-Br, which shows a quantum yield of 12% in CH₂Cl₂ but only one of 3% in benzene. Such behaviour has also been reported for purely organic thioxanthones, which also show strong solvent dependence of the rates of internal conversion and intersystem crossing owing to a modulation of the relative energies of the S₁, S₂, T₁ and T₂ states. 94-96 This effect is also reflected in the energies of the fluorescence and the phosphorescence bands of Pt-Br that are blue shifted by 1530 cm⁻¹ and 1025 cm⁻¹, respectively, when the solvent is changed from CH₂Cl₂ to benzene.

We note that the fluorescence quantum yields and, obviously, the lifetimes of the four complexes vary only very little and are also very similar to those of purely organic thioxanthones. 88,94,96,105 The rate constant of fluorescence can be estimated as ca. 1.2 to 0.3 ns by the established equation: 105

$$\Phi_{\rm f} = k_{\rm f} \tau_{\rm f} \text{ with } k_{\rm f} = 5.12 \times 10^7 \text{ s}^{-1}.$$

This shows that attachment of the {Pt(PEt₃)X}-fragment has an only limited influence on the already high rate of intersystem crossing (ISC) of the Tx dye. The quantum yield of triplet formation of parent thioxanthone has been determined as 66% for acetonitrile and 84% for benzene solutions. 97 Of note is, however, the observation of moderately to rather intense room-temperature phosphorescence from the thioxanthonyl ligand with quantum yields of up to almost 19%. Organic thioxanthones, while being phosphorescent in glassy matrices

at 77 K,89,90,94 usually show only very weak emission under these conditions, if any at all. 97 Intersystem crossing from the excited S₁ to the excited T₂ state of thioxanthones is usually very fast (see Fig. 1) and occurs within 10 ps^{86,95} while the Internal Conversion from the T2 to the T1 state occurs at a rate which is identical to the lifetime of the fluorescence (1.5-2 ns).86 In contrast, the T₁ state is very long-lived with half-lives in the range of 100 to 140 ms, making it very susceptible to quenching and photoreactions such as H atom abstraction. The role of the covalently bonded Pt centre is thus to open a pathway for relatively rapid decay of the T1 state to the ground state with the emission of light due to its high spin-orbit coupling constant, which is not available for thioxanthone itself or its organic derivatives. Pt-Cl, the strongest triplet emitter of the present series, has a phosphorescence half-life of 2.17 µs, which is by five orders of magnitude lower compared to that of purely organic thioxanthones. The lifetime clearly proves that the emission band at 510 nm originates from a triplet state and further indicates that phosphorescence emission now becomes competitive to radiationless deactivation pathways.

Conclusions

Four new square-planar complexes trans-Pt(PEt₃)₂(X)(thioxanthon-2-yl) (X = Cl, Br, CN or I) were successfully synthesised by oxidative addition of 2-bromo-9H-thioxanthen-9-one to Pt-(PEt₃)₄ and subsequent halide exchange. Electronic absorption data in concert with TD-DFT calculations reveal a considerable red-shift of the thioxanthone-centred $\pi \rightarrow \pi^*$ (HOMO \rightarrow LUMO) transition when compared to the free ligand in accordance with the overall donor capabilities of the trans-{Pt(PEt₃)₂X}fragment. All four complexes exhibit dual fluorescence and phosphorescence emission in fluid solution at room temperature arising from excited singlet and triplet states located at the thioxanthonyl ligand. While the fluorescence quantum yield is largely independent on X and very similar to that of other thioxanthones, that of the phosphorescence emission is strongly modulated by the ligand X, ranging from 1.7% for Pt-I to a truly exceptional value of 18.8% for Pt-Cl in nitrogensaturated CH₂Cl₂ solution. In an intuitive ordering, the phosphorescence quantum yield of the halogenido complexes increases with the ligand-field splitting abilities of ligand X. The lower phosphorescence quantum yield of the strongfield cyanido ligand may be due to its high-energy C≡N stretch, which probably provides an efficient non-radiative deactivation channel. Key to the observation of dual emission is that the platinum coordination centre makes only limited contribution to the relevant frontier orbitals of the excited states^{70–72} as is, e.g., seen by the computed spin density distributions of the T_1 state. This attenuated Pt contribution has no noticeable influence on the already fast rate of ISC inherent to thioxanthones. It is, however, still sufficient to open a radiative deactivation channel for the usually very long-lived triplet state that is unattainable for thioxanthones themselves, thus diminishing the lifetime of the excited triplet state of the Tx ligand

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by five orders of magnitude to a value of ca. 2 μ s. This allows for thioxanthone-based phosphorescence to be observed in fluid solution with unprecedentedly high quantum yield. We note here that complexes exhibiting dual fluorescence and phosphorescence emission have advantages over two-component sensors for quantitative detection of singlet-oxygen. The platinum Tx complexes herein display strong phosphorescence and quantitative quenching of that emission at atmospheric O_2 levels while fluorescence emission is fully retained. This makes them promising candidates for such application, and work along these lines is underway.

Experimental section

Materials and general methods

All reactions, except the oxidative addition of 2-bromo-9H-thioxanthen-9-one (Br-Tx) to $Pt(PEt_3)_4$, were performed under air with undried solvents. THF was distilled from blue sodium/benzophenone ketyl and was degassed by three freeze-pump-thaw cycles. All other solvents were used without further purification. 2-Bromo-9H-thioxanthen-9-one (Br-Tx)¹⁰⁶ and [Pt- $(PEt_3)_4$]¹⁰⁷ were synthesised according to literature procedures.

NMR experiments were carried out on a Varian Unity Inova 400, a Bruker Avance III DRX 400 or a Bruker Avance DRX 600 spectrometer. 1 H and 13 C NMR spectra were referenced to the solvent signal. 31 P and 195 Pt NMR spectra were referenced to external standard (85% $_{3}$ PO₄ or saturated solution of $_{3}$ PtCl₆ in D₂O, respectively). NMR spectral data are given as follows: chemical shift ($_{3}$ ppm), multiplicity (m: multiplet, s: singlet, d: doublet, t: triplet, quin: quintet, vm: virtual multiplet,), coupling constant (Hz), integration. In addition to 1D NMR experiments, complete NMR characterisations were performed by 2D NMR experiments ($_{4}$ H, $_{4}$ H gCOSY, $_{4}$ H, $_{4}$ C gHSQC, $_{4}$ H, $_{4}$ C gHMBC and $_{4}$ H, $_{4}$ P gHMBC).

X-ray diffraction analysis was performed at 100 K on a STOE IPDS-II diffractometer equipped with a graphite-monochromated radiation source ($\lambda = 0.71073 \text{ Å}$) and an image plate detection system. A crystal mounted on a fine glass fiber with silicon grease was employed. If not indicated otherwise, the selection, integration and averaging procedure of the measured reflex intensities, the determination of the unit cell dimensions and a least-squares fit of the 2θ values as well as data reduction, LP-correction and space group determination were performed using the X-Area software package delivered with the diffractometer. A semiempirical absorption correction was performed. The structure of Pt-Cl was solved by the heavyatom methods (SHELXS-97), whereas that of Pt-CN was solved by direct methods (SHELXS-97). 118 Solutions of both structures were completed with difference fourier syntheses and refined with full-matrix least-square using SHELXL-97¹⁰⁸ minimising $\omega(F_0^2 - F_c^2)^2$. Weighted R factor (wR²) and the goodness of fit GooF are based on F^2 . All hydrogen atoms were treated according to a riding model. Molecular structures in this work are plotted with ORTEP 3.109 Hydrogen atoms and solvent molecules are omitted for clarity.

UV-Vis spectra were obtained on a TIDAS fiberoptic diode array spectrometer MCS UV/NIR from j&m in HELLMA quartz cuvettes with 1 cm optical path lengths. Emission spectra were measured on a Perkin Elmer luminescence spectrometer LS50 using HELLMA quartz cuvettes modified with an angle valve from Normag. Time-resolved luminescence spectroscopy was measured on a LP920-KS instrument from Edinburgh Instruments, equipped with a R928 photomultiplier and an iCCD camera from Andor using a Starna quartz cuvette modified with an angle valve from Rotaflow. Luminescence was excited with the frequency-doubled output from a Quantel Brilliant b laser and the pulse length of approximately 10 ns. The concentration of samples was ca. 1.5×10^{-5} M.

Electrochemical investigations in degassed dichloromethane were performed in a vacuum-tight one-compartment cell on a BAS CV50 potentiostat using Pt or glassy carbon disk electrodes from BAS as the working electrode, a platinum counter electrode and a silver pseudoreference electrode. Potentials were calibrated against the ferrocene/ferrocenium couple (E^0 Fc/Fc⁺ = 0.000 V).

Computational details

The ground state electronic structures were calculated by density functional theory (DFT) methods using the Gaussian 09¹¹⁰ program packages. Quantum chemical studies were performed without any symmetry constraints. Open shell systems were calculated by the unrestricted Kohn-Sham approach (UKS). 111 Geometry optimization followed by vibrational analysis was made either in vacuum or in solvent media. The quasirelativistic Wood-Boring small-core pseudopotentials (MWB)112,113 and the corresponding optimized set of basis functions¹¹⁴ for Pt and 6-31G(d) polarized double-ζ basis set¹¹⁵ for the remaining atoms were employed together with the Perdew, Burke, Ernzerhof exchange and correlation functional (PBE0). 116,117 Solvent effects were described by the polarizable conductor continuum model (PCM)118-121 with standard parameters for CH2Cl2. Absorption spectra and orbital energies were calculated using time-dependent DFT (TD-DFT)¹²² using the same functional/basis set combination mentioned above. For easier comparison with the experiment, the obtained absorption and emission energies were converted into wavelengths and broadened by a Gaussian distribution (full width at half maximum = 3000 cm⁻¹) using the program Gauss-Sum. 123 Molecular orbitals were visualised with the GaussView program. 124

trans-Bromido(thioxanthon-2-yl)bis(triethylphosphine)platinum(n) (Pt-Br). In a 50 mL Schlenk tube Pt(PEt₃)₄ (1.26 g, 1.89 mmol) and 2-bromo-9H-thioxanthen-9-one (549 mg, 1.89 mmol) were suspended under nitrogen atmosphere in 10 mL of degassed THF. After stirring the yellow suspension for 1.5 h at room temperature complete consumption of Pt(PEt₃)₄ was observed by ³¹P NMR experiments. The solvent was removed and the remaining yellow solid was washed with pentane (3 × 5 mL). The product was extracted from the residue with toluene (3 × 5 mL) and recrystallised from 7 mL of toluene. The needle shaped crystalline material was redis-

solved in benzene and then frozen at 273 K. The benzene was sublimed off in vacuo to yield 810 mg of the yellow product. Yield: 59%. ¹H NMR (600.2 MHz, CD_2Cl_2 , 298 K) δ 8.55 (d, $^3J_{HH}$ = 8.2 Hz, 1H, H9), 8.54 (s, with satellites ${}^{3}J_{PtH}$ = 71.3 Hz, 1H, H13), 7.70 (d, ${}^{3}J_{HH}$ = 8.2 Hz, with satellites ${}^{3}J_{PtH}$ = 63.4 Hz, 1H, H2), 7.61-7.57 (m, 2H, H6, and H7), 7.44 (m, 1H, H8), 7.21 (d, $^{3}J_{HH}$ = 8.2 Hz, 1H, H3), 1.67 (m, 12H, H14), 1.07 (m, 18H, H15). 13 C NMR (150.9 MHz, CD₂Cl₂, 298 K) δ 179.6 (s, C11), 142.7 (t, ${}^{3}J_{PC}$ = 2.2 Hz, with satellites ${}^{2}J_{PtC}$ = 43.8 Hz, C2), 139.7 (t, ${}^{2}J_{PC}$ = 9.0 Hz, with satellites ${}^{1}J_{PtC}$ = 972 Hz, C1), 138.3 (s, C5), 137.5 (t, ${}^{3}J_{PC} = 2.7$ Hz, with satellites ${}^{2}J_{PtC} = 50.4$ Hz, C13), 132.2 (s, C7), 130.7 (t, ${}^{5}J_{PC}$ = 1.4 Hz, C4), 130.3 (s, C10), 130.0 (s, C9), 129.1 (t, ${}^{4}J_{PC}$ = 1.2 Hz, C12), 126.6 (s, C6), 126.2 (s, C8), 125.2 (s, with satellites ${}^{3}J_{PtC} = 72.8$ Hz, C3), 14.8 (vquin, J =16.9 Hz, C14), and 8.1 (s, with satellites ${}^{3}J_{PtC}$ = 24.5 Hz, C15). 31 P NMR (162 MHz, CD₂Cl₂, 298 K) δ 11.91 (s, with satellites $^1\!J_{\rm PtP}$ = 2701 Hz). $^{195}{\rm Pt}$ NMR (85.6 MHz, ${\rm CD_2Cl_2},$ 298 K) δ -4382(t, ${}^{1}J_{PtP}$ = 2701 Hz). Anal. Calcd for C₂₅H₃₇BrOP₂PtS: C, 41.56; 5.16. Found: C, 42.11; H, 5.30%.

trans-Chlorido(thioxanthon-2-yl)bis(triethylphosphine)platinum(II) (Pt-Cl). Pt-Br (46.5 mg, 64.4 µmol) and AgBF₄ (12.6 mg, 64.4 µmol) were suspended in 3 mL of methanol and heated at 65 °C for 10 min. NaCl (8 mg, 137 µmol) was added and the yellow suspension was stirred for 16 h at room temperature. The precipitate was filtered off and solvent was removed from the filtrate. The remaining yellow solid was washed with water (3 × 5 mL) and recrystallised from toluene to give 27.5 mg of yellow, needle shaped crystals. (Yield: 63%). ¹H NMR (600.33 MHz, CD_2Cl_2 , 298 K) δ 8.55 (s, with satellites $^{3}J_{PtH}$ = 69.2 Hz, 1H, H13), 8.54 (m, 1H, H9), 7.71 (dd, $^{3}J_{HH}$ = 8.13 Hz, ${}^{4}J_{HH}$ = 1.24 Hz, with satellites ${}^{3}J_{PtH}$ = 61.9 Hz, 1H, H2), 7.59 (m, 2H, H6, and H7), 7.44 (m, 1H, H8), 7.19 (d, ${}^{3}J_{HH} =$ 8.13 Hz, 1H, H3), 1.62 (m, 12H, H14), 1.08 (m, 18H, H15). ¹³C NMR (151.0 MHz, CD_2Cl_2 , 298 K) δ 180.2 (s, C11), 143.0 (t, ${}^3J_{PC}$ = 2.0 Hz, with satellites ${}^{2}J_{PtC}$ = 43.0 Hz, C2), 138.3 (s, C5), 137.8 $(t, {}^{2}J_{PC} = 9.2 \text{ Hz}, \text{ with satellites } {}^{1}J_{PtC} = 960 \text{ Hz}, C1), 137.7 (t, {}^{3}J_{PC})$ = 2.56 Hz, with satellites ${}^{2}J_{PtC}$ = 48.2 Hz, C13), 132.19 (s, C7), 130.5 (t, ${}^{5}J_{PC}$ = 1.38 Hz, C4), 130.3 (s, C10), 130.0 (s, C9), 129.0 (s, with satellites ${}^{3}J_{PtC} = 70.5$ Hz, C12), 126.6 (s, C6), 126.2 (s, C8), 125.1 (s, with satellites ${}^{3}J_{PtC}$ = 71.1 Hz, C3), 14.2 (vquin, J = 16.9 Hz, C14), 8.1 (s, with satellites ${}^{2}J_{PtC}$ = 23.5 Hz, C15). ${}^{31}P$ NMR (161.84 MHz, CD_2Cl_2 , 298 K) δ 13.93 (s, with satellites $^{1}J_{\text{PtP}} = 2725 \text{ Hz}, 2P$). $^{195}\text{Pt} \text{ NMR} (85.6 \text{ MHz}, \text{CD}_{2}\text{Cl}_{2}, 298 \text{ K})$ δ -4257 (t, ${}^{1}J_{PtP}$ = 2725 Hz). Anal. Calcd for C₂₅H₃₇ClOP₂PtS: C, 44.28; 5.50. Found: C, 44.28; H, 5.37%. Crystals suitable for X-ray diffraction analysis were obtained by slow evaporation of the solvent from a saturated CH₂Cl₂ solution.

trans-Cyanido(thioxanthon-2-yl)bis(triethylphosphine)platinum(II) (Pt–CN). Pt–Br (41 mg, 56.7 μmol) and AgCN (7.6 mg, 56.7 μmol, prepared from AgNO₃ and KCN) were suspended in 0.5 mL of acetone and heated at reflux for 5 min. The obtained solid was filtered off through a syringe filter and the filter cake was washed with CH_2Cl_2 (2 × 2 mL). The solvent was removed from the combined filtrates and the pure yellow product was obtained by recrystallisation from acetone. (Yield: 89%). ¹H NMR (600.33 MHz, CD_2Cl_2 , 298 K) δ 8.55 (d, $^3J_{HH}$ = 8.0 Hz, 1H,

H9), 8.50 (s, with satellites ${}^{3}J_{PtH} = 45.1$ Hz, 1H, H13), 7.61 (d, $^{3}J_{HH}$ = 8.0 Hz, with satellites $^{3}J_{PtH}$ = 40.9 Hz, 1H, H2), 7.60 – 7.57 (m, 2H, H6 and H7), 7.44 (m, 1H, H8), 7.29 (d, ${}^{3}J_{HH} = 8.0 \text{ Hz}$, 1H, H3), 1.73 (m, 12H, H14), and 1.10 (dt, ${}^{3}J_{HH}$ = 7.6 Hz, ${}^{3}J_{PH}$ = 16.8 Hz, 18H, H15). 13 C NMR (151.0 MHz, CD₂Cl₂, 298 K) δ 180.5 (s, C11), 151.3 (t, ${}^{2}J_{PC}$ = 10.5 Hz, C1), 143.5 (t, ${}^{3}J_{PC}$ = 2.2 Hz, with satellites ${}^{3}J_{PtC}$ = 27.9 Hz, C2), 138.5 (t, ${}^{3}J_{PC}$ = 2.6 Hz, with satellites ${}^2J_{\rm PtC}$ = 31.1 Hz, C13), 138.3 (s, C5), 133.7 (t, ${}^2J_{\rm PC}$ = 12.4 Hz, C(CN)), 132.2 (s, C7), 131.3 (s, C4), 130.5 (s, C10), 130.0 (s, C9), 129.0 (s, C12), 126.6 (s, C6), 126.2 (s, C8), 125.2 (s, with satellites ${}^{3}J_{PtC}$ = 49.9 Hz, C3), 16.1 (vquin, J = 17.60 Hz, C14), and 8.3 (s, with satellites ${}^{3}J_{PtC}$ = 24.4 Hz, C15). ${}^{31}P$ NMR (161.84 MHz, CD_2Cl_2 , 298 K) δ 10.73 (s, with satellites ${}^{1}J_{PtP}$ = 2524 Hz, 2P). ¹⁹⁵Pt NMR (85.6 MHz, CD₂Cl₂, 298 K) δ –4674 (t, ${}^{1}J_{\text{PtP}}$ = 2524 Hz). Anal. Calcd for C₂₆H₃₇NOP₂PtS: C, 46.70; H, 5.58; N, 2.09; S, 4.80. Found: C, 46.52; H, 5.60, N, 2.19; S, 4.69%. Crystals suitable for X-ray diffraction analysis were obtained by slow evaporation of the solvent from a saturated CH₂Cl₂ solution.

trans-Iodido(thioxanthonyl)bis(triethylphosphine)platinum (II) (Pt-I). Pt-Br (20 mg, 27.7 μmol) and AgOTf (10.7 mg, 41.5 µmol) were combined in methanol and heated to reflux for 5 min. NaI (8.3 mg, 55.4 µmol) was added and the reaction mixture was stirred at ambient temperature for 1 h. The beigeyellowish precipitate was filtered off and the solvent was removed from the filtrate. The remaining yellow solid was redissolved in benzene and filtered through a syringe filter. The solvent was removed and the vellow product washed with pentane (3 × 1.5 mL) and dried in vacuo. (Yield: 89%). ¹H NMR (600.2 MHz, CD_2Cl_2 , 298 K) δ 8.55 (d, ${}^3J_{HH}$ = 8.0 Hz, 1H, H9), 8.52 (s, with satellites ${}^{3}J_{PtH}$ = 71.9 Hz, 1H, H13), 7.66 (dd, ${}^{3}J_{HH}$ = 8.1 Hz, ${}^{4}J_{HH}$ = 1.5 Hz, with satellites ${}^{3}J_{PtH}$ = 64.0 Hz, 1H, H2), 7.62-7.58 (m, 2H, H6, and H7), 7.45 (m, 1H, H8), 7.23 (d, broad, ${}^{3}J_{HH}$ = 8.1 Hz, 1H, H3), 1.77 (m, 12H, H14), and 1.05 (dt, ${}^{3}J_{HH} = 7.7$ Hz, ${}^{3}J_{PH} = 16.4$ Hz, 18H, H15). ${}^{13}C$ NMR (150.9 MHz, CD_2Cl_2 , 298 K) δ 180.3 (s, C11), 143.7 (t, ${}^2J_{PC}$ = 8.8 Hz, C1), 142.3 (s, with satellites ${}^{2}J_{PtC} = 44.4$ Hz, C2), 138.3 (s, C5), 137.2 (s, with satellites ${}^{2}J_{PtC}$ = 49.9 Hz, C13), 132.3 (s, C8), 130.9 (s, C4), 130.3 (s, C10), 130.0 (s, C9), 129.1 (s,C12), 126.6 (s, C6), 126.3 (s, C8), 125.3 (s, with satellites ${}^{3}J_{PtC} = 74.5$ Hz, C3), 16.0 (vquin, J = 17.4 Hz, C14), and 8.2 (s, with satellites ${}^{3}J_{PtC} =$ 25.0 Hz, C15). ³¹P NMR (161.84 MHz, CD₂Cl₂, 298 K) δ 8.39 (s, with satellites ${}^{1}J_{PtP} = 2668 \text{ Hz}, 2P$). ${}^{195}Pt \text{ NMR} (85.6 \text{ MHz},$ CD_2Cl_2 , 298 K) δ -4602 (t, ${}^{1}J_{PtP}$ = 2668 Hz). Anal. Calcd for C₂₅H₃₇IOP₂PtS: C, 39.02; 4.85. Found: C, 38.46; H, 4.40%.

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References

Dalton Transactions

- 1 M. A. Baldo, D. F. O'Brien, Y. You, A. Shoustikov, S. Sibley, M. E. Thompson and S. R. Forrest, *Nature*, 1998, **395**, 151–154.
- R. D. Costa, E. Ortí, H. J. Bolink, F. Monti, G. Accorsi and N. Armaroli, *Angew. Chem., Int. Ed.*, 2012, 51, 8178–8211.
- 3 R. Visbal and M. C. Gimeno, *Chem. Soc. Rev.*, 2014, 43, 3551-3574.
- 4 L. F. Gildea and J. A. G. Williams, in *Organic Light-Emitting Diodes (OLEDs)*, ed. A. Buckley, Woodhead Publishing, 2013, pp. 77–113, DOI: 10.1533/9780857098948.1.77.
- 5 D.-L. Ma, V. P.-Y. Ma, D. S.-H. Chan, K.-H. Leung, H.-Z. He and C.-H. Leung, *Coord. Chem. Rev.*, 2012, 256, 3087– 3113.
- 6 K. Li, G. Cheng, C. Ma, X. Guan, W.-M. Kwok, Y. Chen, W. Lu and C.-M. Che, *Chem. Sci.*, 2013, 4, 2630–2644.
- 7 D.-L. Ma, H.-Z. He, K.-H. Leung, D. S.-H. Chan and C.-H. Leung, *Angew. Chem., Int. Ed.*, 2013, **52**, 7666–7682.
- 8 K. K.-W. Lo and S. P.-Y. Li, in *Molecular Design and Applications of Photofunctional Polymers and Materials*, 2012, vol. 2, pp. 130–198.
- 9 Q. Zhao, C. Huang and F. Li, Chem. Soc. Rev., 2011, 40, 2508–2524.
- 10 P. K. M. Siu, S.-W. Lai, W. Lu, N. Zhu and C.-M. Che, *Eur. J. Inorg. Chem.*, 2003, 2749–2752.
- 11 V. Guerchais and J.-L. Fillaut, *Coord. Chem. Rev.*, 2011, 255, 2448–2457.
- 12 Q. Zhao, F. Li and C. Huang, Chem. Soc. Rev., 2010, 39, 3007–3030.
- 13 K. K.-W. Lo, A. W.-T. Choi and W. H.-T. Law, *Dalton Trans.*, 2012, **41**, 6021–6047.
- 14 Z. Liu, W. He and Z. Guo, *Chem. Soc. Rev.*, 2013, **42**, 1568–1600.
- 15 H.-B. Xu, L.-X. Shi, E. Ma, L.-Y. Zhang, Q.-H. Wei and Z.-C. Chen, *Chem. Commun.*, 2006, 1601–1603.
- 16 T. K. Ronson, T. Lazarides, H. Adams, S. J. A. Pope, D. Sykes, S. Faulkner, S. J. Coles, M. B. Hursthouse, W. Clegg, R. W. Harrington and M. D. Ward, *Chem. – Eur. J.*, 2006, 12, 9299–9313.
- 17 W. T. Eckenhoff and R. Eisenberg, *Dalton Trans.*, 2012, 41, 13004–13021.
- 18 J. A. G. Williams, Top. Curr. Chem., 2007, 281, 205-268.
- 19 L. J. Rothberg and A. J. Lovinger, *J. Mater. Res.*, 1996, **11**, 3174–3187.
- 20 H. Yersin, Top. Curr. Chem., 2004, 241, 1-26.
- 21 P.-T. Chou, Y. Chi, M.-W. Chung and C.-C. Lin, *Coord. Chem. Rev.*, 2011, 255, 2653–2665.

- 22 J. Schneider, P. Du, P. Jarosz, T. Lazarides, X. Wang, W. W. Brennessel and R. Eisenberg, *Inorg. Chem.*, 2009, 48, 4306–4316.
- 23 J. A. G. Williams, A. Beeby, E. S. Davies, J. A. Weinstein and C. Wilson, *Inorg. Chem.*, 2003, 42, 8609–8611.
- 24 W. J. Brooks, V. Babayan, S. Lemansky, P. I. Djurovich, I. Tsyba, R. Bau and M. E. Thompson, *Inorg. Chem.*, 2002, 41, 3055–3066.
- 25 S. J. Farley, D. L. Rochester, A. L. Thompson, J. A. K. Howard and J. A. G. Williams, *Inorg. Chem.*, 2005, 44, 9690–9703.
- 26 S.-W. Lai, M. C.-W. Chan, T.-C. Cheung, S.-M. Peng and C.-M. Che, *Inorg. Chem.*, 1999, 38, 4046–4055.
- 27 J. H. K. Yip, Suwarno and J. J. Vittal, *Inorg. Chem.*, 2000, 39, 3537–3543.
- 28 S.-W. Lai and C.-M. Che, *Top. Curr. Chem.*, 2004, **241**, 27–63.
- 29 A. S. Ionkin, W. J. Marshall and Y. Wang, Organometallics, 2005, 24, 619–627.
- 30 S. C. F. Kui, I. H. T. Sham, C. C. C. Cheung, C.-W. Ma, B. Yan, N. Zhu, C.-M. Che and W.-F. Fu, *Chem. Eur. J.*, 2007, 7, 417–435.
- 31 J. Schneider, P. Du, X. Wang, W. W. Brennessel and R. Eisenberg, *Inorg. Chem.*, 2009, 48, 1498–1506.
- 32 Y. Chi and P.-T. Chou, Chem. Soc. Rev., 2010, 39, 638-655.
- 33 Z. Wang, E. Turner, V. Mahoney, S. Madakuni, T. Groy and J. Li, *Inorg. Chem.*, 2010, 49, 11276–11286.
- 34 W. Wu, H. Guo, S. Ji and J. Zhao, *Inorg. Chem.*, 2011, **50**, 11446–11460.
- 35 D. P. Rillema, A. J. Cruz, C. Moore, K. Siam, A. Jehan, D. Base, T. Nguyen and W. Huang, *Inorg. Chem.*, 2012, 52, 596–607.
- 36 A. Tronnier, A. Risler, N. Langer, G. Wagenblast, I. Münster and T. Strassner, *Organometallics*, 2012, 31, 7447–7452.
- 37 H. Uesugi, T. Tsukuda, K. Takao and T. Tsubomura, *Dalton Trans.*, 2013, **42**, 7396–7403.
- 38 J. Moussa, T. Cheminel, G. R. Freeman, L.-M. Chamoreau, J. A. G. Williams and H. Amouri, *Dalton Trans.*, 2014, 43, 8162–8165.
- 39 W. Wu, W. Wu, S. Ji, H. Guo, P. Song, K. Han, L. Chi, J. Shao and J. Zhao, J. Mater. Chem., 2010, 20, 9775– 9786.
- 40 Y. Zhang, J. A. Garg, C. Michelin, T. Fox, O. Blacque and K. Venkatesan, *Inorg. Chem.*, 2011, 50, 1220–1228.
- 41 Y. Zhang, J. Clavadetscher, M. Bachmann, O. Blacque and K. Venkatesan, *Inorg. Chem.*, 2014, 53, 756–771.
- 42 Y. Unger, D. Meyer, O. Molt, C. Schildknecht, I. Münster, G. Wagenblast and T. Strassner, *Angew. Chem., Int. Ed.*, 2010, 49, 10214–10216.
- 43 Y. Unger, A. Zeller, M. A. Taige and T. Strassner, *Dalton Trans.*, 2009, 4786–4794, DOI: 10.1039/b900655a.
- 44 W. Lu, M. C. W. Chan, Z. Hui, C.-M. Che, N. Zhu and S.-T. Lee, *J. Am. Chem. Soc.*, 2004, **126**, 4958–4971.
- 45 W. Lu, M. C. W. Chan, N. Zhu, C.-M. Che, C. Li and Z. Hui, *J. Am. Chem. Soc.*, 2004, **126**, 7639–7651.

46 C. E. Whittle, J. A. Weinstein, M. W. George and K. S. Schanze, *Inorg. Chem.*, 2001, **40**, 4053–4062.

- 47 M. Hissler, W. B. Connick, D. K. Geiger, J. E. McGarrah, D. Lipa, R. J. Lachiotte and R. Eisenberg, *Inorg. Chem.*, 2000, 39, 447–457.
- 48 T. J. Wadas, R. J. Lachiotte and R. Eisenberg, *Inorg. Chem.*, 2003, 42, 3772–3778.
- 49 K.-Y. Kim, S. Liu, M. E. Köse and K. S. Schanze, *Inorg. Chem.*, 2006, 45, 2509–2519.
- 50 F. N. Castellano, I. E. Pomestchenko, E. Shikhova, F. Hua, M. L. Muro and N. Rajapakse, *Coord. Chem. Rev.*, 2006, 250, 1819–1828.
- 51 C.-H. Tao and V. W.-W. Yam, *J. Photochem. Photobiol., C*, 2009, **10**, 130–140.
- 52 R. Saha, M. A. Qaium, D. Debnath, M. Younus, N. Chawdhury, N. Sultana, G. Kociok-Kohn, L.-l. Ooi, P. R. Raithby and M. Kijima, *Dalton Trans.*, 2005, 2760–2765, DOI: 10.1039/b505484b.
- 53 W. Wu, J. Zhao, J. Sun, L. Huang and X. Yi, *J. Mater. Chem. C*, 2013, **1**, 705–716.
- 54 I. Fratoddi, C. Battocchio, A. Furlani, P. Mataloni, G. Polzonetti and M. V. Russo, *J. Organomet. Chem.*, 2003, **674**, 10–23.
- 55 K. Gagnon, S. M. Aly, A. Brisach-Wittmeyer, D. Bellows, F.-F. Bérubé, L. Caron, A. S. Abd-El-Aziz, D. Fortin and P. D. Harvey, *Organometallics*, 2008, 27, 2201–2214.
- 56 L. Liu, D. Fortin and P. D. Harvey, *Inorg. Chem.*, 2009, 48, 5891–5900.
- 57 D. Bellows, S. M. Aly, C. P. Gros, M. El Ojaimi, J.-M. Barbe, R. Guilard and P. D. Harvey, *Inorg. Chem.*, 2009, 48, 7613–7629.
- 58 M. S. Khan, M. R. A. A. Al-Mandhary, M. K. Al-Suti, F. R. Al-Battashi, S. Al-Saadi, B. Ahrens, J. K. Bjernemose, M. F. Mahon, P. R. Raithby, M. Younus, N. Chawdhury, A. Köhler, E. A. Marseglia, E. Tedesco, N. Feeder and S. J. Teat, *Dalton Trans.*, 2004, 2377–2385.
- 59 N. J. Long, C. K. Wong and A. J. P. White, *Organometallics*, 2006, 25, 2525–2532.
- 60 L. Liu, W.-W. Wong, J.-X. Shi, K.-W. Cheah, T.-H. Lee and L. M. Leung, J. Organomet. Chem., 2006, 691, 4028– 4041.
- 61 D. Fortin, S. Clément, K. Gagnon, J.-F. Bérubé, M. P. Stewart, W. E. Geiger and P. D. Harvey, *Inorg. Chem.*, 2009, 48, 446–454.
- 62 J. M. Keller, K. D. Glusac, E. O. Danilov, S. McIlroy, P. Sreearuothai, A. R. Cook, H. Jiang, J. R. Miller and K. S. Schanze, J. Am. Chem. Soc., 2011, 133, 11289–11298.
- 63 J. R. Berenguer, E. Lalinde and M. Teresa Moreno, *Coord. Chem. Rev.*, 2010, **254**, 832–875.
- 64 J. E. Rogers, J. E. Slagle, D. M. Krein, A. R. Burke, B. C. Hall, A. Fratini, D. G. McLean, P. A. Fleitz, T. M. Cooper, M. Drobizhev, N. S. Makarov, A. Rebane, K.-Y. Kim, R. Farley and K. S. Schanze, *Inorg. Chem.*, 2007, 46, 6483–6494.
- 65 A. A. Rachford, S. Goeb and F. N. Castellano, *J. Am. Chem. Soc.*, 2008, **130**, 2766–2767.

- 66 L. Liu, D. Huang, S. M. Draper, X. Yi, W. Wu and J. Zhao, Dalton Trans., 2013, 42, 10694–10706.
- 67 H. Guo, M. L. Muro-Small, S. Ji, J. Zhao and F. N. Castellano, *Inorg. Chem.*, 2010, **49**, 6802–6804.
- 68 M.-H. Nguyen and J. H. K. Yip, *Organometallics*, 2012, 31, 7522-7531.
- 69 M.-H. Nguyen, V. H. Nguyen and J. H. K. Yip, *Organometallics*, 2013, **32**, 7283–7291.
- 70 D. N. Kozhevnikov, V. N. Kozhevnikov, M. Z. Shafikov, A. M. Prokhorov, D. W. Bruce and J. A. Gareth Williams, *Inorg. Chem.*, 2011, 50, 3804–3815.
- 71 Y. Liu, H. Guo and J. Zhao, *Chem. Commun.*, 2011, 47, 11471–11473.
- 72 Y. Y. Chia and M. G. Tay, *Dalton Trans.*, 2014, 43, 13159–13168.
- 73 J. Hu, R. Lin, J. H. K. Yip, K.-Y. Wong, D.-L. Ma and J. J. Vittal, *Organometallics*, 2007, 26, 6533– 6543.
- 74 A. Singh and P. R. Sharp, J. Am. Chem. Soc., 2006, 128, 5998-5999.
- 75 H. B. Lee and P. R. Sharp, Organometallics, 2005, 24, 4875–4877.
- 76 S. Lentijo, G. Aullon, J. A. Miguel and P. Espinet, *Dalton Trans.*, 2013, 42, 6353–6365.
- 77 M.-H. Nguyen and J. H. K. Yip, Organometallics, 2010, 29, 2422–2429.
- 78 W. Y. Heng, J. Hu and J. H. K. Yip, *Organometallics*, 2007, **26**, 6760–6768.
- 79 B.-Y. Wang, A. R. Karikachery, Y. Li, A. Singh, H. B. Lee, W. Sun and P. R. Sharp, *J. Am. Chem. Soc.*, 2009, 131, 3150–3151.
- S. Lentijo, J. A. Miguel and P. Espinet, *Inorg. Chem.*, 2010,
 49, 9169–9177.
- 81 W. G. Herkstroeter, A. A. Lamola and G. S. Hammond, J. Am. Chem. Soc., 1964, 86, 4537–4540.
- 82 D. E. Damschen, C. D. Merritt, D. L. Perry,
 G. W. Scott and L. D. Talley, *J. Phys. Chem.*, 1978, 82, 2268–2272.
- 83 R. W. Anderson Jr., R. M. Hochstrasser, H. Lutz and G. W. Scott, *Chem. Phys. Lett.*, 1974, **28**, 153–157.
- 84 R. M. Hochstrasser, H. Lutz and G. W. Scott, *Chem. Phys. Lett.*, 1974, **24**, 162–167.
- 85 D. Wöll, J. Smirnova, W. Pfleiderer and U. E. Steiner, *Angew. Chem., Int. Ed.*, 2006, **45**, 2975–2978.
- 86 D. Wöll, S. Laimgruber, M. Galetskaya, J. Smirnova, W. Pfleiderer, B. Heinz, P. Gilch and U. E. Steiner, *J. Am. Chem. Soc.*, 2007, 129, 12148–12158.
- 87 G. Amirzadeh and W. Schnabel, *Makromol. Chem.*, 1981, 182, 2821–2835.
- 88 M. G. Neumann, M. H. Gehlen, M. V. Encinas, N. S. Allen, T. Corrales, C. Peinado and F. Catalina, *J. Chem. Soc., Faraday Trans.*, 1997, 93, 1517–1521.
- 89 M. Aydin, N. Arsu, Y. Yagci, S. Jockusch and N. J. Turro, *Macromolecules*, 2005, 38, 4133–4138.
- 90 L. Cokbaglan, N. Arsu, Y. Yagci, S. Jockusch and N. J. Turro, *Macromolecules*, 2003, 36, 2649–2653.

91 T. Corrales, C. Peinado, F. Catalina, M. G. Neumann, N. S. Allen, A. M. Rufs and M. V. Encinas, *Polymer*, 2000, 41, 9103–9109.

Dalton Transactions

- 92 X. Jiang and J. Yin, Macromol. Chem. Phys., 2008, 209, 1593–1600.
- 93 H. Tar, D. Sevinc Esen, M. Aydin, C. Ley, N. Arsu and X. Allonas, *Macromolecules*, 2013, 46, 3266–3272.
- 94 J. C. Dalton and F. C. Montgomery, *J. Am. Chem. Soc.*, 1974, **96**, 6230–6232.
- 95 C. Ley, F. Morlet-Savary, P. Jacques and J. P. Fouassier, Chem. Phys., 2000, 255, 335–346.
- 96 T.-i. Lai and E. C. Lim, Chem. Phys. Lett., 1980, 73, 244– 248.
- 97 X. Allonas, C. Ley, C. Bibaut, P. Jacques and J. P. Fouassier, *Chem. Phys. Lett.*, 2000, 322, 483–490.
- 98 P. K. Monaghan and R. J. Puddephatt, *Organometallics*, 1985, 4, 1406–1412.
- 99 A. Abo-Amer, M. S. McCready, F. Zhang and R. J. Puddephatt, *Can. J. Chem.*, 2011, **90**, 46–54.
- 100 G. W. Parshall, J. Am. Chem. Soc., 1966, 88, 704-708.
- 101 A. Pidcock, R. E. Richards and L. M. Venanzi, *J. Chem. Soc. A*, 1966, 1707–1710, DOI: 10.1039/J19660001707.
- 102 J. A. Pople and D. P. Santry, Mol. Phys., 1964, 8, 1-18.
- 103 T. Sajoto, P. I. Djurovich, A. Tamayo, M. Yousufuddin, R. Bau, M. E. Thompson, R. J. Holmes and S. R. Forrest, *Inorg. Chem.*, 2005, 44, 7992–8003.
- 104 S. K. Doorn, R. B. Dyer, P. O. Stoutland and W. H. Woodruff, J. Am. Chem. Soc., 1993, 115, 6398–6405.
- 105 D. Burget and P. Jacques, J. Lumin., 1992, 54, 177-181.
- 106 D. W. Wöll, PhD Dissertation, University of Konstanz, 2006.
- 107 T. Yoshida, T. Matsuda, S. Otsuka, G. W. Parshall and W. G. Peet, in *Inorganic Syntheses*, John Wiley & Sons, Inc., 2007, ch. 32, pp. 122–123, DOI: 10.1002/9780470132593.
- 108 G. M. Sheldrick, SHELXL-97, Program for Crystal Structure Solution and Refinement, Universität Göttingen, 1997.
- 109 L. J. Farrugia, J. Appl. Crystallogr., 2013, 45, 849-854.
- 110 M. J. Frisch, G. 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. Blonio, 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, Т. J. A. Montgomery Jr., J. E. Peralta, 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, Yazyev, A. J. Austin, R. Cammi, C. Pomelli, 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, Gaussian09, Version 8.0, Gaussian Inc., Wallingford CT, 2009.
- 111 O. Gunnarsson and B. I. Lundqvist, *Phys. Rev. B: Solid State*, 1976, **13**, 4274.
- 112 W. Küchle, M. Dolg, H. Stoll and H. Preuss, *J. Chem. Phys.*, 1994, **100**, 7535–7532.
- 113 M. Dolg, H. Stoll and H. Preuss, *J. Chem. Phys.*, 1989, **90**, 1730–1734.
- 114 D. Andrae, U. Haeussermann, M. Dolg, H. Stoll and H. Preuss, *Theor. Chim. Acta*, 1990, 77, 123.
- 115 P. H. Hariharan and J. A. Pople, *Theor. Chim. Acta*, 1973, 28, 213–222.
- 116 J. P. Perdew, K. Burke and M. Enzerhof, *Phys. Rev. Lett.*, 1996, 77, 3865–3868.
- 117 C. Adamo and V. Barone, *J. Chem. Phys.*, 1999, **110**, 6158–6170.
- 118 E. Cancés, B. Menucci and J. Tomasi, J. Chem. Phys., 1997, 107, 3032–3041.
- 119 B. Mennucci and J. Tomasi, *J. Chem. Phys.*, 1997, **106**, 5151–5158.
- 120 M. Cossi, N. Rega, G. Scalmani and V. Barone, *J. Comput. Chem.*, 2003, 24, 669–681.
- 121 G. Scalamani and M. J. Frisch, *J. Chem. Phys.*, 2010, **132**, 114110–114124.
- 122 E. Runge and K. U. G. E, *Phys. Rev. Lett.*, 1984, **52**, 997–1000.
- 123 N. M. O'Boyle, A. L. Tenderholt and K. M. Langner, *J. Comput. Chem.*, 2008, **29**, 839–845.
- 124 T. Keith and J. Millam, *GaussView, Version 3*, Semichem Inc., Shawnee Mission KS, 2009.