Mini Review

Luminescent metal complexes derived from carbene and related ligands: tuning excited-state properties with metal–carbon multiple bonds

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Abstract

This account begins with a report on the photophysical responses to pH exhibited by the luminescent chelating bis(carbene) platinum(II) complexes [(C10H21N4)Pt(CN)(CN’Bu)] (1) and [(C10H21N4)Pt(CN’Bu)2] (2), and concludes with an overview of recent work by our group regarding luminescent platinum and rhenium derivatives bearing metal–carbon multiple bonds. © 2001 Elsevier Science B.V. All rights reserved.

Keywords: Carbene; Photoluminescence; Platinum; Protonation; Rhenium; Sensors

1. Introduction

As research into the preparation of metal carbene complexes and their roles as stoichiometric and catalytic reagents have proliferated [1], their luminescent characteristics have largely been neglected. Yet such investigations are potentially rewarding, because the population of charge-transfer excited states can transform the nature and reactivity of the metal–carbene moiety. By analogy to the rich photochemistry displayed by d2 derivatives containing metal–ligand multiple bonds [2,3], the metal–carbene interaction may be weakened or modified in the excited state to yield species capable of mediating photo-induced electron transfer or carbene group transfer reactions.

The photochemistry of the Group 6 Fischer carbene complexes [(CO)3M=C(OR)]’ (M = Cr, W) have been studied in some detail [4] and applications in organic synthesis have been developed [5]. Yam et al. have described the luminescent behavior of ruthenium(II) bipyridine complexes with orthometalated aminocarbene moieties [6]. The first observation of a metal→π*(carbene) charge transfer excited state in solution at ambient temperatures was reported by our group for the hexanuclear platinum(II) macrocycle [Pt(CN)2(C10H21N4)6] bearing chelating bis(carbene) and bridging cyanide ligands [7]. Our endeavors in developing luminescent metal–carbon multiple-bonded species have also proved fruitful for low-valent Pt(II) [8] and Re(I) [9] carbene derivatives supported by diimine auxiliaries, while the preparation and excited-state properties of high-valent Re(V) [10,11] and -(VI) [12] alkylidyne luminophores have been undertaken; an overview of this body of work is given at the end of this account. Mayr et al. [13] first noted fluid luminescence from alkylidyne complexes in 1985, but reports of further examples remain sparse [14,15].

The new study which constitutes the main focus of this account was inspired by early reports that protonation of chelating bis(carbene) platinum(II) complexes...
such as \([\text{C}_4\text{H}_9\text{N}_4\text{Pt(CNCH}_3\text{)}_2]^{+}\) occurred at the heterocyclic bis(carbene) moiety [16]. We perceived these species as potential pH sensors because the reaction was reversible upon treatment with base and the protonation was accompanied by a color change from red to yellow. In addition, their photoluminescent properties are hitherto unknown. Significantly, a number of Pt(II) complexes have been identified as prospective luminescent chemical sensors [17]. Compared to luminescent octahedral d\(^6\) metal species, square planar d\(^8\) Pt(II) derivatives can offer superior chemical sensing capabilities because the coordinatively unsaturated geometry facilitates outer- and inner-sphere interactions with substrates. We now described the synthesis and spectroscopic properties of the chelating bis(carbene) Pt(II) analogs \([\text{C}_{10}\text{H}_{21}\text{N}_4\text{Pt(CN)(CNBu)}]^{+}\) (1) and \([\text{C}_{10}\text{H}_{21}\text{N}_4\text{Pt(CNBu)}_2]^{+}\) (2), together with the crystal structure of 1. Their photophysical responses upon treatment with tetrafluoroboric acid, and in particular observations of photoluminescence enhancement, are also presented. Kaiwar et al. have recently examined the impact of protonation upon the excited-state parameters of heterocycle-substituted platinum–1,2-enedithiolate complexes [18].

2. Experimental

2.1. General procedures

K\(_2\)PtCl\(_4\) (Strem), tert-butylisocyanide (Aldrich) and hydrazine hydrate (Aldrich) were used as received. \(\text{(Caution: perchlorate salts are potentially explosive and should be handled with care and in small amounts.)}\) Solvents used for synthesis were of analytical grade and purified according to conventional methods. Fast atom bombardment (FAB) mass spectra were obtained on a Finnigan Mat 95 mass spectrometer. \(^1\)H (300 MHz) and \(^13\)C (126 MHz) spectra were recorded on DPX 300 and 500 Bruker FT-NMR spectrometers, respectively, with chemical shift (in ppm) relative to tetramethylsilane. Elemental analysis was performed by the Institute of Chemistry at the Chinese Academy of Sciences, Beijing. Infrared (IR) spectra were recorded as Nujol mulls on a BIO RAD FTIR spectrometer. UV–vis absorption spectra were recorded on a Perkin–Elmer Lambda 19 UV–vis spectrophotometer.

Steady-state emission spectra were recorded on a SPEX 1681 Fluorolog-2 series F111AI spectrophotometer. Low-temperature (77 K) emission spectra for solid-state samples were recorded in 5-mm diameter quartz tubes which were placed in a liquid nitrogen Dewar equipped with quartz windows. The emission spectra were corrected for monochromator and photomultiplier efficiency and for xenon lamp stability.

Sample and standard solutions were degassed with at least three freeze–pump–thaw cycles. The emission quantum yield was measured with \([\text{Ru(bpy)}_3]^{2+}\) in degassed acetonitrile as the standard (\(\Phi_s = 0.062\)) and calculated by: \(\Phi_s = \Phi_l(\varepsilon_s/\varepsilon_l)(n_m/n_s)(D_s/D_l)\), where the subscripts s and r refer to sample and reference standard solution, respectively, \(n\) is the refractive index of the solvents, \(D\) is the integrated intensity, and \(\Phi\) is the luminescence quantum yield. The quantity \(B = 1 - 10^{-\alpha L}\) where \(\lambda\) is the absorbance at the excitation wavelength and \(L\) is the optical path length.

Emission lifetimes and flash-photolysis measurements were performed with a Quanta Ray DCR-3 pulsed Nd:YAG laser system (pulse output 355 nm, 8 ns). The emission signals were detected by a Hamamatsu R928 photomultiplier tube and recorded on a Tektronix model 2430 digital oscilloscope. Error limits are estimated for: \(\lambda(\pm 1\text{nm})\), \(t(\pm 10\%)\), \(\phi(\pm 10\%)\).

2.2. Syntheses

2.2.1. \([\text{C}_{10}\text{H}_{21}\text{N}_4\text{Pt(CN)(CNBu)}]^{+}\) (1) and \([\text{C}_{10}\text{H}_{21}\text{N}_4\text{Pt(CNBu)}_2]^{+}\)ClO\(_4\) (2ClO\(_4\))

To a stirred solution of K\(_2\)PtCl\(_4\) (0.20 g, 0.48 mmol) in H\(_2\)O (20 ml) at room temperature was added tert-butylisocyanide, tBuN=C (0.20 g, 2.41 mmol). A colorless solution of the [Pt(CNBu)]\(^{2+}\) cation was formed. An excess of hydrazine hydrate (1 ml) was added to give a yellow solution, which was stirred at 40°C for 2 h. The resultant yellow precipitation was collected, washed with water and dried to yield complex 1 as a yellow solid: yield 0.04 g, 15 %.

To the yellow filtrate was added excess LiClO\(_4\) to yield 2(ClO\(_4\)) as a yellow solid: yield 0.20 g, 63%. Recrystallization of 1 and 2(ClO\(_4\)) by diffusion of diethyl ether into chloroform solutions yielded yellow crystals.

2.2.1.1. Compound 1. Anal. Calc. for C\(_{14}\)H\(_{30}\)N\(_6\)Pt: C, 38.32; H, 6.03; N, 16.76. Found: C, 38.22; H, 5.98; N, 16.87%. FAB-MS: \(m/z\) 502 [M\(^{+}\), 445 [M\(^{+}+\)tBu]. IR (Nujol): \(\nu = 1997, 2132\) (C=N), 1562, 1534 (C=N) cm\(^{-1}\).

**1H-NMR (CD\(_3\)CN):** 1.37 (s, 18H, tBu), 1.59 (s, 9H, tBuNC), 4.17 (broad s, 1H, NH), 5.55 (broad s, 1H, NH), 6.55 (broad s, 1H, NH). 13C\(^{1}\)H-NMR (CD\(_3\)CN): 28.1, 28.2, 29.1 (C\(_{tBu}\)), 4.17 (broad s, 1H, NH), 5.55 (broad s, 1H, NH), 6.55 (broad s, 1H, NH).

2.2.1.2. Compound 2(ClO\(_4\)). Anal. Calc. for C\(_{20}\)H\(_{38}\)N\(_6\)Cl\(_2\)O\(_4\)Pt: C, 36.50; H, 5.97; N, 12.77. Found: C, 36.45; H, 5.81; N, 12.86%. FAB-MS: \(m/z\) 558 [M\(^{+}\), 502 [M\(^{+}+\)tBu]. IR (Nujol): \(\nu = 2229, 2206\) (C=N), 1571, 1531 (C=N) cm\(^{-1}\).

**1H-NMR (CD\(_3\)CN):** 1.37 (s, 18H, tBu), 1.58 (s, 18H, tBuNC), 4.92 (broad s, 2H, NH), 8.69 (broad s, 1H, NH), 13C\(^{1}\)H-NMR (CD\(_3\)CN): 28.9, 29.9 (C\(_{Me}\)), 54.5, 61.0 (C\(_{Me}\)); remaining signals not resolved.
2.2.2. [(C10H22N4)Pt(CN)(CN‘Bu)]BF4 (3(BF4)) and (C10H22N4)Pt(CN‘Bu)2[BF4]2 (4(BF4)2)

HBF4 (50 %) was added dropwise to an acetonitrile solution (15 ml) of 1 (0.1 g, 0.20 mmol) until a colorless solution was afforded. The solvent volume was reduced and addition of diethyl ether gave a white solid. This was collected, dried, and recrystallized by slow diffusion of diethyl ether into an acetonitrile solution to yield white crystals of 3(BF4). This procedure was adopted using 2(ClO4) (0.10 g, 0.15 mmol) to afford 4(BF4)2.

2.2.2.1. Compound 3(BF4). Yield 0.09 g, 75%. Anal. Calc. for C16H31N6BF4Pt: C, 32.61; H, 5.30; N, 14.26. Found: C, 32.80; H, 5.01; N, 14.19 %. FAB-MS: m/z 503 [M+], 446 [M+–Bu]. IR (Nujol): ν = 2223, 2140 (C=N), 1567, 1529 (C≡N) cm⁻¹. 1H-NMR (CDCl3): 1.46, 1.47 (two singlets, 18H, tBu), 1.66 (s, 9H, tBuNC), 5.70 (s with 195Pt satellites, 1H, 3JpH = 26.4 Hz, tBuNH), 7.18 (s with 195Pt satellites, 1H, 3JpH = 33.5 Hz, tBuNF), 9.92, 9.99 (two singlets with overlapping 195Pt satellites, 2H, 3JpH = 67.5 and 71.5 Hz, heterocyclic NH). 13C{1H}-NMR (CD3CN): 27.1, 28.5 (CMe3); remaining signals not resolved.

2.2.2.2. Compound 4(BF4)2. Yield 0.08 g, 70%. Anal. Calc. for C20H40N6B2F8Pt: C, 32.76; H, 5.50; N, 11.46. Found: C, 32.84; H, 5.46; N, 11.28 %. FAB-MS: m/z 559 [M+], 502 [M+–Bu]. IR (Nujol): ν = 2230 (C≡N), 1597, 1571 (C≡N) cm⁻¹. 1H-NMR (CD3CN): 1.45 (s, 18H, tBuNH), 1.62 (s, 18H, tBuNC), 6.43 (broad s, 2H, 3JpH = 26.4 Hz, tBuNH), 9.61 (s with 195Pt satellites, 2H, 3JpH = 63.8 Hz, heterocyclic NH). 13C{1H}-NMR (CD3CN): 28.1, 28.9 (CMe3), 53.1, 59.4 (CMe3); remaining signals not resolved.

2.3. Procedure for examining effects of pH on photophysical properties

A solution of 2(ClO4) was prepared by dissolving the complex (7.17 mg, 0.01 mmol) in aqueous NaCl solution (0.1 M, 50 ml). The pH of the solution was adjusted by addition of aqueous HBF4 (3 × 10⁻² M). The pH value was monitored with a pH meter and the absorption spectra at various pH values were recorded.

2.4. X-ray crystallography

Yellow crystals of [(C10H22N4)Pt(CN)(CN‘Bu)] (1) were obtained by vapor diffusion of diethyl ether into a chloroform solution. Crystal data: C16H30N6Pt, M = 501.53, crystal size = 0.40 × 0.30 × 0.26 mm, orthorhombic, P212121, a = 9.392(2), b = 12.925(2), c = 17.211(2) Å, V = 2089.4(6) Å³, Z = 4, Dcalc = 1.594 g cm⁻³, μ = 6.723 mm⁻¹, F(000) = 984, T = 295 K. A total of 2712 independent reflections were collected on a Nonius CAD4 diffractometer (λ(Mo-Kα) = 0.71073 Å, θ range = 197–27.50 °). The structure was solved by direct methods and refined by full-matrix least-squares treatment on F² using SHELXTL [19]. All H atoms were generated in idealized positions and refined in riding model together with attached atoms (the atoms H(3) and H(4) each have occupation numbers of 0.5 since C(7)–N(3), C(8)–N(4) and C(7)–N(5), C(8)–N(6) are indistinguishable): R = 0.029, wR = 0.061, GoF = 1.035 for 2281 absorption-corrected (semi-empirical from ϕ-scans, transmission 0.161–0.253) reflections with I > 2σ(I) and 209 parameters (residual ρ = + 1.59, 0.66 e Å⁻³).

3. Results and discussion

3.1. Synthesis and characterization of mononuclear platinum(II) chelating bis(carbene) complexes

The synthesis of mononuclear platinum(II) complexes with a heterocyclic monoanionic planar bis(carbene) ligand by treating a concentrated aqueous solution of K2PtCl₄ with alkyl isocyanide then hydrazine has long been known [16a]. In this work, reaction of K2PtCl₄·BuNC and hydrazine in H₂O at 40°C resulted in the precipitation of [(C₁₀H₂₂N₄)Pt(CN)(CN‘Bu)] (1) in low yield (15%). Addition of excess LiClO₄ to the filtrate afforded [(C₁₀H₂₂N₄)Pt(CN‘Bu)₂]-ClO₄ (2(ClO₄)) as a yellow solid in 63 % yield. Protonation of 1 and 2(ClO₄) with aqueous HBF₄ solution gave [(C₁₀H₂₂N₄)Pt(CN)(CN‘Bu)]BF₄ (3(BF₄)) and
Complex [(C10H22N4)Pt(CN)2]34 (15 300), 282 (14 100), 298 (6700), 311 ppm. After protonation, the 1H-NMR spectrum of three very broad peaks at 4.17, 6.55 and 8.11 (NH) 1.41 (inequivalent Pt–C(2)–N(2) 174.9(8)
Pt–C(1)–N(1) C(7)–N(5)–C(9)179.2(9) 132.1(7)
C(1)–Pt–C(8) C(7)–N(3)–N(4) 96.3(3) 119.5(6)

[(C10H22N4)Pt(CN)2]227 (12 500), 270 (9500), 288 (6700), 310 ppm. After protonation, the 1H-NMR spectrum of three very broad peaks at 4.17, 6.55 and 8.11 (NH) and 1.59 (CN)

Table 1
Selected bond lengths (Å) and angles (°) for 1

<table>
<thead>
<tr>
<th>Bond Length/Angle</th>
<th>Pt–C(1)</th>
<th>Pt–C(2)</th>
<th>Pt–C(7)</th>
<th>Pt–C(8)</th>
<th>C(1)–N(1)</th>
<th>C(2)–N(2)</th>
<th>C(7)–N(3)</th>
<th>C(8)–N(4)</th>
<th>C(1)–Pt–C(2)</th>
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<td></td>
<td>2.006(8)</td>
<td>1.991(8)</td>
<td>2.007(7)</td>
<td>2.009(7)</td>
<td>1.16(1)</td>
<td>1.12(1)</td>
<td>1.12(1)</td>
<td>1.12(1)</td>
<td>113.7(5)</td>
<td>113.7(5)</td>
</tr>
<tr>
<td></td>
<td>Pt–C(7)–N(3)</td>
<td>C(7)–N(5)</td>
<td>C(8)–N(4)</td>
<td>C(7)–N(3)–N(4)</td>
<td>113.7(5)</td>
<td>113.7(5)</td>
<td>113.7(5)</td>
<td>113.7(5)</td>
<td>113.7(5)</td>
<td>113.7(5)</td>
</tr>
</tbody>
</table>

Table 2
UV–vis spectral data for 1–4 in acetonitrile at 298 K

<table>
<thead>
<tr>
<th>Complex</th>
<th>λmax (nm) (ε (dm3 mol−1 cm−1))</th>
</tr>
</thead>
<tbody>
<tr>
<td>[(C10H22N4)Pt(CN)2]</td>
<td>234 (20 300), 282 (10 100), 326 (2700), 390 (2480)</td>
</tr>
<tr>
<td>(CN(Bu)) (1)</td>
<td>229 (17 300), 271 (10 500), 292 (sh, 5400), 325 (1900)</td>
</tr>
<tr>
<td>[(C10H22N4)Pt(CN)(CN(Bu))2] (2)</td>
<td>234 (15 300), 282 (14 100), 298 (6700), 311 (5700), 330 (2700)</td>
</tr>
<tr>
<td>[(C10H21N4)Pt(CN)2]3+ (3)</td>
<td>227 (12 500), 270 (9500), 288 (6700), 310 (6800), 330 (3750)</td>
</tr>
<tr>
<td>[(C10H21N4)Pt(CN)(CN(Bu))2] (4)</td>
<td>234 (20 300), 282 (10 100), 326 (2700), 390 (2480)</td>
</tr>
</tbody>
</table>

The molecular structure of the neutral Pt(II) complex [(C10H21N4)Pt(CN)(CN(Bu))2] (1) has been determined by X-ray crystallography and is shown in Fig. 2; selected bond lengths and angles are listed in Table 1. A distorted square-planar environment around the metal core and a planar PtC2N2 metallacycle [7,16] are observed (in the refinement process, half occupancies have been assigned to two hydrogens attached to the two nitrogen atoms because they are crystallographically indistinguishable). The Pt–C(carbene) distances (2.007(7) and 1.991(8) Å for Pt–C(7) and Pt–C(8), respectively) are similar to those in [Pt(CN)(C10H21N4)]6 [7] (1.97(1)–2.01(1) Å) and other Pt(II) carbene complexes with or without heteroatom stabilization [8] (and references therein). Such Pt–C(carbene) interactions are expected to be predominantly σ in nature with minimal π character, and the observation of comparable Pt–C (cyanide) and Pt–C(isocyanide) bond lengths (2.006(8) and 1.991(8) Å, respectively) in 1 reinforces this notion. The short C(1)–N(1) and C(2)–N(2) distances (1.16(1) and 1.12(1) Å, respectively) are consistent with carbon–nitrogen triple bonds in the isocyanide and cyanide ligands.

3.2. Absorption and emission spectroscopy:
 pH-dependent behavior

The UV–vis spectrum of 2(ClO4) in CH3CN displays a moderately intense absorption band at 406 nm (εmax = 2150 dm3 mol−1 cm−1, Table 2). This band is solvatochromic in nature [e.g. 384 nm (εmax = 2000 dm3 mol−1 cm−1) in deionized water] and is tentatively assigned as a charge transfer transition with Pt→π*(carbene) MLCT character. This is consistent with the observed blue shift for the methylisocyanide-derived Pd(II) analog (345 nm in CH3CN) relative to the Pt(II) congener (410 nm) [20]. Assignment to Pt→π*(CN−)/π*(CNR) transitions is not favored, since Pt(II) cyanide and alkylisocyanide derivatives bearing π-acid ligands do not absorb strongly at λ > 300 nm. For example, the lowest energy absorption band for [Pt(dppe)(CN)]3 (dppe = 1,2-bis(diphenylphosphino)ethane) and [Pt(CN(Bu))2(CN)]2 appear at 294 and 282 nm, respectively in methanol [21]. Furthermore, the Pt→π*(CN−) transition of [Pt(CN)4]2− occurs at 280 nm [22]. The 390 nm band for 1 is similarly ascribed to a MLCT-type transition.

The photophysical responses by 2(ClO4) as a function of pH have been examined. The UV–vis spectral behavior of 2(ClO4) at pH 2 to 11 in deionized water is

Fig. 2. Perspective view of 1 (50% probability ellipsoids).
monitored, and a well-defined transformation is detected between pH 3 and 6. Outside this pH range, the spectral changes are more complex and a secondary process other than protonation/deprotonation is presumably occurring. The UV–vis spectral traces for 2(ClO₄) at pH 6.08 to 3.10 upon addition of HBF₄ are shown in Fig. 3. Upon decreasing the pH, the intensity of the absorption band at \( \lambda_{\text{max}} \) 384 nm decreases while those at \( \lambda_{\text{max}} \) 312 and 328 (sh) nm increase. Isosbestic points are evident at 286 and 356 nm and a color change from yellow to colorless is observed. These spectral changes are reversible as the 384 nm band is recovered upon addition of hydrazine. The UV–vis spectra of 3(BF₄) and 4(BF₄)₂ in acetonitrile (Table 2) both show a moderately intense shoulder at \( \lambda_{\text{max}} \) 330 nm (\( \varepsilon_{\text{max}} \sim 3 \times 10^3 \text{ dm}^3 \text{ mol}^{-1} \text{ cm}^{-1} \)). This further demonstrates that protonation of the heterocyclic bis(carbene) moiety produces a blue shift for the lowest energy absorption.

The luminescence of complexes 1 and 2(ClO₄) in solution at room temperature is extremely weak, but their solid-state emissions are more pronounced. Upon excitation at 350 nm, crystalline 1 emits at \( \lambda_{\text{max}} \) 508 and 536 nm (\( \tau = 0.27 \mu s \)) at 298 K, while a well-resolved vibronic structure (470, 502 (max), 537, 579 nm) is displayed at 77 K (Fig. 4). The observed vibrational spacing of ca. 1350 cm⁻¹ correlates with the C=N stretching of the chelating bis(carbene) moiety, and signifies its involvement in the excited state. We therefore tentatively assign these excited states as \(^3\text{MLCT} \text{Pt} \rightarrow \pi^*(\text{carbene})\) in nature.

The potential employment of 1 and 2(ClO₄) as luminescent pH sensors has been investigated. The photoluminescence of complex 1 is extremely weak in acetonitrile at 298 K (\( \lambda_{\text{max}} \sim 490 \text{ nm}, \phi < 5 \times 10^{-4} \)). However, emission enhancement (ca. 12-fold) accompanied by a blue shift to \( \lambda_{\text{max}} \) 442 nm is detected upon addition of excess HBF₄ (\( \lambda_{\text{ex}} \) 350 nm, Fig. 5). Likewise, 2(ClO₄) is weakly emissive in acetonitrile (\( \lambda_{\text{max}} \sim 520 \text{ nm}, \phi < 1 \times 10^{-3} \)), but addition of HBF₄ yields an increase in the emission intensity plus a blue shift to \( \lambda_{\text{max}} \) 461 nm. Importantly, these responses are reversible upon treatment with hydrazine. The observed high-energy emissions in acidic acetonitrile solutions of 1 and 2(ClO₄) are assigned to the triplet intraligand excited state of the heterocyclic bis(carbene) moiety.

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1. This is presumably due to solvent interactions with the coordinatively unsaturated Pt(II) center in the excited state, like for platinum diimine complexes.
2. The 298 K fluid emissions of the protonated derivatives 3(BF₄) and 4(BF₄)₂ in acetonitrile appear at \( \lambda_{\text{ex}} \) 430 and 445 nm (\( \tau \sim 0.3 \mu s \)), respectively. As suggested by a reviewer, the discrepancies between these emission maxima and those observed for the in situ protonation of 1 and 2(ClO₄) may be attributed to ion-pairing effects resulting in equilibrating processes in the latter.
ment with hydrazine. Limitations of this system with respect to pH-sensing applications are clear: secondary reactions/processes are apparently taking place under highly acidic and basic conditions, while the degree of emission enhancement is noticeable yet not dramatic. Nevertheless, the intention of this report is to introduce Fischer-type carbene complexes as potential luminescent chemosensors.

4.2. Luminescent Pt(II) diamino-carbene complexes [8]

A series of luminescent diimine-supported platinum(II) diamino-carbene complexes, namely [(CNN)Pt(C(NHR)(NHR))]<sup>+</sup> (A, HCNN = 6-phenyl-2,2'-bipyridine; for R<sup>1</sup> = tBu, 2,6-Me<sub>2</sub>C<sub>6</sub>H<sub>3</sub>; R<sup>2</sup> = Me, NH<sub>2</sub>, CH<sub>3</sub>Ph), have been synthesized by nucleophilic attack of amines at the coordinated isocyanide ligands of [(CNN)Pt(C≡NRR')]<sup>+</sup>. The binuclear bridging bis(carbene) derivative [(CNN)Pt(tBu-μ-(CNN)Pt(μ-CN))]<sup>+</sup> (B) was also prepared (Fig. 6). All compounds display structureless emissions in acetonitrile at room temperature (λ<sub>max</sub> 528–558 nm) and these are assigned to 3MLCT excited states. The isocyanide derivatives emit with substantially higher quantum yields (ca. 0.1), while low energy bands (λ<sub>max</sub> > 600 nm) assigned to π–π excimeric or metal–metal-to-ligand charge transfer (3MMLCT) excited states have been observed in concentrated glassy solutions.

4.3. Luminescent Re(I) N-heterocyclic carbone complexes [9]

Rhenium(I) N-heterocyclic carbone complexes of the type [HNCH<sub>2</sub>CH<sub>2</sub>NHC≡N(L-L)(CO)]<sup>+</sup> (C, Fig. 7) have been prepared [L-L = 4,4'-dimethoxy-2,2'-bipyridine, 4,4'-bis(t-butyl)-2,2'-bipyridine, 2,2'-bipyridine, 4,4'-dichloro-2,2'-bipyridine, 4,4'-bis(carboxymethoxy)-2,2'-bipyridine, 5-phenyl-1,10-phenanthroline, o-phenylenebis(diphenylphosphine)]. All diimine derivatives are emissive in fluid solution at 298 K and in frozen glasses at 77 K. The combination of detailed spectroscopic studies and theoretical calculations suggested that the emitting state at room temperature is 3MLCT in nature; namely [d(Re) → π*(diimine)] with the latter exhibiting partial π*(carbene) parentage. Subtle modification of the excited state energies (e.g. λ<sub>max</sub> 565–635 nm in acetonitrile) and redox potentials have been demonstrated using diimine auxiliaries with different electron-donating/accepting abilities.

4.4. Luminescent Re(V) and -(VI) benzylidyne complexes

Efficient methodologies for the synthesis of two series of luminescent rhenium(V) benzylidyne complexes,
namely (i) 2,4,6-trimethylbenzylidyne species bearing various phosphorus or nitrogen donor ligands (D), and (ii) para-substituted benzylidyne derivatives supported by o-phenylenediphenylphosphine (E, Fig. 7), have been developed [10,11]. The rhenium(V) benzylidyne complexes display photoluminescence at room temperature and 77 K. Data from electronic absorption and emission spectra, electrochemistry, and molecular orbital calculations provided evidence to suggest that the excited state for diimine-supported species is d(Re) \rightarrow \pi^*(X_2-bpy). In contrast, the emitting state for the remaining derivatives were assigned as [\pi(Re=CAr) \rightarrow \pi^*(Re=CAr)] with d(Re) \rightarrow \pi^*(\pi=CAr) MLCT character. Significantly, the excited-state properties associated with this class of chromophores have been modified with different benzylidyne and ancillary ligands. Hence the emission energy can be tuned from 520 to 610 nm in CH_2Cl_2; a range of 2840 cm^{-1}. The excited state lifetimes can be adjusted from 20 ns to 4.84 \mu s, many of which are noticeably longer than those observed in related Mo, W and Os alkylidyne/carbonyl systems [13–15].

The first examples of emissive d^1 complexes containing a metal–carbon multiple bond, namely Re^V(e=CAr)(PPh_3) (H_2) O X_2 (X = Cl and Br, Ar = 2,4,6-Me_3 C_6 H_2), were prepared from [Re^V(e=CAr)(PPh_3) (CO)(H_2) O] Cl _2 by treatment with HCl in chloroform and by photooxidation in dibromomethane respectively [12]. The crystal structure for X = Cl revealed a Re=C distance of 1.738(6) Å. Excitation of the Re(VI) benzylidyne species at \lambda > 300 nm in CH_2Cl_2 produced red emission (\lambda_{max} = 696 and 728 nm for X = Cl and Br, respectively) with lifetimes of ca. 0.4 \mu s.

5. Supplementary material

Crystallographic data for the structures reported in this paper have been deposited with the Cambridge Crystallographic Data Centre, CCDC no. 150179 for complex I. Copies of this information may be obtained free of charge from the Director, CCDC, 12 Union Road, Cambridge, CB2 1EZ, UK (fax: + 44-1223-336033; e-mail: deposit@ccdc.cam.ac.uk or http://www.ccdc.cam.ac.uk).

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