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Synthesis and redox properties of mono-, di- and tri-metallic Platinum-ethynyl complexes based on the trans-Pt(C₆H₄N{C₆H₄OCH₃-4}₂)(C≡CR)(PPh₃)₂ motif.†

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† Dedicated to Professor Claude Lapinte, in recognition of his outstanding career,  
and with thanks for his generosity and kindness over many years of fruitful and  
enjoyable collaboration.
Abstract

The Pt-halide complex trans-PtI\{C_6H_4NAr_2\}(PPh_3)_2 (Ar = C_6H_4OMe-4, 3) was prepared by oxidative addition of N(C_6H_I)Ar_2 (2) to Pt(PPh_3)_4. Reactions of trans-PtI\{C_6H_4NAr_2\}(PPh_3)_2 (3) with 1-alkynes under CuI catalysed dehydrohalogenation conditions allows the preparation of a range of platinum ethynyl compounds containing up to four redox-active triarylamine centres. The compounds trans-Pt(C≡CAr)(C_6H_4NAr_2)(PPh_3)_2 (4a), trans-Pt(C≡CC_6H_4NAr_2)(C_6H_4NAr_2)(PPh_3)_2 (4b), \{trans-Pt(C_6H_4NAr_2)(PPh_3)_2\}_2(\mu-C≡C-1,4-C_6H_4C≡C) (5) and N\{C_6H_4C≡CPt(C_6H_4NAr_2)(PPh_3)_2\}_3 (6) undergo a single electrochemical event for each chemically distinct type of triarylamine in the molecular backbone. The complete reversibility of the larger systems means that they can be used for charge storage materials capable of releasing up to four electrons. A combination of electrochemical, spectroelectrochemical and quantum chemical analyses reveal weak electronic coupling between the amine moieties in the redox products derived from one-electron oxidation.

Keywords mixed-valence; spectroelectrochemistry; DFT; triarylamine; platinum

Introduction

Complexes of general form PtX(Ar')(PR_3)_2 (X = halide, pseudo halide; Ar' = aryl; R = aromatic, alkyl) have a long history, and can be prepared by numerous synthetic methods, including reactions of [PtX_2\{bis(olefin)\}] with aryl Grignard reactions and phosphine,[1-3] or Pt(X)_2(PR_3)_2 with ArLi,[4] by disproportionation reactions of PtCl(MPh_2)(PPh_3)_2 (M = Sn, Pb),[5] reaction of cis-PtPh(PPh_3)_2(PbPh_3) with Br_2 or HBr,[6] decarbonylation of PtX(COAr)(PR_3)_2,[7] ligand exchange reactions of {Pt(μ-Cl(tht))(C_6F_5)_2 \[8\], and perhaps most simply by oxidative addition of arylhalides to
Pt(PR$_3$)$_n$ (R = Ph, Et; n = 3, 4)[9-13]. These aryl platinum(II) complexes have served as models through which to explore key reaction steps in various catalytic transformations,[14-17] and as structurally well defined building blocks for the assembly of molecular nanostructures[18-20], scaffolds and redox-innocent end-caps for polyynes, some of quite extraordinary length.[3, 8, 21-26] Alkynyl complexes trans-Pt(C≡CR)(Ar')(PR$_3$)$_2$ have also demonstrated a range of interesting and potentially useful optoelectronic properties, leading to the design of soluble and processable materials with low band-gaps for solar cell applications,[27-34], and significant two-photon absorption cross-sections,[35], whilst the heavy atom effect leads to effective intersystem crossing[36] and efficient triplet sensitized processes such as visible light induced ring-closing of Irie-style molecular switches,[37] efficient optical limiting [38] and (electro)phosphorescence[39-42]. The preparation of a penta-(platinum alkynyl) complex of corannulene further demonstrates the vast scope for use of the trans-PtXAr'(PR$_3$)$_2$ motif to assemble and stabilise complex structures.[43]

We were attracted to the synthetic, structural and electronic properties of Pt(II) aryl complexes as potential scaffolds for the construction of structures in which two or more organic electrophores, specifically triarylamine moieties, could be linked into larger structures of well defined geometry with a view to exploring further examples of metal-bridged organic mixed-valence compounds.[44, 45] [46] We report here the synthesis of the redox-active building block trans-PtI(C$_6$H$_4$NAr$_2$)(PPh$_3$)$_2$ (3, Ar = C$_6$H$_4$OMe-4), and its use in the preparation of mono-, bi- and trimetallic complexes from CuI catalysed reactions with terminal alkynes. Electrochemical and spectroelectrochemical investigations, supported by DFT calculations on
representative examples, have been used to explore the redox chemistry and
electronic structures of these complexes and their redox products. These studies
reveal sequential oxidation of the \{Ar_2NC_6H_4^-\} and \{-C=C-C_6H_4NAr_2\} centres,
which are only weakly coupled through the \textit{trans}-Pt(PPh_3)_2 bridge.

\section*{Results and Discussion}

\textit{Syntheses} The key triarylamine building blocks 1[47] and 2 [48, 49] were assembled
using the sequence of Ullmann and iodination reactions as illustrated in Scheme 1.
These compounds were characterised in the usual manner by \(^1\)H and \(^{13}\)C NMR
spectroscopy and EI-mass spectrometry, with the data fully consistent with that
previously reported.

\begin{scheme}
\centering
\includegraphics[width=\textwidth]{scheme1.png}
\caption{The preparation of 1 and 2}
\end{scheme}

Compound 2 reacted smoothly with Pt(PPh\(_3\))\(_4\) [50] in refluxing toluene to give \textit{trans}-
PtI(C\(_6\)H\(_4\)NAr\(_2\))(PPh\(_3\))\(_2\) (3) in excellent (95\%) isolated yield (Scheme 2). The trans
geometry of 3 was established by the observation of a singlet in the \(^{31}\)P NMR
spectrum at \(\delta 20.67\) ppm, with Pt satellites (\(J_{Pt-P} = 3074\) Hz). The complex was further
characterised by \(^1\)H and \(^{13}\)C NMR spectroscopy, which were fully assigned on the
basis of NOESY, COSY, HSQC and HMBC methods. MALDI-MS and elemental
analytical results were also fully consistent with the proposed structure.
Scheme 2. The preparation of 3, 4a, 4b, 5 and 6. (a) CuI (cat) / NHEt2 / HC≡CC6H4R.
(b) CuI (cat) / NHEt2 / HC≡CC6H4C≡CH. (c) CuI (cat) / NHEt2 / N(C6H4C≡CH-4).

Complexes of general form *trans*-PtX(Ar')(PR3)2 are known to undergo CuI-catalysed
dehydrohalogenation reactions with terminal alkynes in the presence of amine
solvents (X = Cl[2, 3, 8, 21-24, 26, 41, 42, 51, 52]; I[11]) In keeping with these
general observations, reaction of 3 with 4-ethynyl anisole or 4-
ethynylenephenylenedi-p-anisylamine (prepared from 2 by Sonogashira cross-
coupling with trimethylsilylacetylene and subsequent desilylation)[53] in NHEt2 and
catalytic (ca. 15%) CuI gave the alkynyl complexes *trans*
Pt(C≡CAr)(C6H4NAr2)(PPh3)2 (4a, 87%) and *trans*
Pt(C≡C6H4NAr2)(C6H4NAr2)(PPh3)2 (4b, 63%) as precipitates of analytical purity
(Scheme 2). The acetylide ligand gave rise to a strong ν(C≡C) band at 2107 (4a) or
2105 (4b) cm⁻¹, with whilst the *trans* geometry was again confirmed by the
observation of a singlet in the $^{31}$P NMR spectrum in each case ($4a$ 20.84 ($J_{Pt-P}$ 2985 Hz); $4b$ 20.95 ($J_{Pt-P}$ 2981 Hz)). The alkyne carbons were not observed in the $^{13}$C NMR spectra, with the weak signals from these quaternary carbons presumably being further reduced by coupling to both $^{31}$P and $^{195}$Pt. The chemically distinct OMe groups on both aryl and arylacetylide ligands were, however, readily observed in both $^1$H and $^{13}$C NMR spectra, and further confirm the presence of the alkyne ligand.

In an entirely analogous manner, the CuI-catalysed reaction of two equivalents of $3$ with 1,4-diethynyl benzene gave the bimetallic complex $\{trans-$Pt(C$_6$H$_4$NAr$_2$)(PPh$_3$)$_2\}$$_2$(µ-C≡C-1,4-C$_6$H$_4$C≡C) (5, 54%) whilst 3:1 stoichiometric reaction of $3$ with redox-active tris(4-ethynylphenyl)amine gave trimetallic N$\{C$_6$H$_4$C≡CPt(C$_6$H$_4$NAr$_2$)(PPh$_3$)$_2\}$$_3$ (6, 67%) (Scheme 2). The identity of these multimetallic systems was also readily confirmed by multinuclear NMR and IR spectroscopies, MALDI-MS and elemental analysis, or in the case of 5, ESI-HRMS. In addition to the observation of the [M]$^+$ (5) or [M+H]$^+$ (6) ions, the presence of the fully metallated bis- and tris-alkynyl ligands was confirmed by the observation of a single ν(C≡C) band at 2103 (5) or 2105 (6) cm$^{-1}$ and the absence of ν(≡C-H) bands in the IR spectra of these complexes. The protons of the aryl rings of the bridging ligand were observed as a singlet (5: 5.72 ppm) or doublets (6: 5.93, 6.31 ppm; $J_{HH}$ = 9 Hz) whilst the other spectroscopic features were similar to those of the monometallic complexes $4a$ and $4b$.

*Molecular structures* Pale yellow crystals of $4a$ (Figure 1) and $4b$ (Figure 2) that were suitable for study by X-ray diffraction were grown by slow diffusion of EtOH in to a
CH₂Cl₂ solution of the compound. Selected bond lengths (Å), bond and torsion angles (°) are given in Table 1.

Figure 1. A plot of a molecule of 4a showing the atom labelling scheme, with thermal ellipsoids plotted at 50%. Hydrogen atoms have been omitted for clarity.
Figure 2. A plot of a molecule of 4b showing the atom labelling scheme, with thermal ellipsoids plotted at 50%. The disorder in the C91 ring is shown (see Experimental), and hydrogen atoms have been omitted for clarity.

The molecular structures of 4a and 4b confirm the anticipated connectivity and trans-geometry of the complexes. The Pt centre displays the expected square-planar arrangement whilst the triarylamine fragments are arranged in the usual propeller-like geometry. [54-56] The greatest significant differences in the two structures arise in the alkynyl fragment with some evidence for a degree of cumulenic character in the N2-C26…C22-C21 portion of 4b, although the relatively low precision of the later structure prohibits detailed analysis. The Pt1-C21 bond in 4a is slightly shorter than in 4b, perhaps reflecting a greater degree of electrostatic attraction between Pt1-C1 brought about by the electron donating OMe group at C26.
Table 1. Selected bond lengths (Å), bond and torsion angles (°) for 4a and 4b

<table>
<thead>
<tr>
<th></th>
<th>4a</th>
<th>4b</th>
<th>4a</th>
<th>4b</th>
</tr>
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<tbody>
<tr>
<td>Pt1-P1</td>
<td>2.2944(7)</td>
<td>2.286(3)</td>
<td>C1-Pt1-C21</td>
<td>178.66(10)</td>
</tr>
<tr>
<td>Pt1-P2</td>
<td>2.2892(7)</td>
<td>2.313(3)</td>
<td>P1-Pt1-P2</td>
<td>169.73(3)</td>
</tr>
<tr>
<td>Pt1-C1</td>
<td>2.057(3)</td>
<td>2.144(5)</td>
<td>C1-Pt1-P1</td>
<td>94.14(7)</td>
</tr>
<tr>
<td>Pt1-C21</td>
<td>2.072(3)</td>
<td>1.993(10)</td>
<td>C1-Pt1-P2</td>
<td>93.44(7)</td>
</tr>
<tr>
<td>N1-C4</td>
<td>1.438(4)</td>
<td>1.464(9)</td>
<td>P1-Pt1-C21</td>
<td>87.20(7)</td>
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<tr>
<td>N1-C7</td>
<td>1.415(4)</td>
<td>1.431(10)</td>
<td>P2-Pt1-C21</td>
<td>85.22(7)</td>
</tr>
<tr>
<td>N1-C14</td>
<td>1.438(4)</td>
<td>1.418(9)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>C21-C22</td>
<td>1.138(4)</td>
<td>1.236(12)</td>
<td>C2-C1-C23-C24</td>
<td>41.49</td>
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<tr>
<td>C22-C23</td>
<td>1.479(4)</td>
<td>1.435(10)</td>
<td>C3-C4-C7-C8</td>
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</tr>
<tr>
<td>N2-C26</td>
<td>1.457(8)</td>
<td></td>
<td>C3-C4-C14-C15</td>
<td>42.10</td>
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<tr>
<td>N2-C29</td>
<td>1.460(8)</td>
<td></td>
<td>C25-C26-C29-C30</td>
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<tr>
<td>N2-C36</td>
<td>1.414(9)</td>
<td></td>
<td>C25-C26-C36-C37</td>
<td>121.26</td>
</tr>
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</table>

Electrochemical characterisation The triarylamine moiety is a well-known organic electrophore,[54-56] [57] which features prominently in the design of organic hole-transporting materials [58]and mixed-valence compounds.[44] [46, 59-61] The presence of one or more triarylamine groups in compounds 4 - 6 prompted consideration of their electrochemical response. Cyclic voltammograms were recorded in CH₂Cl₂ / 0.1M NBu₄[PF₆] solutions at Pt microdot working electrodes, and referenced against an internal decamethylferrocene / decamethylferrocenium couple to ferrocene [62] (Figure 3). In each case peak currents displayed a linear relationship with ν¹/² (for scan rates ν = 100, 200, 400, 800 mV s⁻¹), with peak current
ratios $0.98 < i_{pa} / i_{pc} < 1.0$. Taking 4a as a model system, the presence of the $\sigma$-bonded Pt moiety results in less positive oxidation potential for the triaryl amine moiety than in simpler organic derivatives [56]. The forward and reverse waves display a separation $\Delta E_p(1)$ of 87 mV which is comparable to that of the internal FeCp$^\ast_2 / [\text{FeCp}^\ast_2]^+$ reference couple (84 mV) (Table 2). The introduction of a second, ethynyl substituted triarylamine in 4b causes a small (+40 mV) shift in the half-wave potential of the metallated triarylamine (Table 2), with a second reversible wave arising from the ethynyl triarylamine at 189 mV. In the case of the bis(triarylamine) derivative 5, only a single redox wave could be observed for the two equivalent triarylamine moieties, with the characteristics of a one-electron process pointing to the complete independence of these moieties through the Pt-$\equiv$CC$_6$H$_4$C≡C-Pt chain. In keeping with this proposition, the voltametric wave shape was unchanged in the very weakly ion-pairing electrolyte NBu$_4$[B{C$_6$H$_3$-3,5-(CF$_3$)$_2$}]$_4$. [63-66] The electrochemical response of the branched molecule 6 was consistent with the points noted for the smaller analogues, with an initial oxidation event at +38 mV ($\Delta E_p(1) = 80$ mV) being followed by a second process of 1/3rd the peak current at +204 mV ($\Delta E_p(2) = 76$ mV). These processes can clearly be associate with the near simultaneous oxidation of the three external, metallated triarylamine moieties, and the internal tris(ethynyl)triarylamine core.
Figure 3. Plots of the cyclic voltammograms of 4a, 4b, 5 and 6 in CH$_2$Cl$_2$ with 0.1 M NBu$_4$[PF$_6$] at a scan rate of 100 mV/s and referenced against [FeCp*$_2$/FeCp*$_2$]$^+$ at -0.48 V vs FeCp$_2$ / [FeCp$_2$]$^+$ = +0.0 V.

Table 2: Oxidation potentials for platinum complexes 4 - 6 in CH$_2$Cl$_2$ with 0.1 M NBu$_4$[PF$_6$] at a scan rate of 100 mV/s and referenced against [FeCp*$_2$/FeCp*$_2$]$^+$ at -0.48 V vs FeCp$_2$ / [FeCp$_2$]$^+$ = +0.0 V.

| Compound | $E_{1/2}$(1)/mV | $\Delta E_p$(1) | $E_{1/2}$(2)/mV | $\Delta E_p$(2) | $|E_2-E_1|$/mV |
|----------|----------------|----------------|----------------|----------------|----------------|
| 4a       | 4              | 87             |                |                |                |
| 4b       | 44             | 84             | 189            | 83             | 145            |
| 5        | 12             | 93             |                |                |                |
| 6        | 38             | 80             | 204            | 76             | 166            |
Spectroelectrochemical investigations To further explore the redox products derived from 4 - 6 a series of IR (Figure 4, Table 3) and UV-vis-NIR (Figure 5, Figure 6) spectroelectrochemical experiments were conducted. IR spectroelectrochemical investigations carried out with ca. 1 mM solutions in 0.1 M NBu₄[PF₆] / CH₂Cl₂ show that the neutral complexes 4a, 4b, 5 and 6 all give rise to single ν(C≡C) band near 2100 cm⁻¹ (Table 3), which provides a convenient spectroscopic marker for redox induced structural changes in this series of compounds.[67] Oxidation of the reference compound 4a to [4a]⁺ results only a decrease in the intensity of the ν(C≡C) band, with no discernable change in frequency. An entirely analogous decrease in intensity with the most modest of shifts to higher wavenumber accompanies oxidation of 5 to [5]²⁺, which is consistent with the conclusions drawn from the electrochemical analysis concerning the independence of the amine sites.

Figure 4: IR spectra of (a) [4a]ⁿ⁺, (b) [4b]ⁿ⁺, (c) [5]ⁿ⁺ and (d) [6]ⁿ⁺ showing the ν(C≡C) band in the various oxidation states generated by in situ electrochemical oxidation in CH₂Cl₂ / 0.1 M NBu₄PF₆ in an OTTLE cell
In contrast, oxidation of 4b to [4b]$^+$ results in both a small decrease in intensity of the initial $\nu$(C=\textit{C}) band, together with the appearance of a broad, poorly structured $\nu$(\textit{C}C) band envelope between 2071 - 2027 cm$^{-1}$. Further oxidation to [4b]$^{2+}$ results in a substantial increase in the intensity of this lower energy band envelope. These IR features are accompanied by the appearance of a weak NIR band unique to [4b]$^+$ with an apparent band centre at 8000 cm$^{-1}$, which collapses on oxidation to [4b]$^{2+}$ (Figure 4) and therefore suggestive of \{Ar$_2$NC$_6$H$_4$C≡\textit{C}\} $\rightarrow$ \{PtC$_6$H$_4$N$^+$Ar$_2$\} IVCT character and hence description of [4b]$^+$ as a weakly coupled mixed valence complex (rudimentary analysis of the NIR band using the Hush expressions and taking the crystallographically determined N1-N2 distance as a proxy for the electron-transfer distance gives H$_{ab} = 170$ cm$^{-1}$).

Figure 5. An expansion of the NIR region of [4b]$^+$ on oxidation to [4b]$^{2+}$ showing the collapse of the IVCT band
Similarly, initial three-electron oxidation of 6 to [6]$^{3+}$ gives the same pattern of behaviour, with oxidation accompanied by a decrease in intensity of the $\nu$(C≡C) band and a subtle shift to higher wavenumbers, but now with a weaker $\nu$(CC) band envelope at lower wavenumbers, together with a NIR band ($\nu_{\text{max}} = 8700$ cm$^{-1}$) of now appreciable intensity. However, further oxidation to [6]$^{4+}$ results in the NIR region being masked by the tail of a higher energy optical band. Nevertheless, assuming the NIR feature in [6]$^{3+}$ is due to an IVCT-like transition, a $H_{\text{ab}}$ value of ca. 130 cm$^{-1}$ (allowing for the three possible transitions) is obtained. The $\nu$(CC) IR band envelope of [6]$^{4+}$ is similar to that of the linear model [4b]$^{2+}$. The observation of multiple $\nu$(CC) features in these highly charged systems could be a result of Fermi coupling, as often observed by Lapinte in studies of open-shell iron complexes,[68] or may arise from contributions to the spectral profile from multiple conformers.[69, 70]

Table 3: Summary of IR spectra from compounds [4 - 6]$^{n+}$ in their electrochemically accessible redox states.

<table>
<thead>
<tr>
<th>Compound</th>
<th>n = 0</th>
<th>n = 1</th>
<th>n = 2</th>
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<tbody>
<tr>
<td>[4a]$^{n+}$</td>
<td>2107(s)</td>
<td>2107(m)</td>
<td></td>
</tr>
<tr>
<td>[4b]$^{n+}$</td>
<td>2105(s)</td>
<td>2105(m)</td>
<td>2071(vs)</td>
</tr>
<tr>
<td></td>
<td>2071(s)</td>
<td>2052(sh)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>2052(sh)</td>
<td>2027(sh)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>2027(sh)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>[5]$^{n+}$</td>
<td>2103(s)</td>
<td></td>
<td>2105(m)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>n = 0</td>
<td>n = 3</td>
<td>n = 4</td>
</tr>
<tr>
<td>[6]$^{n+}$</td>
<td>2105(s)</td>
<td>2107(m)</td>
<td>2068(s)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>2027(sh)</td>
</tr>
</tbody>
</table>
To complement the IR spectra and to further investigate the electronic structures of these complexes, UV-vis-NIR spectroelectrochemical studies of 4a, 4b, 5 and 6 were undertaken from ca. 1 mM solutions in 0.1 M NBu₄PF₆ / CH₂Cl₂ (Figure 6). The complexes 4 – 6 all feature similar electronic absorption spectra with bands near 33000 cm⁻¹ and 25000 cm⁻¹, arising from the two N→π* transitions commonly observed in triarylamine complexes of general form NArAr’ (Figure 6). [54-57]. As a result of the first oxidation events, in each case this band envelope is shifted somewhat to lower energy, with a new band envelope between 20000 - 10000 cm⁻¹ characteristic of an {NArAr’}⁺ fragment and arising from π→N⁺ transitions, possibly admixed with some Pt→N⁺ character in these cases [46], in addition to the low energy IVCT bands noted earlier for [4b]⁺ and [6]³⁺. Further oxidation of the complexes [4b]⁺ to [4b]²⁺, and [6]³⁺ to [6]⁴⁺ resulted in a collapse of the IVCT feature (more clearly evident in the case of [4b]⁺→[4b]²⁺) and the appearance of a second set of π→N⁺ / Pt→N⁺ bands at lower energy.
Figure 6: UV-vis-NIR spectra of (a) \([4a] \rightarrow [4a]^+\), (b) \([4b] \rightarrow [4b]^+\), (c) \([4b]^+ \rightarrow [4b]^{2+}\), (d) \([5] \rightarrow [5]^{2+}\), (e) \([6] \rightarrow [6]^{3+}\) and (f) \([6]^{3+} \rightarrow [6]^{4+}\) showing the changes in the various oxidation states generated by \textit{in situ} electrochemical oxidation in CH\(_2\)Cl\(_2\) / 0.1 M NBu\(_4\)PF\(_6\) in an OTTLE cell.
Quantum chemical calculations. To complete the description of these redox-active Pt-ethynyl/triarylamine assemblies, we turned to quantum-chemical calculations using the BLYP35 functional and COSMO (dichloromethane) solvent model. The combination of a high amount of direct exchange in the functional and the inclusion of a solvent model has been shown to allow the accurate description of charge localization/delocalization now in a wide range of organic and organometallic mixed-valence complexes spanning the weakly to strongly coupled regime.

As expected, complexes [4a]+ and [4b]+ exhibit spin densities localized at one triarylamine unit in their monocationic forms, and in the case of [4b]+ this localization takes place at the triarylamine moiety which is directly connected to the platinum (Figure 7). This amine-localised redox behaviour is in agreement with previous studies of platinum-bridged mixed-valent triarylamine systems. Vibrational analysis within a harmonic framework provides excellent agreement between the computed ν(C≡C) frequency at 2075 cm⁻¹ and the experimental at 2071 cm⁻¹ for [4b]+ after scaling by an empirical factor of 0.95. In contrast for [4a]+ no intensity is calculated for ν(C≡C), as the ethynyl-unit is not involved in the redox event, consistent with the decrease in intensity of this band observed experimentally.
To characterise the UV-vis-NIR transitions we turned to TDDFT calculations employing the same BLYP35/COSMO(CH$_2$Cl$_2$) combination as for the ground-state analysis. We note in passing that despite the elaborately discussed failure of TDDFT in describing charge transfer excitations, this quantum-chemical approach has proven to yield reliable results for Class II systems.[71] All complexes exhibit localized charge distributions in the ground-state, and the reasonable agreement between experimental bands and calculated excitations is consistent with previous observations at this level for Class II systems. A recently published study has extensively discussed the general electronic and spectroscopic features of platinum alkynyl complexes featuring triarylamine redox centres.[46] We therefore refrain from a detailed discussion of [4a]$^+$ and [4b]$^+$. We note simply that these two complexes exhibit a typical $\beta$-HOMO - $\beta$-SOMO transition at 9996 cm$^{-1}$ ([4a]$^+$, exp. ca. 10,000 cm$^{-1}$), and at 6514 cm$^{-1}$ ([4b]$^+$, exp. ca. 7000 cm$^{-1}$), respectively. These transitions arise from
charge transfer between the alkynyl ligand and its substituent to the oxidised triarylamine moiety directly coordinated to Pt. At higher energies characteristic excitations associated with the oxidised triarylamine unit and MLCT transitions are computed (see SI). Interestingly the ethynyl unit contributes more to the orbitals involved in the excitations for $[4a]^+$ than $[4b]^+$ (see SI).

Although $[5]^+$ was not sufficiently thermodynamically stable to be observed in the spectroelectrochemical experiments, some information concerning this unusual mixed-valence complex can be obtained from quantum chemical calculations using a truncated model $\{\text{trans-Pt}(C_6H_4NAr_2)(\text{PMe}_3)_2\}_2(\mu-\text{C≡C-1,4-C}_6\text{H}_4\text{C≡C}) ([5-\text{Me}]^+)$. The compound $[5-\text{Me}]^+$ is calculated to exhibit a $N\rightarrow N^+$ IVCT transition at 6692 cm$^{-1}$ with only modest intensity ($\mu_{\text{trans}} = 0.8$ D) as expected for a weakly coupled system. The $\beta$-HOMO–1 - $\beta$-SOMO charge transfer excitation at 9808 cm$^{-1}$ ($\mu_{\text{trans}} = 3.8$ D) from the oxidized triarylamine moiety to the diethynylbenzene bridge is responsible for the next lowest-energy excitation (Figure 8). The two most intense transitions are computed at 14966 cm$^{-1}$ ($\mu_{\text{trans}} = 7.4$ D) and 16258 cm$^{-1}$ ($\mu_{\text{trans}} = 5.5$ D) as is commonly observed for oxidised triarylamine compounds. The first corresponds to a $\pi\rightarrow N^+$ CT excitation with one triarylamine unit (43%), the neighbouring platinum (20%) and the diethynylbenzene (19%) contributing significantly to the $\pi$-type orbital. The second excitation is best described in terms of an IC excitation at the charged triarylamine.
Figure 8. Isosurface plots (± 0.03 a.u.) of the β-SOMO (top) and β-HOMO–1 (bottom) of [5-Me]⁺. Hydrogen atoms have been omitted for clarity.

For [5-Me]²⁺, which was observed in the spectroelectrochemical experiments (Figure 5d), the calculations provide almost degenerate broken-symmetry (BS, “open-shell singlet”) and triplet state, with the former being slightly lower with spin desnisty distribution illustrated in Figure 9. Given that hybrid functionals with increased exact exchange admixture tend to favor high-spin states, the negligible energy difference of only 0.04 kJ/mol (calculated using the Yamaguchi spin projection procedure) in combination with the onset of spin-contamination ($<S^2>_{\text{singlet}} = 1.06$) indicate that
both states likely contribute significantly to the spectra. But as expected, the two spin states exhibit very similar spectral features both in the ground- (e.g. \( \nu(C=C)_{\text{singlet}} = 2124 \text{ cm}^{-1} \) and \( \nu(C=C)_{\text{triplet}} = 2124 \text{ cm}^{-1} \)) and the excited state. TDDFT gives very similar transition energies for the BS (e.g. 11057 cm\(^{-1}\), \( \mu_{\text{trans}} = 5.9 \text{ D} \)) and triplet state (e.g. 11073 cm\(^{-1}\), \( \mu_{\text{trans}} = 6.1 \text{ D} \)), which likely correspond to the low energy shoulder observed near 10,000 cm\(^{-1}\) (Figure 6d). But while excitations arise mainly from and to \( \alpha \)-orbitals for the BS state, transitions involve almost exclusively \( \beta \)-orbitals for the triplet. The lowest-energy transition arises in both cases from orbitals located at the diethynylbenzene unit. But while the charge is transferred upon excitation from one triarylamine in the case of the BS state, both triarylamine moieties contribute significantly (30% and 41%) to the orbital involved in the transition for the triplet.

![Figure 9](image)

**Figure 9.** Spin density isosurface plot (\( \pm 0.002 \text{ a.u.} \)) of the (broken-symmetry) low-spin state of [5-Me]\(^{2+}\). Hydrogen atoms have been omitted for clarity.

With this background understanding of the spectroscopic features of the smaller model complexes [4a]\(^{+}\), [4b]\(^{+}\) and the calculated results from [5-Me]\(^{+}\) and [5-Me]\(^{2+}\) it
is possible to readily assign the spectroscopic features in the larger systems $[6]^{3+}$ and $[6]^{4+}$. The observation of the characteristic intense triarylamine bands between 10,000 – 20,000 cm$^{-1}$ together with the current ratios described in the electrochemistry section are consistent with the oxidation of the peripheral amine moieties in $[6]^{3+}$, with the less intense NIR feature at 5500 cm$^{-1}$ arising from IVCT like processes from the inner to outer amines, all of which are similar to the analogous transitions in $[4b]^{+}$. Further oxidation of the central amine moiety in $[6]^{4+}$ gives rise to a second set of amine based transitions similar to those observed in $[4b]^{2+}$.

**Conclusion**

The synthesis of mono-ethynyl platinum species with redox active triarylamine ligands, $4a$, $4b$, $5$ and $6$, has successfully been achieved through the oxidative addition of the triarylamino iodide with the platinum precursor Pt(PPh$_3$)$_4$ and then CuI-catalysed coupling to the respective 1-alkyne. These compounds have been studied electrochemically and spectroelectrochemically, revealing weak interactions between the amine electrophores when bridged by the {-Pt(C≡C–)(PPh$_3$)$_2$} moiety.

**Experimental**

*Synthesis*

All reactions were carried out under an atmosphere of nitrogen using standard Schlenk techniques as a matter of routine, although no special precautions were taken to exclude air or moisture during work-up. Reaction solvents were purified and dried using an Innovative Technology SPS-400, and degassed before use, but no special
precautions were taken during work up, isolation or crystallisation. The compounds 4-ethynylenephenylene-diphenylamine,[53] Pt(PPh₃)₄ [50] and Pd(PPh₃)₄ [72] were prepared by the literature methods.

NMR spectra were recorded on a Bruker Avance (¹H 400.13 MHz, ¹³C 100.61 MHz, ³¹P 161.98 MHz) spectrometer from CDCl₃ solutions unless otherwise indicated, and referenced against solvent resonances (CDCl₃ ¹H 7.26 ¹³C 77.0). IR spectra (CH₂Cl₂) were recorded using a Nicolet 6700 spectrometer from cells fitted with CaF₂ windows. Electrospray ionisation mass spectra were recorded using Thermo Quest Finnigan Trace MS-Trace GC or WATERS Micromass LCT spectrometers. Samples in dichloromethane (1 mg/mL) were 100 times diluted in either methanol or acetonitrile, and analysed with source and desolvation temperatures of 120 °C, with cone voltage of 30 V. ASAP mass spectra were recorded from solid aliquots on LCT Premier XE mass spectrometer (Waters Ltd, UK) or Xevo QToF mass spectrometer (Waters Ltd, UK) in which the aliquot is vaporized using hot N₂, ionized by a corona discharge and carried to the TOF detector (working range 100-1000 m/z).

**Synthesis of N(C₆H₅)(C₆H₄OMe-4)₂ (1) [47]**

![Diagram of N(C₆H₅)(C₆H₄OMe-4)₂ (1)](image-url)
An oven dried Schlenk flask was charged with o-xylene (30 mL) and the solvent degassed. To this solution, 4-iodoanisole (15.00 g, 64.1 mmol), CuCl (0.201 g, 2.03 mmol), 1,10-phenanthroline (0.274 g, 1.5 mmol) and aniline (2.78 mL, 30.5 mmol) were added, the mixture was heated at reflux for 35 minutes, KOH (13.70 g, 244 mmol) was added and the mixture heated at reflux for 28 hours, cooled, poured in to H2O (150 mL) and extracted with CH2Cl2 (3 x 60 mL). The organic layers were combined and dried over MgSO4, filtered and the solvent removed in vacuo. The residue was suspended in hexane and purified by silica column chromatography eluting with hexane increasing to hexane:CH2Cl2 (50:50), removal of the solvent and crystallisation from hot hexane gives the title compound as a white solid. Yield 3.29 g, 32 %. 1H NMR (400 MHz, Chloroform-d) δ 7.16 (d, J = 7 Hz, 2H), 7.05 (d, J = 9 Hz, 4H), 6.94 (d, J = 7 Hz, 2H), 6.90 - 6.85 (m, 1H), 6.82 (d, J = 9 Hz, 4H), 3.80 (s, 6H).

Literature: [47] 1H NMR (CDCl3): δ = 7.25–6.79 (m, 13H, Ar), 3.79 ppm (s, 6H, OCH3). ESI-MS: 305.6 [M]+

**Synthesis of N(C6H4I-4)Ar2 (2) [73]**
An oven dried Schlenk flask was charged with chloroform (90 mL) and the solvent degassed. To this solution, 1 (3.00 g, 9.8 mmol) and NIS (2.430 g, 10.8 mmol) were added with the exclusion of light followed by acetic acid (60 mL) and the mixture stirred at room temperature for 30 hours and quenched with aqueous sodium thiosulphate (300 mg in 30 mL H$_2$O) and extracted with CH$_2$Cl$_2$ (3 x 30 mL). The organic layers were combined and dried over MgSO$_4$, filtered and the solvent removed in vacuo to give an off-white solid. Yield 3.14 g, 75 %). $^1$H NMR (400 MHz, CDCl$_3$) δ 7.40 (d, $J$ = 8 Hz, 2H), 7.02 (d, $J$ = 8 Hz, 4H), 6.82 (d, $J$ = 8 Hz, 4H), 6.67 (d, $J$ = Hz, 2H), 3.79 (s, 6H). Literature: $^1$H NMR (250 MHz, CDCl$_3$): δ 7.40/6.67 (m, AA’, 2 H/m, BB’, 2H; I-C$_6$H$_4$), 7.03/6.82 (m, AA’+BB’, 8H; MeO-C$_6$H$_4$), 3.79 (s, 6H; MeO) ESI-MS: 431.5 [M]$^+$

**Synthesis of trans-PtI(C$_6$H$_4$NAr$_2$)(PPh$_3$)$_2$ (3)**

An oven dried Schlenk flask was charged with dry toluene (20 cm$^3$) and the solvent degassed. To this solution, Pt(PPh$_3$)$_4$ (1.00 g, 0.804 mmol) and 4-iodophenylene-$p$-dianisylamine (0.693 g, 1.60 mmol) were added and the mixture stirred for 16 hours at reflux. The mixture was cooled to ambient temperature and added to vigorously
stirred hexane (150 cm³) and the off-white precipitate was filtered and washed with hexane (2 x 10 mL). Yield 880 mg, 95 %. ¹H NMR (CD₂Cl₂) δ 3.76 (s, 6H, Ha), 5.98 (d, J = 8Hz, 2H, Hh), 6.71 (m, 10H, Hc,d and g) 7.34 (vt, J = 8Hz, 12H, Hk), 7.41 (t, J = 8Hz, 6H, Hm), 7.62 (m, 12H, Hl). ³¹P NMR (CD₂Cl₂): 20.67 (JPt-P = 3074Hz). ¹³C NMR (CD₂Cl₂): 154.72 (Cb, s), 142.22 (Ce, s), 141.78 (Cf, s), 136.17 (Cg, s), 135.08 (Cl, t, Jc-P = 5Hz), 131.80 (Cj, t, Jc-P = 28Hz), 129.88 (Cm, s), 127.58 (Ck, t, Jc-P = 5Hz), 125.19 (Cc, s), 122.99 (Ch, s), 113.99 (Cd, s), 55.39 (Ca, s). MALDI-MS(+) m/z: 1151.1 [M+H]⁺. Analysis found (calculated) %: C 58.60 (58.44), H 4.04 (4.20), N 1.29 (1.22).

**Synthesis of trans-Pt(C≡CAr)(C₆H₄NAr₂)(PPh₃)₂ (4a)**

![Chemical Structure](image)

An oven dried Schlenk flask was charged with dry HNEt₂ (10 cm³) and the solvent degassed. To this solution, 4-ethynylanisole (0.017 g, 0.13 mmol), 3 (150 mg, 0.13 mmol) and CuI (4 mg) were added and the solution stirred at room temperature for 17 hours. The precipitate was filtered, washed with ethanol (3 x 5 cm³) and methanol (3 x 5 cm³) and dried under airflow for 1 h. Yield 131 mg, 87 %. ¹H NMR (CD₂Cl₂): δ 3.64 (s, 3H, Ht), 3.76 (s, 6H, Ha), 6.11 (d, J = 8Hz, 2H, Hh), 6.15 (d, J = 9Hz, 6H, Hr), 6.43 (d, J = 9Hz, 2H, Hq), 6.71 (d, J = 8Hz, 2H, Hg), 6.73 (m, 8H, Hc and d) 7.32 (vt, J = 8Hz, 12H, Hk), 7.40 (t, J = 8Hz, 6H, Hm), 7.64 (m, 12H, Hj). ³¹P NMR
(CD₂Cl₂): 20.84 (Jₚ-ₚ = 2985Hz). ¹³C NMR (CD₂Cl₂): 156.77 (Cs, s), 154.29 (Cb, s), 142.31 (Ce, s), 141.55 (Cf, s), 139.55 (Cg, s), 134.84 (Cl, t, Jₗₚ₋ₚ = 5Hz), 131.79 (Cj, t, Jₗₚ₋ₚ = 28 Hz), 131.35 (Cq, s), 129.75 (Cm, s), 127.58 (Ck, t, Jₗₚ₋ₚ = 5Hz), 124.10 (Cc, s), 123.94 (Ch, s), 121.74 (Cp, s), 113.92 (Cd, s), 112.71 (Cr, s), 55.40 (Ca, s), 54.96 (Ct, s). MALDI-MS(+) m/z: 1155.3 [M+H]+. IR (CH₂Cl₂) ν(C≡C) 2107 cm⁻¹. Analysis found (calculated) %: C 67.45 (67.58), H 4.74 (4.80), N 1.17 (1.21).

**Synthesis of trans-Pt(C≡C₆H₄NAr₂)(C₆H₄NAr₂)(PPh₃)₂ (4b)**

An oven dried Schlenk flask was charged with 4-ethynylephenylene-ₚ-anisylamine (0.043 g, 0.13 mmol) in dry CH₂Cl₂ and the solvent removed in vacuo. Dry HNET₂ (10 cm³) was added and the solvent degassed. To this solution, PtI(C₆H₄N(C₆H₄OCH₃-4)₂)(PPh₃)₂ (150 mg, 0.130 mmol) and CuI (4 mg) were added and the solution stirred at room temperature for 2 hours. The precipitate was filtered, washed with methanol (3 x 5 cm³), ethanol (3 x 5 cm³) and hexane (3 x 5 cm³) and dried under airflow for 1 h. Yield 109 mg, 63 %. ¹H NMR (CD₂Cl₂): δ 3.76 (s, 12H, Ha and x), 6.03 (d, J = 8Hz, 2H, Hq), 6.11 (d, J = 8Hz, 2H, Hh), 6.44 (d, J = 8Hz, 2H, Hr), 6.68 (d, J = 8Hz, 2H, Hg), 6.73 (m, 8H, Hc and d), 6.76 (d, J = 8Hz, 4H, Hv),
6.89 (d, J = 8Hz, 4H, Hu), 7.32 (vt, J = 8Hz, 12H, Hk), 7.39 (t, J = 8Hz, 6H, Hm),
7.64 (m, 12H, Hf). $^{31}$P NMR (CD$_2$Cl$_2$): 20.95 (J$_{Pt-P}$ = 2981Hz). $^{13}$C NMR (CD$_2$Cl$_2$):
155.48 (Cw, s), 154.29 (Cb, s), 145.33 (Cr, s), 142.31 (Ce, s), 141.59 (Cf, s), 141.12
(Cs, s), 139.53 (Cg, s), 134.84 (Cl, t, J$_{C-P}$ = 5Hz), 131.781 (Cj, t, J$_{C-P}$ = 28 Hz), 130.97
(Cq, s), 129.76 (Cm, s), 127.59 (Ck, t, J$_{C-P}$ = 5Hz), 125.86 (Cu, s), 124.11 (Cc, s),
123.93 (Ch, s), 120.45 (Cr, s), 114.37 (Cv, s), 113.92 (Cd, s), 55.40 (Ca, s), 55.36
(Cx, s). MALDI-MS(+) m/z: 1352.3 [M+H]$^+$. IR (CH$_2$Cl$_2$) ν(C≡C) 2105 cm$^{-1}$.
Analysis found (calculated)%: C 69.05 (69.27), H 4.84 (4.92), N 2.11 (2.07).

**Synthesis of {trans-Pt(C$_6$H$_4$NAr$_2$)(PPh$_3$)$_2$}$_2$(µ-C≡C-1,4-C$_6$H$_4$C≡C) (5)**

An oven dried Schlenk flask was charged with dry HNEt$_2$ (10 cm$^3$) and the solvent
degassed. To this solution, PtI(C$_6$H$_4$N(C$_6$H$_4$OCH$_3$-4)$_2$)(PPh$_3$)$_2$ (100 mg, 0.087 mmol),
CuI (3 mg) and 1,4-diethynylbenzene (5 mg, 0.043 mmol) were added and the
solution stirred at room temperature for 16 hours. The precipitate was filtered, washed
with ethanol (3 x 5 cm$^3$), hexane (3 x 5 cm$^3$) and methanol (3 x 5 cm$^3$) and dried
under airflow for 1 h. Yield 50 mg, 54 %. $^1$H NMR (CD$_2$Cl$_2$): δ 3.74 (s, 12H, Ha),
5.72 (s, 4H, Hq), 6.08 (d, J = 8Hz, 4H, Hh), 6.67 (d, J = 8Hz, 4H, Hg), 6.71 (m, 16H,
Hc and d) 7.28 (vt, J = 8Hz, 24H, Hk), 7.37 (t, J = 8Hz, 12H, Hm), 7.59 (m, 24H, Hf).
$^{31}$P NMR (CD$_2$Cl$_2$): 20.73 (J$_{Pt-P}$ = 2971Hz). $^{13}$C NMR (CD$_2$Cl$_2$): 154.27 (Cb, s),
142.29 (Ce, s), 141.42 (Cf, s), 139.50 (Cg, s), 134.78 (Cl, t, J$_{C-P}$ = 5Hz), 131.70 (Cj, t,
$J_{C,P} = 28 \text{ Hz}$, 129.71 (C$q$, s), 129.23 (C$m$, s), 127.55 (C$k$, t, $J_{C,P} = 5 \text{Hz}$), 124.08 (C$c$, s), 123.90 (C$h$, s), 113.90 (C$d$, s), 55.39 (C$a$, s). MALDI-MS(+) $m/z$: 2170.6 $[M]^+$. IR (CH$_2$Cl$_2$) $\nu$(C≡C) 2103 cm$^{-1}$. ESI-HRMS (m/z) found 1085.8011, calculated 1085.7977

**Synthesis of N\{C$_6$H$_4$C≡CPt(C$_6$H$_4$NAr$_2$)(PPh$_3$)$_2$\}_3 (6)**

An oven dried Schlenk flask was charged with dry HNEt$_2$ (10 cm$^3$) and the solvent degassed. To this solution, PtI(C$_6$H$_4$N(C$_6$H$_4$OCH$_3$-4)$_2$)(PPh$_3$)$_2$ (150 mg, 0.130 mmol), CuI (3 mg) and triethynylphenylamine (13 mg, 0.043 mmol) were added and the solution stirred at room temperature for 16 hours. The precipitate was filtered, washed
with ethanol (3 x 5 cm³), hexane (3 x 5 cm³) and methanol (3 x 5 cm³) and dried under airflow for 1 h. Yield 98 mg, 67 %. ¹H NMR (CD₂Cl₂): δ 3.76 (s, 18H, Ha), 5.93 (d, J = 9Hz, 6H, Hr), 6.11 (d, J = 8Hz, 6H, Hh), 6.31 (d, J = 9Hz, 6H, Hq), 6.69 (d, J = 8Hz, 6H, Hg), 6.73 (m, 24H, Hc and d) 7.32 (vt, J = 8Hz, 36H, Hk), 7.39 (t, J = 8Hz, 18H, Hm), 7.62 (m, 36H, Hl). ³¹P NMR (CD₂Cl₂): 20.93 (J_P-P = 2998Hz). ¹³C NMR (CD₂Cl₂): 154.29 (Cb, s), 142.30 (Ce, s), 143.90 (Cs, s), 141.66 (Cf, s), 139.55 (Cg, s), 134.83 (Cl, t, J_C-P = 5Hz), 131.76 (Cj, t, J_C-P = 28 Hz), 130.96 (Cr, s), 129.78 (Cm, s), 127.60 (Ck, t, J_C-P = 5Hz), 124.11 (Cc, s), 123.93 (Ch, s), 122.73 (Cq, s), 113.92 (Cd, s), 55.40 (Ca, s). MALDI-MS(+) m/z: 3385.6 [M+H]+. IR (CH₂Cl₂) ν(C≡C) 2105 cm⁻¹. Analysis found (calculated) %: C 67.87 (68.10), H 4.48 (4.64), N 1.69 (1.65).

**Computational Details**

Full structure optimizations and analysis of all ground and excited state[74-76] properties were performed using a locally modified version of the TURBOMOLE 6.4 program code[77] enabling the use of the BLYP35 hybrid functional[78] based on

\[ E_{XC} = 0.65(E_{X}^{LSDA} + \Delta E_{X}^{88}) + 0.35E_{X}^{exact} + E_{C}^{LYP} \]

which has been shown to give accurate results for organic[78-82] and inorganic[83-86] MV systems. Split-valence def2-SVP basis sets were employed on all lighter atoms, together with the corresponding def2-SVP effective-core potential and a corresponding valence basis set for platinum.[87-89] Computed harmonic vibrational frequencies were scaled by an empirical factor of 0.95.[90, 91] To account for solvent
effects, the conductor-like-screening (COSMO) solvent model was employed for
ground state structure optimizations and analysis as well as in subsequent TDDFT
calculations of excitation energies and transition dipole moments.[92] Dichloromethane ($\varepsilon = 8.93$) was used, as experimental data were collected in this
solvent (non-equilibrium solvation was assumed in the TDDFT calculations). Spin-
density and molecular-orbital isosurface plots were generated with the GaussView
were replaced by PMe$_3$, was used to reduce computational cost.

**Crystallographic Details**
The X-ray single crystal data for compounds **4a** and **4b** have been collected at 120.0K
on an Agilent XCalibur diffractometer (graphite monochromator, $\lambda$MoK$\alpha$, $\lambda$
=0.71073Å) equipped with a Cryostream (Oxford Cryosystems) open-flow nitrogen
cryostat. The structures was solved by direct method and refined by full-matrix least
squares on $F^2$ for all data using Olex2 [94] and SHELXTL [95] software. All non-
disordered non-hydrogen atoms were refined anisotropically, hydrogen atoms were
placed in the calculated positions and refined in riding mode. During the preliminary
stages of refinement, the extremely elongated thermal ellipsoids of carbon atoms of
one of the Ph-rings in the structure **4b** indicated the disorder of the group. The
different parts of the disordered group differ by their orientation around the P2-C91
bond. The disordered atoms were split in two groups and the group occupancies were
allowed to refine freely while the isotropic a.d.p. of the atoms remained fixed at equal
values. Then the occupancies of the disordered groups were fixed at the refined values
and both components of the disordered group were refined as regular hexagons (AFIX 66) and with isotropic a.d.p. for all carbon atoms.

**Crystal Data for 4a:** C_{65}H_{55}NO_3P_2Pt, M = 1155.13, triclinic, space group P-\(1\), \(a = 10.2066(3)\), \(b = 14.5412(4)\), \(c = 18.8204(6)\) Å, \(\alpha = 70.667(2)\), \(\beta = 85.206(2)\), \(\gamma = 85.756(2)\)°, \(V = 2623.33(12)\) Å\(^3\), \(Z = 2\), \(\mu(\text{MoK}\alpha) = 2.784\) mm\(^{-1}\), \(D_{\text{calc}} = 1.462\) g/mm\(^3\), 41162 reflections measured, 12659 unique (\(R_{\text{int}} = 0.0472\)) were used in all calculations. The final \(R_1\) was 0.0293 (>2\(\sigma(I)\)) and \(wR_2\) was 0.0641 (all data).

**Crystal Data for 4b:** C_{78}H_{66}N_2O_4P_2Pt, M = 1352.36, monoclinic, space group P2_1/c, \(a = 22.8460(15)\), \(b = 9.9786(5)\), \(c = 31.3485(17)\) Å, \(\beta = 110.108(7)\)°, \(V = 6710.9(7)\) Å\(^3\), \(Z = 4\), \(\mu(\text{MoK}\alpha) = 2.189\) mm\(^{-1}\), \(D_{\text{calc}} = 1.338\) g/mm\(^3\), 70643 reflections measured, 15348 unique (\(R_{\text{int}} = 0.2724\)) were used in all calculations. The final \(R_1\) was 0.0882 (6961>2\(\sigma(I)\)) and \(wR_2\) was 0.1729 (all data). The structure contains a severely disordered solvent molecules (H\(_2\)O/methanol). Their contribution to the structural factors was taken into account by applying MASK procedure of OLEX2 program package.

**Appendix A Supplementary data**

Tables of calculated excited state parameters: UV-vis-NIR transition energies \(E_{\text{trans}}\), transition dipole moments \(\mu_{\text{trans}}\) and main MO contributions for [4a]\(^+\), [4b]\(^+\), [5-Me]\(^+\), 1[5-Me]\(^{2+}\), 3[5-Me]\(^{2+}\). ESI-HRMS of 5 showing the predicted and observed spectra for [M]\(^{2+}\). CCDC 989928 and 989929 contain the supplementary crystallographic data for 4a and 4b. These data can be obtained free of charge via [http://www.ccdc.cam.ac.uk/conts/retrieving.html](http://www.ccdc.cam.ac.uk/conts/retrieving.html), or from the Cambridge Crystallographic Data Centre, 12 Union Road, Cambridge CB2 1EZ, UK; fax: (+44) 1223-336-033; or e-mail: deposit@ccdc.cam.ac.uk.
References

[77] in.