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Dinuclear platinum(\parallel) complexes emitting through TADF: new ligand design to minimise aggregation and the S_1-T_1 energy gap†

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Dinuclear platinum(III) complexes of a new, ditopic, bis-tridentate NCN-NCN-coordinating ligand, appended with four mesityl groups, are reported. The high radiative rate constants and correspondingly efficient luminescence of the complexes involves thermally activated delayed fluorescence (TADF), thanks to a near-zero energy gap between the S_1 and T_1 states. The mesityl groups also serve to hinder the aggregation that was detrimental to electroluminescence efficiency in previous studies, allowing a \sim 4-fold increase in OLED efficiency to be achieved (i.e. from 2.3% previously to 10% in this work). Oxidation of one of the Pt(II) complexes led to a dinuclear Pt(IV) complex of unprecedented structure.

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Introduction

Organoplatinum(II) complexes are widely used as the luminophore or sensitizer in diverse applications, often complementing their iridium(III) counterparts. 1-3 Important uses include bioimaging, 4-9 photodynamic therapy, 10 photocatalysis, 11-14 and organic light-emitting diodes (OLEDs). 15-18 The luminescence displayed by such complexes is normally considered phosphorescence, whereby the high spin-orbit coupling associated with the heavy metal relaxes the spin selection rule, accelerating the rate of the otherwise forbidden $T_1 \rightarrow S_0$ process.^{2,19} However, recent findings by some of the present authors have revealed that another mechanism may be at work in at least some such complexes, involving thermally activated delayed fluorescence (TADF).20-22 Dinuclear Pt(II) complexes of ditopic ligands featuring a pyrimidine bridge were found to have a small energy gap $\Delta E_{\rm ST}$ between the lowest singlet S₁ and triplet T1 excited states.21 It leads to a shortening of the decay lifetime by thermal activation of T_1 to S_1 and subsequent emission through the allowed $S_1 \rightarrow S_0$ transition (i.e., following

In the present work, we took our original ligand design and appended it with four 2,4,6-trimethylphenyl (mesityl) groups, with a view to improving the solubility of the resulting complex and reducing aggregation. We also show how this seemingly otherwise insignificant structural change leads to a reduction in $\Delta E_{\rm ST}$, which in turn enhances the TADF contribution to emission and substantially improves performance.

Results and discussion

Synthesis

The new ditopic NCN–NCN proligand H_2L (compound 4 in Scheme 1) was prepared by a sequence of Pd-catalysed cross-coupling reactions from readily obtainable starting materials; full experimental details and the characterisation of new compounds are given in the ESI.† Compound H_2L was platinated using K_2PtCl_4 in acetic acid to give the dinuclear complex $L(Pt-Cl)_2$ (5), from which the iodo derivative $L(Pt-I)_2$ (6) was prepared by metathesis of the monodentate ligand upon treatment with $Ag(SO_3CF_3)$ followed by KI. We were unable to obtain crystals of 5 or 6 of sufficient quality for an X-ray diffraction study, but the oxidation of 5 with PhICl₂ led cleanly to

the TADF model). 23,24 A similar phenomenon has since been observed by others in a mononuclear $Pt(\pi)$ complex, 25 and the future discovery of many further examples seems likely. While preparing this manuscript, another work reporting both mono- and dinuclear $Pt(\pi)$ complexes showing TADF has been published, reporting high OLED efficiency and low efficiency roll-off. This further underscores the potential advantages that TADF complexes of this metal offer.

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$$\begin{array}{c} B_{1} \\ B_{2} \\ B_{3} \\ B_{4} \\ B_{5} \\ B_{7} \\ B_{7} \\ B_{8} \\ B_{8} \\ B_{8} \\ B_{7} \\ B_{8} \\ B_{9} \\ B_{1} \\ B_{2} \\ B_{2} \\ B_{1} \\ B_{2} \\ B_{2} \\ B_{1} \\ B_{2} \\$$

Scheme 1 Synthetic procedure for complexes 5, 6 and 7: (i) toluene/EtOH/H₂O (2:1:1 v/v), Pd(PPh₃)₄, Na₂CO₃, 70 °C (2 h) \rightarrow 90 °C (18 h), 35%; (ii) toluene, Pd(PPh₃)₄, Cs₂CO₃, 90 °C (18 h), 67%; (iii) dioxane, Pd(dppf)Cl₂·CH₂Cl₂, KOAc, 90 °C (18 h); (iv) toluene, Pd(PPh₃)₄, Cs₂CO₃, 90 °C (18 h), 35%; (v) AcOH, reflux, 65%; (vi) step 1: acetone, RT (1.5 h); step 2: RT (2 h); 67%; (vii) CHCl₃, RT (18 h), 78%. Inset: The previously studied complexes 8 and 9 incorporating a related NCN-NCN-coordinating ligand.

L(Pt-Cl₃)₂ (7), a dinuclear Pt(IV) complex of unprecedented structure that was amenable to crystallography. Although the thrust of the current work is the Pt(II) systems, there is growing interest in the less widely explored +4 oxidation state, ^{27,28} and 7 represents an interesting structure in that context for future elaboration by replacement of the chloride ligands.²⁹ Here, it offers insight into the likely structures of 5 and 6. The structure (Fig. 1) shows the two Pt centres in very similar pseudooctahedral environments, each bound to an NCN unit with three chlorides completing the coordination sphere. The torsion angles between the mesityl and phenyl rings are in the range 69 \pm 3° and, between the pyridyl and mesityl rings, 69 \pm 5° . These angles are a little less than that of 80° in the related mesityl-appended NCHN proligand, 30 suggesting a slightly greater degree of conjugation across the rings upon complexation, attenuating the steric preference for orthogonality.

EQE = 10.0%

DFT and TD-DFT calculations

The excited states in 5 and 6 were probed by DFT/TD-DFT at the B3LYP/def2-SVP level of theory using ORCA 5.0.3, 31,32 and by quasi-degenerate perturbation theory (QDPT). 33,34 The opti-

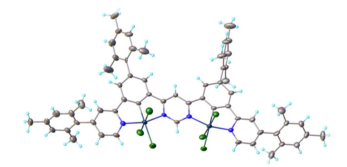


Fig. 1 The molecular structure of the dinuclear Pt(IV) complex L(Pt-Cl₃)₂, 7.

mised T_1 geometries (Fig. 2) show torsions between the plane of the NCN-NCN unit and the pendent mesityl rings that are similar to those observed experimentally in the Pt(IV) complex (atomic coordinates for T_1 and S_0 are given as separate ESI†). The mesityl groups thus form a congested shield around the complex, which is expected to inhibit intermolecular interactions between NCN-NCN planes.

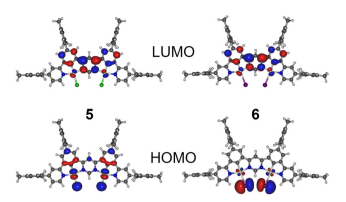


Fig. 2 HOMO and LUMO of complexes 5 and 6 calculated using the B3LYP/def2-SVP/B3LYP/def2-TZVP level of theory.

The frontier molecular orbital isosurfaces of 5 and 6 (Fig. 2) resemble those of 8 and 9,21,22 with the LUMO distributed over the pyrimidine linker and the neighbouring benzene rings, but also including a small admixture of d orbitals from both Pt(II) centres. The HOMO comprises d orbitals of both the Pt centres and p orbitals of their respective monodentate halide ligands. For 6, there are no significant contributions from other parts of the molecule, such that the degree of HOMO-LUMO overlap is small. In 5, the HOMO includes additional contributions from π orbitals of the organic ligand, which serves to increase the HOMO-LUMO overlap. In both complexes, the S_1 and T_1 are associated mainly with the HOMO \rightarrow LUMO transition, and thus may be classed as predominantly MXLCT or metal-halogen-to-ligand charge-transfer in character. Complex 6 appears to display a stronger MXLCT character of the excited state, while 5 shows a rather MXLCT + LC (LC ligand-centred) character due to a clear contribution of the NCN-NCN ligand to HOMO.

When spin–orbit coupling (SOC) is included in the calculations, excited states of predominantly triplet or singlet nature are identified (Fig. S4.1, Tables S4.1 and S4.2†). The lowest triplet state splits into three closely separated levels, the separation between the first and third, ΔE_{1-3} , being the zero-field splitting (ZFS). Values of 2 meV (13 cm⁻¹) and 15 meV (119 cm⁻¹) are calculated for 5 and 6 respectively. In 5, state 7 is the first excited state with predominant singlet character (68%; states 4–6 are associated with T₂); thus $\Delta E_{\rm ST} = \Delta E_{1-7}$, which is calculated to be 291 meV (2345 cm⁻¹). For 6, the first predominantly singlet excited state (81.4%) is state 4, and $\Delta E_{\rm ST} = \Delta E_{1-4} = 19$ meV (149 cm⁻¹). The trend to smaller $\Delta E_{\rm ST}$ upon changing X from Cl to I is consistent with that observed previously for 8 and 9.

Solution-state photophysics

The solvent of choice for photophysical measurements is toluene (PhMe) (based on **9** having previously shown the highest radiative rate in this solvent²²). Here, we studied **6** in toluene, but **5** has poorer solubility and intermolecular interactions suppress its emission leading to a lower Φ_{PL} . We believe that the larger diameter of the X = I as compared to X =

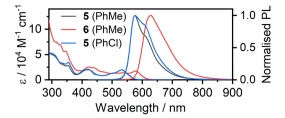


Fig. 3 Absorption and photoluminescence spectra of 5 and 6 in dilute ($c=10^{-5}$ M) toluene (PhMe) and (for 5) chlorobenzene (PhCl) solutions.§¶.

Cl hinders close $\pi\cdots\pi$ and $\text{Pt}\cdots\text{Pt}$ intermolecular interactions, leading to the better overall solubility of **6**. This is in line with our previous work where the X = I auxiliary ligand was found to disrupt aggregation of the closely related mono-Pt(II) complexes of the Pt(NCN)-X type. The full characterisation of **5** was therefore performed in chlorobenzene (PhCl), where such interactions were essentially absent. There is substantial overlap between the photoluminescence (PL) spectrum and the lowest-energy absorption band in both complexes (Fig. 3) – a clear indication that the PL originates not from the lowest-energy T_1 state but rather from a higher state, and a strong clue for the involvement of TADF.

The PL spectrum of **6** is very similar to that of the analogous complex **9** ($\lambda_{\text{max}}^{\text{PL}} = 627$ and 628 nm respectively), ²² but the emission spectrum of **5** ($\lambda_{\text{max}}^{\text{PL}} = 576$ nm) is drastically different from that of **8** ($\lambda_{\text{max}}^{\text{PL}} = 617$ nm), Fig. 4. The difference between the PL of **5** and **8** in chlorobenzene can be attributed to a significantly smaller ΔE_{ST} in the former with TADF dominating the spectrum at RT as opposed to **8** with phosphorescence being the dominant component. As the PL onsets are similar in either case, the S₁ energy is likely comparable in both complexes, but the structural differences between the two luminophores lead to a higher T₁ energy in **5** (Table S5.2†).

The PL spectra of the complexes in solution are sensitive to temperature (Fig. S5.1 and S5.2†), in line with the TADF mechanism. For 5 in PhCl, the interplay between phosphorescence and TADF at intermediate temperatures is evident, as in **8**. For **6**, however, the PL spectrum blue shifts very slightly at lower temperatures, indicating a visible influence of suppressed molecular motion. Even at 160 K, there is no evidence of phosphorescence (in stark contrast to the behaviour of **9** for example), suggesting that $\Delta E_{\rm ST}$ in **6** is very small indeed, with TADF consequently predominating even at low temperatures.

Complexes 5 and 6 display unusually large radiative decay rate constants $k_{\rm r}$ for platinum(II) complexes: $\tau = 0.34~\mu s$, $\Phi_{\rm PL} =$

 \S A slight difference between the PL spectra of 5 in toluene and chlorobenzene can be ascribed to a variation in the $\Delta E_{\rm ST}$, in line with the previous report of complex 8. ²¹ Here, 5 displays a larger $\Delta E_{\rm ST}$ in the higher polarity chlorobenzene than in toluene, leading to a visible phosphorescence contribution in chlorobenzene, manifest on the low-energy side of the spectrum.

¶Complex 7 does not show any detectable emission, as expected based on earlier studies. 28,29 The absorption spectrum of 7 is included in the ESI for completeness (Fig. S5.3†).

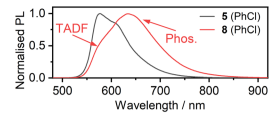


Fig. 4 Comparison of photoluminescence spectra of compounds 5 and 8 recorded in dilute PhCl.

0.11, $k_{\rm r}=3.3\times10^5~{\rm s}^{-1}$ for 5 in PhMe [$\tau=2.1~{\rm \mu s},~\Phi_{\rm PL}=0.45,~k_{\rm r}=2.1\times10^5~{\rm s}^{-1}$ in PhCl]; $\tau=0.40~{\rm \mu s},~\Phi_{\rm PL}=0.23,~k_{\rm r}=5.7\times10^5~{\rm s}^{-1}$ for 6 in PhMe.|| The $k_{\rm r}$ values are significantly higher than for 8 and 9 (Table S5.1†), suggesting more efficient TADF. We use the method of Strickler and Berg³⁶ to estimate the singlet-state radiative rate constants $k_{\rm r}^S$ and $f({\rm S}_1\to{\rm S}_0)$ oscillator strengths, from the lowest energy absorption bands. ^{37,38} For 5 in PhCl, $k_{\rm r}^S=5.3\times10^7~{\rm s}^{-1}$ and $f({\rm S}_1\to{\rm S}_0)=0.12$; corresponding values for 6 in PhMe are $3.3\times10^7~{\rm s}^{-1}$ and 0.09. The values are two orders of magnitude higher than the experimental values, showing (as expected) that the emission is not fluorescence, but rather pointing to the likely involvement of TADF.

Solid-state photophysics

More conclusive evidence of a TADF mechanism comes from the temperature dependence of the complexes' PL spectra in a dilute polystyrene (PS) film, over the range 295-80 K (Fig. 5). The spectra display a clear change, shifting to the red as the temperature decreases, from which it is apparent that TADF dominates at 295 K, while phosphorescence from the lowerenergy T₁ state dominates at 80 K. The evolution of the spectra correlates with a change in the radiative lifetime (Fig. 6, Fig. S5.4 and S5.5†). Fitting of the lifetime data of 6 to an established Boltzmann-based expression (eqn (S2)†) gives a natural fluorescence lifetime $\tau_0 \approx 32$ ns, phosphorescence lifetime $\tau_{\rm PH} \approx 13$ µs, and $\Delta E_{\rm ST} \approx 66$ meV, not dissimilar to the value calculated by TD-DFT. The $k_r^{\rm S}$ figures for both complexes $(\sim 10^7 \text{ s}^{-1})$ are in good agreement with the estimates obtained using the Strickler-Berg method. It is striking that the radiative lifetime at 295 K is an order of magnitude shorter than at 80 K thanks to the involvement of TADF, underscoring how TADF can benefit Pt(II) complexes despite the heavy atom effect that might otherwise be expected to funnel emission through phosphorescence.

OLED devices

Owing to the good PL properties of 6 and its low susceptibility to aggregative quenching (in contrast to 5), it was tested as an emitter in a solution-processed organic light-emitting diode (OLED). Three pairs of device architectures (Tables S6.1 and

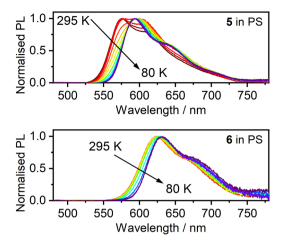


Fig. 5 PL spectra of 5 and 6 in polystyrene dispersion (c = 0.2% w/w) over the temperature range indicated.

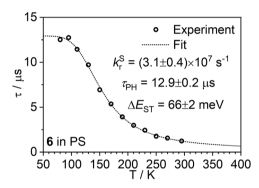


Fig. 6 Variation of the PL decay lifetime τ of 6 as a function of temperature T.

S6.3†) were trialled to optimise the efficiency: Devs 1, 3, and 5 at 3% w/w loading of the complex in the host, and Devs 2, 4, and 6 at 5%, to probe the effect of concentration. The pair of devices 1 and 2 use a simple structure with a relatively thick TCTA:PO-T2T host emissive layer.³⁹ Devices 3 and 4 feature a mCP:PO-T2T host. Devices 5 and 6 employ a mCP:PBD host with a poly(vinyl-carbazole) hole-transporting and electron-blocking layer (as previously used with 9 and hence serving as a reference²²). The OLED data are presented in Fig. 7, Fig. S6.1–S6.8 and Table S6.1.†

Comparison of Devices 5 and 6 with those reported previously using 9 shows that the use of the new complex leads to a \sim 4-fold higher EQE (up to 10%) than complex 9, and a \sim 2-4-fold higher luminance (up to 8700 cd m $^{-2}$). Although several factors may have contributed to this result, the main one appears to be suppression of emitter aggregation in the solid state by the mesityl substituents. This effect can be identified from a comparison of the respective EL spectra (Fig. 8). Despite identical emitter loading and OLED structure, Device 6 displays a visibly narrower EL spectrum, lacking the long wavelength tail >700 nm that the previously reported Device 7

 $[\]parallel k_{\rm r}$ is estimated from the lifetime and quantum yield, assuming unit population of the emissive state, where $k_{\rm r}=\Phi/\tau$. The quoted τ and Φ values were measured in deoxygenated solution at 295 K.

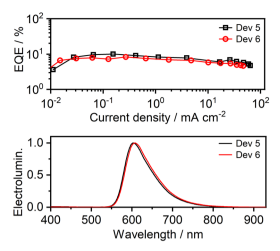


Fig. 7 Characteristics of OLEDs 5 and 6: (top) External quantum efficiency (EQE); (bottom) EL spectra.

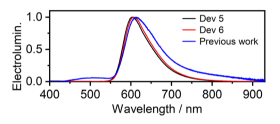


Fig. 8 Comparison of the EL spectra of OLEDs 5 and 6 with the EL spectrum of OLED 7 (using complex 9) from previous work.²²

shows (using complex 9). That tail is due to dimeric or oligomeric species formed through aggregation; it is detrimental both to the overall OLED EQE and to colour purity.** Thus, the encumbered molecular design is successful in promoting efficiency in OLEDs using dinuclear Pt(II) TADF emitters.

Conclusions

In conclusion, the design of ligand 4 – featuring a ditopic *NCN-NCN* core decorated with four mesityl units – successfully limits aggregation of complex 6, inhibiting the detrimental effect on EL in an OLED. Complex 6 is highly soluble in toluene and other solvents and displays a lower susceptibility to aggregation in the solid state than the previously reported complex 9. The hexyl chains and t-butyl substituents in 9 are insufficient to impede intermolecular interactions, whereas the mesityl groups of 6 do so. The solution-processed OLEDs reach an EQE of 10% with a maximum luminance of 5400 cd m⁻² (Dev 5), or EQE of 9.2% with a maximum luminance of 8700 cd m⁻² (Dev 3). These figures exceed by \sim 4-fold the

values reported for the structurally similar complex **9**, and are the highest values reported for a TADF-based Pt(π) emitter. Moreover, the use of ligand **4** also leads to a significant reduction of the $\Delta E_{\rm ST}$ (*e.g.*, to 0.07 eV in **5** in PS, compared to 0.20 eV reported for complex **8**). This is achieved by increasing the T₁ energy in complex **5**.

The high solubility attained using ligand 4 could also be exploited to oxidise the dinuclear Pt(II) 5 complex to the corresponding dinuclear Pt(IV) complex 7. Analogous attempts to prepare such materials from related, less substituted *NCN-NCN* ligands have led to intractable mixtures, probably due to poor solubility.

In summary, this study has shown how modifying the design of rigid, ditopic, bis-tridentate ligands can simultaneously lead to improvements in several properties of the corresponding Pt_2 complexes, including enhanced solubility, lower propensity to aggregation, and a reduction in $\Delta E_{\rm ST}$ that serves to accelerate radiative decay through TADF. We expect TADF to be implicated in the emission of many other $3^{\rm rd}$ row phosphors and that these findings will help inform the future design of such molecules.

Author contributions

P. P. – Conceptualization, funding acquisition, project administration, formal analysis, investigation (synthesis, photophysics, OLED devices, computations), visualization, writing – original draft, writing – review & editing; Y. M. D. – investigation (synthesis), writing – review & editing; E. V. P. – conceptualization, investigation (synthesis), writing – review & editing; J. A. G. W. – conceptualization, funding acquisition, resources, project administration, supervision, writing – original draft, writing – review & editing.

Data availability

Our supporting research data is available from the Durham Research Online DATAsets Archive (DRO-DATA) open data repository. DOI: https://doi.org/10.15128/r2tt44pm914.

Conflicts of interest

There are no conflicts of interest to declare.

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^{**}Some deleterious interactions nevertheless remain, as the 3%-loaded devices are generally more efficient than the 5% devices, a trend also widely observed with, for example, Ir(III) emitters.

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