Stable Platinum(IV) Corroles: Synthesis, Molecular Structure, and Room-Temperature Near-IR Phosphorescence

Abraham B. Alemayehu, Laura J. McCormick, Kevin J. Gagnon, Sergey M. Borisov, and Abhik Ghosh

INTRODUCTION

The 5d metallocorroles constitute a unique class of size-mismatched complexes that incorporate a large 5d transition-metal ion within a sterically constrained macrocyclic ligand. Despite a steric mismatch inherent in their structures, the majority of them exhibit remarkable chemical and photochemical stabilities. A number of them also exhibit room-temperature near-IR phosphorescence, which has led to applications as oxygen sensors and as photosensitizers in photodynamic therapy and dye-sensitized solar cells. Platinum(IV) corroles, of which there has been only a single report, are particularly intriguing because of their potential for axial reactivity. They are, however, only accessible via a low-lying, serendipitously discovered reaction, which involves the interaction of a free-base corrole and Pt4(OAc)8 in benzonitrile at high temperature. The initially formed Pt(IV)-PhCN products reveal very short equatorial Pt-N distances of 1.94−1.97 Å, an axial Pt-C distance of 2.03 Å, and an axial Pt-N distance of 2.22 Å. The complexes exhibit Soret maxima at ~430 nm, which are essentially independent of the meso-aryl para substituents, and strong Q bands with the most intense peak at 595−599 nm. The substituent-independent Soret maxima are consistent with an innocent PtIV−corrole description for the complexes. The low reduction potentials (−1.45 ± 0.08 V vs saturated calomel reference electrode) also support a highly stable Pt(IV) ground state as opposed to a noninnocent corrole description. The reductions, however, are irreversible, which suggests that they involve concomitant cleavage of the Pt−aryl bond. Unlike Pt(IV) porphyrins, two of the complexes, PtIV[TPXPC]/(m-C6H4CN)(py) (X = CF3 and CH3), were found to exhibit room-temperature near-IR phosphorescence with emission maxima at 813 and 826 nm, respectively. The quantum yield of ~0.3% is comparable to those observed for six-coordinate Ir(III) corroles.

RESULTS AND DISCUSSION

As mentioned above, the Pt(IV) corroles PtIV[TPXPC]/(m-p-C6H4CN)(py) (X = CF3, H, and CH3) were obtained rather simply by the addition of pyridine to the reaction mixture at the end of the Pt insertion. For all compounds, purity and composition were established via thin-layer chromatography, X-ray structure determination. Here, we report that in situ exposure of the initially formed Pt(IV)-PhCN products to pyridine leads to a new class of stable, nonradical Pt(IV) corroles with the general formula PtIV[TPXPC]/(m-p-C6H4CN)(py), which have been variously characterized with single-crystal X-ray structure determination, electrochemical studies, and UV−vis−NIR absorption and emission spectroscopy (Figure 1). Although the results represent modest progress from a synthetic viewpoint, the physicochemical measurements afford significant insight into the electronic properties of a rare and elusive class of substances.
high-resolution electrospray ionization mass spectrometry, and 
1H NMR spectroscopy (Figures 2 and 3). Elemental analyses, 
however, could not be obtained because of the very small 
quantities available. Single-crystal X-ray structures could be 
obtained for two of the complexes, providing unambiguous 
proof of structure (Table 1). Both structures revealed a Pt 
atom located exactly or nearly exactly in the mean plane of a 
planar corrole ligand. For PtIV[TpCF3PC](m-C6H4CN)(py), 
the two axial ligands, m-C6H4CN and pyridine, were found to 
occupy symmetry-equivalent sites in the crystal, each with 50% 
occupancy, and were modeled such that the atoms of the two 
six-membered rings were superimposed (Figure 4). Accord-
ingly, the axial Pt−C/N distances for this structure only 
represent an average of the “true” Pt−C and Pt−N distances. 
Fortunately, the second structure, PtIV[TPC](m-C6H4CN)- 
(py) (TPC = triphenylcorrolato), was found to be fully 

Figure 1. Current status of Pt–corrole chemistry; the complexes prepared in the course of this study are schematically depicted in blue.

Figure 2. Representative 1H NMR spectrum: Pt[TpCF3PC](m-C6H4CN)(py).
ordered (Figure 5). The structures exhibit some of the shortest Pt–N distances known, which for the equatorial nitrogens are 1.955 ± 0.015 Å, reflecting the sterically constrained character of 5d metallocorroles. The axial Pt–C and Pt–N distances in the TPC complex are longer, 2.033(7) and 2.216(6) Å, respectively.

All six complexes exhibit slightly split Soret bands (Table 2 and Figures 6 and 7), which are essentially unaffected by the para substituents on the meso-aryl groups as well as strong Q bands. Over a long series of studies, we have shown that such substituent-insensitive Soret maxima are indicative of an innocent, nonradical corrole macrocycle, which is typical of the great majority of stable 4d and 5d metallocorroles, including MoO,9 RuN,10 OsN,11 TcO,12 ReO,13 and Au14−17 corroles as well as Mo18 and W biscorroles.19 In contrast, the Soret maxima of the Pt IV[TpXPC2−](m-/p-C6H4CN)(Ar) were found to redshift dramatically in response to increasing electron-donating character of the para substituent X, a phenomenon that is also observed for other noninnocent metallocorroles, such as MnCl,20,21 FeCl,22,23 FeNO,24,25 Fe2(μ-O),26 and Cu corroles.27

Cyclic voltammetry measurements were carried out for the meta-cyanophenyl series Pt IV[TpXPC](m-C6H4CN)(py), which could be obtained in somewhat higher yields than the para series (Figure 8 and Table 2). Given the instability of the Pt(V) state, the oxidation potentials, which range from 0.56 V (for X = CH3) to 0.74 V (for X = CF3), underscore the high stability of the PtIVAr−corrole unit toward reduction. That said, although the electrochemical HOMO-LUMO gap of 2.1 eV is typically indicative of a redox-inactive metal center and of ligand-centered oxidation and reduction,10−13,16,36 the fact that the reduction is irreversible suggests concomitant cleavage of the Pt−Ar bond.

Photophysical measurements were carried out on two of the complexes, Pt IV[TpXPC](m-C6H4CN)(py) for X = CF3 and CH3 (Table 3 and Figures 9 and 10). Both are clearly phosphorescent, which was confirmed by almost complete quenching of the emission in the presence of oxygen (Figure 9b,d), measurement of the decay time (Figure 10) and by acquisition of luminescence excitation spectra (Figure 9a,c). The latter are essentially identical to the absorption spectra; the small deviations are due to nonlinearities ascribable to strong absorption in the Soret region (the concentration used was necessary for obtaining high-quality emission spectra with excitation in the Q-band). The NIR phosphorescence is rather weak, but the quantum yields are in the same order of magnitude as those observed for Ir(III) corroles.2 This observation is interesting, considering that Pt(IV) porphyrins,57 in contrast to Pt(II) porphyrins,58−61 have been reported to be nonemissive.52 Weak red fluorescence (not quenchable by oxygen) was also clearly detected for the two compounds studied. The quantum yields for the fluorescence were estimated to be about an order of magnitude lower than

Figure 3. Representative 1H NMR spectrum: Pt[TpCF3PC](p-C6H4CN)(py).
those for the phosphorescence. Upconversion with a triplet annihilator, which proved feasible with OsN corroles, was found to be very weak due to the relatively low energy of the triplet state and the short triplet state decay times.

**CONCLUSIONS**

In what is only the second report on platinum corroles, we have described the synthesis of the first set of stable Pt(IV) complexes, in which the corrole is thought to be an innocent ligand (i.e., without radical character). These have the general formula PtIV[TpXPC](m/p-C6H4CN)(py), where X = CF3, H, and CH3. Although the yields are low (typically <5%), the compounds could be characterized with the standard spectroscopic methods and in two cases single-crystal X-ray crystallography providing rare insight into an elusive class of molecules. The structures revealed short equatorial Pt−N distance of 1.94−1.97 Å, an axial Pt−C distance of ∼2.03 Å, and an axial Pt−N distance of ∼2.22 Å. The UV−vis spectra revealed Soret maxima at ∼430 nm, which are essentially independent of the meso-aryl para substituents and strong Q bands with the most intense peak at 595−599 nm. The substituent-independent Soret maxima are consistent with an innocent PtIV−corrole description for the new complexes.

The low reduction potentials (−1.45 ± 0.08 V vs saturated calomel reference electrode (SCE)) also support a highly stable Pt(IV) ground state and rule out a corrole−2− description. The reductions, however, were found to be irreversible, which suggests that they involve concomitant cleavage of the Pt−aryl bond. Somewhat to our surprise and unlike Pt(IV) porphyrins, two of the complexes, PtIV[TpXPC](m-C6H4CN)(py) (X = CF3 and CH3), were found to exhibit room-temperature near-IR phosphorescence with emission maxima at 813 and 826 nm, respectively. The quantum yield of

<table>
<thead>
<tr>
<th>sample</th>
<th>Pr<a href="Ar">TpPC</a>(py)</th>
<th>Pr<a href="Ar">TpCF3PC</a>(py)</th>
</tr>
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<tr>
<td>chemical formula</td>
<td>C49H32N6Pt</td>
<td>C52H29F9N6Pt</td>
</tr>
<tr>
<td>formula mass</td>
<td>899.89</td>
<td>1103.90</td>
</tr>
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<td>monoclinic</td>
</tr>
<tr>
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<td>P1</td>
<td>P2_1/c</td>
</tr>
<tr>
<td>a (Å)</td>
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<td>0.8857</td>
</tr>
<tr>
<td>b (Å)</td>
<td>9.4792(15)</td>
<td>18.9584(10)</td>
</tr>
<tr>
<td>c (Å)</td>
<td>12.0922(19)</td>
<td>16.8577(8)</td>
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<tr>
<td>α (deg)</td>
<td>16.675(3)</td>
<td>14.0096(7)</td>
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<tr>
<td>β (deg)</td>
<td>109.102(3)</td>
<td>90</td>
</tr>
<tr>
<td>γ (deg)</td>
<td>95.415(3)</td>
<td>111.553(3)</td>
</tr>
<tr>
<td>Z</td>
<td>2</td>
<td>4</td>
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<tr>
<td>V (Å³)</td>
<td>1795.9(5)</td>
<td>4164.3(4)</td>
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<td>173(2)</td>
<td>100(2)</td>
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<td>density (g/cm³)</td>
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<td>1.761</td>
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<tr>
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<td>11,271</td>
<td>4,807</td>
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<td>restraints</td>
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<td>58</td>
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<tr>
<td>Rint</td>
<td>0.1151</td>
<td>0.0544</td>
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<tr>
<td>Rint, wRint, all data</td>
<td>0.0583, 0.1336</td>
<td>0.0482, 0.0746</td>
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<tr>
<td>δ (Goof) all data</td>
<td>1.033</td>
<td>1.037</td>
</tr>
<tr>
<td>max/min res. dens. (e/Å³)</td>
<td>3.436/−1.496</td>
<td>0.968/−0.872</td>
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“Ar = m-C6H4CN.”

![Figure 4](image_url)

Figure 4. Thermal ellipsoid plot for PtIV[TpCF3PC](m-C6H4CN)(py). Selected distances (Å): Pt1−N1 1.950(3), Pt1−N2 1.966(5), Pt1−N3 1.955(6), Pt1−N4 1.944(5), Pt1−N101 2.216(6), and Pt1−C201 2.033(7).

![Figure 5](image_url)

Figure 5. Selected distances (Å): Pt1−N1 1.944(5), Pt1−N2 1.944(5), Pt1−N3 1.955(6), Pt1−N4 1.944(5), Pt1−N101 2.216(6), and Pt1−C201 2.033(7).
∼0.3% is in the same order of magnitude as those of six-coordinate Ir(III) corroles.

**EXPERIMENTAL SECTION**

**Materials.** Free-base *meso*-triarylcorroles were synthesized according to a literature procedure. Platinuim(II) chloride was purchased from Sigma-Aldrich and used to synthesize tetranuclear platinum(II) acetate, as described in the literature. Platinum insertion reactions were carried out in a Biotage microwave reactor using 20 mL of microwave vials. Silica gel 60 (0.04−0.063 mm particle size, 230−400 mesh, Merck) was used for flash chromatography, and silica gel 60 preparative thin-layer chromatography (PTLC) plates (20 cm × 20 cm, 0.5 mm thick, Merck) were used for final purification of all complexes.

**Instrumental Methods.** UV−visible−NIR spectra were recorded on an HP 8454 spectrophotometer. ¹H NMR spectra were recorded on a 400 MHz Bruker AVANCE III HD spectrometer equipped with a 5 mm BB/¹H SmartProbe at 298 K in CDCl₃ and referenced to residual CHCl₃ at 7.26 ppm. Mass spectra were recorded on a Thermo Scientific LTQ Orbitrap XL spectrometer with an Ion-Max electrospray ion source. Cyclic voltammetry was carried out at 298 K with an EG&G model 263A potentiostat having a three-electrode system: a glassy carbon working electrode, a platinum wire counter electrode, and a saturated calomel reference electrode (SCE). Anhydrous CH₂Cl₂ (Aldrich) was used as solvent and tetrakis(n-butyl)ammonium perchlorate, recrystallized twice from absolute ethanol and dried in a desiccator for at least 2 weeks, was used as the supporting electrolyte. The reference electrode was separated from the bulk solution using a fritted glass bridge filled with the solvent/supporting-electrolyte mixture. The electrolyte solution was purged with argon for at least 2 min, and all measurements were carried out under an argon blanket. All potentials were referenced to the SCE. Emission and excitation spectra were acquired on a FluoroLog 3 spectrophotometer (Horiba Scientific) equipped with a NIR-sensitive R2658 photomultiplier (Hamamatsu). Relative quantum yields at room temperature were estimated using a solution of Pt(II) tetraphenyltetrabenzoporphyrin in toluene as a reference (∆Φ = 21%).

**Table 2. Spectroscopic and Electrochemical Properties: UV−vis λ_max (nm) and E_{1/2} Values (V) of Pt[TpXPC](m/p-C₆H₄CN)(py)**

<table>
<thead>
<tr>
<th>complex</th>
<th>λ_max (Soret)</th>
<th>λ_max (Q)</th>
<th>E_{1/2(ox2)}</th>
<th>E_{1/2(ox1)}</th>
<th>E_{1/2(red1)}</th>
<th>ΔE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pt<a href="m-C%E2%82%86H%E2%82%84CN">TpCF₃PC</a>(py)</td>
<td>430</td>
<td>569, 595</td>
<td>1.37</td>
<td>0.74</td>
<td>−1.37</td>
<td>2.11</td>
</tr>
<tr>
<td>Pt<a href="m-C%E2%82%86H%E2%82%84CN">TpPC</a>(py)</td>
<td>427, 437</td>
<td>567, 596</td>
<td>1.12</td>
<td>0.61</td>
<td>−1.49</td>
<td>2.10</td>
</tr>
<tr>
<td>Pt<a href="m-C%E2%82%86H%E2%82%84CN">TpCH₃PC</a>(py)</td>
<td>427, 438</td>
<td>567, 599</td>
<td>1.11</td>
<td>0.56</td>
<td>−1.53</td>
<td>2.09</td>
</tr>
<tr>
<td>Pt<a href="p-C%E2%82%86H%E2%82%84CN">TpCF₃PC</a>(py)</td>
<td>430</td>
<td>571, 595</td>
<td>1.37</td>
<td>0.74</td>
<td>−1.37</td>
<td>2.11</td>
</tr>
<tr>
<td>Pt<a href="p-C%E2%82%86H%E2%82%84CN">TpPC</a>(py)</td>
<td>427, 437</td>
<td>568, 597</td>
<td>1.11</td>
<td>0.56</td>
<td>−1.53</td>
<td>2.09</td>
</tr>
<tr>
<td>Pt<a href="p-C%E2%82%86H%E2%82%84CN">TpCH₃PC</a>(py)</td>
<td>427, 438</td>
<td>567, 599</td>
<td>1.11</td>
<td>0.56</td>
<td>−1.53</td>
<td>2.09</td>
</tr>
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</table>

**Table 3. Summary of Photophysical Properties Measured in Deoxygenated Toluene at 25 °C**

<table>
<thead>
<tr>
<th>complex</th>
<th>λ_max,em (nm)</th>
<th>Φ (%)</th>
<th>decay time (μs)</th>
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<tbody>
<tr>
<td>PtIV<a href="m-C%E2%82%86H%E2%82%84CN">TpCF₃PC</a>(py)</td>
<td>813</td>
<td>0.27</td>
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<tr>
<td>PtIV<a href="m-C%E2%82%86H%E2%82%84CN">TpCH₃PC</a>(py)</td>
<td>826</td>
<td>0.19</td>
<td>17.5</td>
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were recorded on 400 MHz Bruker AVANCE III HD spectrometer equipped with a 5 mm BB/¹H SmartProbe at 298 K in CDCl₃ and referenced to residual CHCl₃ at 7.26 ppm. Mass spectra were recorded on a Thermo Scientific LTQ Orbitrap XL spectrometer with an Ion-Max electrospray ion source.

Cyclic voltammetry was carried out at 298 K with an EG&G model 263A potentiostat having a three-electrode system: a glassy carbon working electrode, a platinum wire counter electrode, and a saturated calomel reference electrode (SCE). Anhydrous CH₂Cl₂ (Aldrich) was used as solvent and tetrakis(n-butyl)ammonium perchlorate, recrystallized twice from absolute ethanol and dried in a desiccator for at least 2 weeks, was used as the supporting electrolyte. The reference electrode was separated from the bulk solution using a fritted glass bridge filled with the solvent/supporting-electrolyte mixture. The electrolyte solution was purged with argon for at least 2 min, and all measurements were carried out under an argon blanket. All potentials were referenced to the SCE.

Emission and excitation spectra were acquired on a FluoroLog 3 spectrophotometer (Horiba Scientific) equipped with a NIR-sensitive R2658 photomultiplier (Hamamatsu). Relative quantum yields at room temperature were estimated using a solution of Pt(II) tetraphenyltetrabenzoporphyrin in toluene as a reference (∆Φ = 21%). The dye solutions were
deoxygenated in a screw-cap cuvette (Hellma, Mülheim, Germany) by bubbling argon through the solution for 12 min. Phosphorescence decay times were acquired in time domain on the FluoroLog 3 spectrofluorometer using a 390 nm SpectraLED (Horiba) as the excitation source.

General Procedure for the Synthesis of Pt[TpXPC](m/p-C6H4CN)(py), Where X = CF3, H, C2H5.

To a 20 mL microwave vial containing PhCN (5 mL) and a magnetic stirring bar were added a free-base corrole H3[TpXPC] (0.114 mmol) and Pt4(OAc)8·2HOAc (1 equiv). The vial was sealed and heated for 2 h at 150 °C under microwave irradiation. Upon completion of the reaction, pyridine (0.5 mL) was added and the contents of the vial were transferred to a round-bottom flask (50 mL) and evaporated to dryness. The resulting solid was dissolved in dichloromethane (5 mL) and loaded onto a silica gel column and eluted with a mixture of dichloromethane and n-hexane (the exact ratio of which is stated below for each case). All fractions containing Pt[TpXPC](m/p-C6H4CN)(py), characterized by a Soret λmax between 426 and 430 nm, were collected and evaporated to dryness. The product thus obtained was separated into the meta and para regioisomers with PTLC using a dichloromethane/n-hexane mixture as eluent, as indicated below.

Synthesis and Separation of Pt[TpCF3PC](m/p-C6H4CN)(py). The crude reaction product was chromatographed on a silica gel column with 3:2 dichloromethane/n-hexane as eluent. The fractions with a UV−vis λmax of 430 nm were collected and evaporated to dryness, resulting in a combined yield of 3.69 mg (6.6%) for the Pt[TpCF3PC](m/p-C6H4CN)(py) regioisomers. PTLC with 1:1 dichloromethane/n-hexane as eluent was then used to separate the m- and p-isomers; the top band was identified as Pt[TpCF3PC](m-C6H4CN)(py) and the lower band as Pt[TpCF3PC](p-C6H4CN)(py) based on 1H NMR analysis. Dark purple X-ray quality crystals of the meta isomer were grown by slow evaporation of a dichloromethane/n-hexane solution over a period of 15 days. Spectroscopic characterization data for the two isomers are as follows (see also Figures 2 and 3).

**Figure 9.** Optical properties of Pt(IV) corroles: (a, c) absorption and luminescence excitation spectra of the PtIV[TpCH3PC](m-C6H4CN)(py) and PtIV[TpCF3PC](m-C6H4CN)(py), respectively, in toluene solution at 25 °C; (b, d) luminescence spectra of PtIV[TpCH3PC](m-C6H4CN)(py) and PtIV[TpCF3PC](m-C6H4CN)(py), respectively, in toluene under anoxic and air-saturated conditions at 25 °C.

**Figure 10.** Phosphorescence decay for Pt(IV) corroles in anoxic toluene (25 °C, detected at 815 ± 7 nm).
hexane as eluent. The fractions with a graphed on a silica gel column with 2:1 dichloromethane/

δ doublets, 2H, 3o (PhCN) regioisomers. PTLC with 3:2 dichloromethane/

o 2H, 3o (PhCN) (py). Yield 1.17 mg (2.22%). UV–vis (CH2Cl2) λmax (nm, ε × 10^−4 M−1 cm−1): 427 (9.67), 438 (7.79), 496 (0.46), 528 (0.62), 567 (1.40), 599 (3.21). 1H NMR δ: 7.56 (d, 2H, 10-H), 7.22 (td, 2H, 7-H), 7.13 (m, 15-H), 7.03 (d, 2H, 8-H), 6.83 (d, 2H, 5-H), 2.67 (s, 3H, 10-CH3), 1.76 (br s, 2H, 2,6-py). Yield 1.08 mg (2.16%). UV–vis (CH2Cl2) λmax (nm, ε × 10^−4 M−1 cm−1): 427 (9.02), 496 (0.61), 528 (0.78), 568 (1.78), 597 (3.72). 1H NMR δ: 8.00 (d, 2H, 10-H), 7.83 (d, 2H, 10-H), 7.10 (d, 2H, 8-H), 6.89 (d, 2H, 5-H), 2.65 (s, 3H, 10-CH3), 1.72 (br s, 2H, 1,4-py). Yield 1.21 mg (2.36%). UV–vis (CH2Cl2) λmax (nm, ε × 10^−4 M−1 cm−1): 427 (9.19), 438 (7.59), 496 (0.41), 528 (0.62), 567 (1.47), 599 (3.28). 1H NMR δ: 7.98 (d, 2H, 10-H), 7.55 (d, 2H, 8-H), 7.02 (d, 2H, 5-H), 2.60 (s, 3H, 10-CH3), 1.70 (br s, 2H, 2,6-py). PT([T3PC](m-p-C6H4CN)(py)). Yield 2.3 mg (4.41%). UV–vis (CH2Cl2) λmax (nm, ε × 10^−4 M−1 cm−1): 427 (9.19), 438 (7.59), 496 (0.41), 528 (0.62), 567 (1.47), 599 (3.28). 1H NMR δ: 7.98 (d, 2H, 10-H), 7.55 (d, 2H, 8-H), 7.02 (d, 2H, 5-H), 2.60 (s, 3H, 10-CH3), 1.70 (br s, 2H, 2,6-py). Yield 1.08 mg (2.16%). UV–vis (CH2Cl2) λmax (nm, ε × 10^−4 M−1 cm−1): 430 (10.02), 498 (0.56), 531 (0.70), 571 (1.64), 595 (3.23). 1H NMR δ: 7.16 (d, 2H, JHH = 3.68 Hz, β-H), 8.92 (d, 2H, JHH = 5.20 Hz, β-H), 8.73 (d, 2H, JHH = 4.28 Hz, β-H), 8.68 (d, 2H, JHH = 5.02 Hz, β-H), 8.43 (d, 2H, JHH = 7.96 Hz, 5,15-o1-Pt). 8.35 (d, 2H, JHH = 7.96 Hz, 5,15-o2-Pt); 8.24 (overlapping doublets, 2H, JHH = 8.12 Hz, 10-o1-Pt and 10-o2-Pt); 8.06 (d, 4H, JHH = 8.12 Hz, 15-o1-Pt and 15-o2-Pt); 8.00 (overlapping doublets, 2H, JHH = 8 Hz, 10-o1-Pt and 10-o2-Pt); 6.33 (br s, 1H, 4-py); 5.46 (br s, 2H, 3,5-py); 5.19 (d, 2H, JHH = 6.88 Hz, C6H4CN ortho); 1.70 (br s, 2H, 2,6-py). 0.46 (d, 2H, JHH = 8.16 Hz, C6H4CN meta). HRMS (major isopomer): M+ = 1103.2118 (expt), 1103.1952 (calc for C49H32N6Pt). Synthesis and Separation of Pt([T3PC](m-p-C6H4CN)(py)). The crude reaction product was initially chromatographed on a silica gel column with 2:1 dichloromethane/n-hexane as eluent. The fractions with a δmax of 427 nm were collected and evaporated to dryness, resulting in combined yield of 4.17 mg (8.0%) for the Pt([T3PC](m-p-C6H4CN)(py)) regioisomers. PTLC with 3:1 dichloromethane/n-hexane as eluent was then used to separate the isomers; the top band was identified as Pt([T3PC](m-p-C6H4CN)(py)) and the lower band as Pt([T3PC](m-p-C6H4CN)(py)) based on 1H NMR analysis. Yield 1.08 mg (2.16%). UV–vis (CH2Cl2) λmax (nm, ε × 10^−4 M−1 cm−1): 430 (10.02), 498 (0.56), 531 (0.70), 571 (1.64), 595 (3.23). 1H NMR δ: 7.16 (d, 2H, JHH = 3.68 Hz, β-H), 8.92 (d, 2H, JHH = 5.20 Hz, β-H), 8.73 (d, 2H, JHH = 4.28 Hz, β-H), 8.68 (d, 2H, JHH = 5.02 Hz, β-H), 8.43 (d, 2H, JHH = 7.96 Hz, 5,15-o1-Pt). 8.35 (d, 2H, JHH = 7.96 Hz, 5,15-o2-Pt); 8.24 (overlapping doublets, 2H, JHH = 8.12 Hz, 10-o1-Pt and 10-o2-Pt); 8.06 (d, 4H, JHH = 8.12 Hz, 15-o1-Pt and 15-o2-Pt); 8.00 (overlapping doublets, 2H, JHH = 8 Hz, 10-o1-Pt and 10-o2-Pt); 6.33 (br s, 1H, 4-py); 5.46 (br s, 2H, 3,5-py); 5.19 (d, 2H, JHH = 6.88 Hz, C6H4CN ortho); 1.70 (br s, 2H, 2,6-py). 0.46 (d, 2H, JHH = 8.16 Hz, C6H4CN meta). HRMS (major isopomer): M+ = 1103.2118 (expt), 1103.1952 (calc for C49H32N6Pt). Synthesis and Separation of Pt([T3PC](m-p-C6H4CN)(py)). The crude reaction product was initially chromatographed on a silica gel column with 2:1 dichloromethane/n-hexane as eluent. The fractions with a δmax of 427 nm were collected and evaporated to dryness, resulting in combined yield of 4.17 mg (8.0%) for the Pt([T3PC](m-p-C6H4CN)(py)) regioisomers. PTLC with 3:1 dichloromethane/n-hexane as eluent was then used to separate the isomers; the top band was identified as Pt([T3PC](m-p-C6H4CN)(py)) and the lower band as Pt([T3PC](m-p-C6H4CN)(py)) based on 1H NMR analysis. Yield 1.08 mg (2.16%). UV–vis (CH2Cl2) λmax (nm, ε × 10^−4 M−1 cm−1): 430 (10.02), 498 (0.56), 531 (0.70), 571 (1.64), 595 (3.23). 1H NMR δ: 7.16 (d, 2H, JHH = 3.68 Hz, β-H), 8.92 (d, 2H, JHH = 5.20 Hz, β-H), 8.73 (d, 2H, JHH = 4.28 Hz, β-H), 8.68 (d, 2H, JHH = 5.02 Hz, β-H), 8.43 (d, 2H, JHH = 7.96 Hz, 5,15-o1-Pt). 8.35 (d, 2H, JHH = 7.96 Hz, 5,15-o2-Pt); 8.24 (overlapping doublets, 2H, JHH = 8.12 Hz, 10-o1-Pt and 10-o2-Pt); 8.06 (d, 4H, JHH = 8.12 Hz, 15-o1-Pt and 15-o2-Pt); 8.00 (overlapping doublets, 2H, JHH = 8 Hz, 10-o1-Pt and 10-o2-Pt); 6.33 (br s, 1H, 4-py); 5.46 (br s, 2H, 3,5-py); 5.19 (d, 2H, JHH = 6.88 Hz, C6H4CN ortho); 1.70 (br s, 2H, 2,6-py). 0.46 (d, 2H, JHH = 8.16 Hz, C6H4CN meta). HRMS (major isopomer): M+ = 1103.2118 (expt), 1103.1952 (calc for C49H32N6Pt).
C₆H₄CN)(py) were found to be twinned, the components were separated using the CELL_NOW program.⁴₅ Absorption corrections were applied with SADABS⁴⁶ for Pt[TpCPF₆]C₆H₄CN(py) and with TWINABS⁴⁷ for Pt[TPC]C₆H₄CN(py). The structures were solved by intrinsic phasing (SHELXT)⁴⁸ and refined by full-matrix least squares on F² (SHELXL-2014)⁴⁹ using the ShelXle GUI.⁵₀ All non-hydrogen atoms were refined anisotropically. Hydrogen atoms were geometrically calculated and refined as riding atoms.

The two axial ligands in Pt[TpCPF₆]C₆H₄CN(py), pyridine and C₆H₄CN, were found to occupy symmetry-equivalent sites within the crystal, each with 50% occupancy, and were modeled such that the atoms of the two six-membered rings were superimposed. The C and N atoms that coordinate to the Pt center (C101 and N101) were constrained to have identical x, y, and z coordinates via the EXYZ command in SHELX and were refined under separate PART instructions. Each of the remaining five atoms of the aromatic ring was modeled as common to both orientations with full occupancies, since attempts to independently model the two rings were unsuccessful. The CN and H substituents bound to C105 were refined under the same PART instructions as C101 and N101, respectively. The disordered axial ligands led to disorder in the unique C₆H₄CF₃ substituent, causing the CF₃ group to be positionally disordered over two symmetry-equivalent sites. The atoms belonging to this CF₃ group were refined with an occupancy of 0.5, but no attempt was made to model disorder in the aromatic ring of this substituent. Rotational disorder was also found for the CF₃ groups on the other two C₆H₄CF₃ substituents, and each CF₃ group was accordingly modeled over two orientations with complementary occupancies. Equivalent disordered atoms (e.g., N101/C101) were constrained to have equal Uᵦ values via the EADP command in SHELX. Additional crystallographic information has been summarized in Table 1, and full details can be found in the Crystallographic Information File provided as Supporting Information.

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**REFERENCES**


**ASSOCIATED CONTENT**

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsomega.8b01149.

Electrospray ionization mass spectra (PDF)

Crystal structures reported in this paper have been deposited at the Cambridge Crystallographic Data Centre and assigned the deposition numbers CCDC 1846980-1846981 (CIF).

**AUTHOR INFORMATION**

Corresponding Author

*E-mail: abhik.ghosh@uit.no.*

ORCID

Laura J. McCornick: 0000-0002-6634-4717
Sergey M. Borisov: 0000-0001-9318-8273
Abhik Ghosh: 0000-0003-1161-6364

Notes

The authors declare no competing financial interest.


