A total of three distinctive main group and transition metal complexes containing the 2-pyridyl pyrazolate (pypz) ligand were prepared, namely, [B(C6F5)3(pypz)] (1), [Ru(CO)2-(pypz)2] (2), and [Os(CO)2(pypz)2] (3), where (pypz)H = 3-trifluoromethyl-5-(2-pyridyl)pyrazole. Single-crystal X-ray diffraction studies were carried out on complexes 2 and 3, revealing octahedral coordination geometry with two CO ligands located at cis dispositions. While the pypz ligand arrangement for complex 2 is in cis-(Npy,Npy) and trans-(Npz,Npz), complex 3 reveals a different configuration, cis-(Npz,Npy) and trans-(Npy,Npy) (Npy for pyridine-N and Nz for pyrazolate donor sites). Similar to that of the in-situ-prepared pypz anion, the boron complex [B(C6F5)3(pypz)] (1) exhibits a strong emission centered at 380 nm, which is unambiguously assigned to fluorescence derived from the S1→S0 transition. In contrast to the nonluminescent behavior for Ru complex 2, the Os complex 3 displays unique, strong room-temperature phosphorescence, showing vibronic progressions at 430, 457, and 480 nm. The remarkable differences in photophysical properties were rationalized by a combination of π-electron accepting CO ligand, relative pypz orientation, and heavy-atom-enhanced spin–orbit coupling effects.

1. Introduction

Third-row transition metal complexes incorporating simple polypyridine ligands, such as 2,2′-bipyridine (bpy) or 1,10-phenanthroline (phen), and cyclometalated ligands, such as 2-phenylpyridine and benzoquinoline, have attracted a great deal of interest in recent years. Research in this area was principally motivated by the use of these complexes in the study of excited-state electron and energy transfer as well as the potential applications in the fabrication of organic light-emitting diodes (OLEDs). Isoelectronic transition metal ions such as Re(I), Os(II), and Ir(III), possessing a unique d6-electron configuration, are particularly attractive for these two metal–ligand interaction, long-lived excited states, and high luminescence efficiencies, which significantly improve the likelihood of energy transfer prior to radiative or nonradiative relaxation. The strong spin–orbit coupling expected for these heavy metal ions, with atomic numbers Z = 75–77, would lead to an efficient intersystem crossing from the singlet excited state to the triplet manifold. Furthermore, mixing singlet and triplet excited states via spin–orbit coupling, to a great extent, would also remove the spin-forbidden nature of the T1 → S0 radiative relaxation, resulting in highly intense phosphorescent emission. On the basis of these concepts, polypyridine and cyclometalated types of ligands tend to form rigid molecular frameworks with the aforementioned metal ions and then give rise to desirable absorption and emission characteristics. More specifically, they normally display intense ligand-centered π–π* chromophores in the ultraviolet region and a weaker metal-to-ligand charge transfer (MLCT) transition in the lower energy, visible region. The strong absorption and emission characteristics derived from the π–π* and MLCT energy levels

Figure 1. Structures of various B(III), Ru(II), and Os(II) complexes in this study.

In this paper, we wish to report the synthesis and characterization of a new series of 5-(2-pyridyl)pyrazole (pypz) metal complexes with formula $[M(CO)_2(pypz)_2]$ where $M = \text{Ru}$ (2) or Os (3-5) (see Figure 1). The structural and chemical characteristics of the 2-pyridyl pyrazole (pypz)H are akin to those of 2,2′-bipyridine (bipy). The pyrazole fragment to form a stable anionic ligand, subsequently providing neutral chelate complexes which, in sharp contrast to typical bipyridine ruthenium and osmium complexes reported in the literature, have a high tendency to form ionic metal complexes. The strong $\pi$-donor property of the pyrazolate, together with the $\pi$-accepting ability of the second pyridyl fragment, may provide a synergism of the electron delocalization so that the electron density is transferred from the pyrazolate to the metal ion and back to the pyridyl side of the ligand, enhancing the metal chelate interaction. For another comparison, although 2-phenylpyridine is effective in forming stable cyclometalated complexes with Rh(III), Ir(III), and Pt(II) ions, it was rather inert to the Ru(II) and Os(II) metal ions and particularly, to our knowledge, failed to afford any Os(II) complexes. Alternatively, 2-pyridyl pyrazole is expected to extend the chemistry of 2-phenylpyridine, providing a great potential to isolate the respective Ru(II) and Os(II) complexes. On the basis of this strategic design, the resulting new series of 2-pyridyl pyrazole incorporating Os(II) and Ru(II) complexes revealed remarkably different photophysical properties from previously reported analogues in that a strong ligand phosphorescence was resolved for $[\text{Os(CO)}_2(pypz)_2]$ complexes in room-temperature solution phase. Detailed results and discussion are elaborated in the following sections.

2. Experimental Section

2.1. General Information and Materials. Mass spectra were obtained on a J EOL SX-102A instrument operating in electron impact (EI) mode or fast atom bombardment (FAB) mode. $^1H$ and $^{13}C$ NMR spectra were recorded on Varian Mercury-400 or INova-500 instruments; chemical shifts are quoted with respect to the internal standard tetramethylsilane for $^1H$ and $^{13}C$ NMR data. Elemental analyses were carried out at the NSC Regional Instrumentation Center at National Chiao Tung University, Hsinchu, Taiwan. The pyrazole chelate ligands, 3-trifluoromethyl-5-(2-pyridyl)pyrazole, (pypz)H, 3-pentafluoromethyl-5-(2-pyridyl)pyrazole, (pypz)H, and 3-trifluoromethyl-5-(4-methyl-2-pyridyl)pyrazole, (pypz)H, were prepared according to the methods reported in the literature. All reactions were performed under a nitrogen atmosphere using anhydrous solvents or solvents treated with an appropriate drying reagent.

Synthesis of Complex 1. A 50 mL reaction flask was charged with 200 mg of $\text{B(C}_6\text{F}_5)_3$ (0.38 mmol) and 30 mL of anhydrous THF solvent. To this solution was added 70 mg of 3-trifluoromethyl-5-(2-pyridyl)pyrazole, (pypz)H (0.34 mmol), and the mixture was stirred at room temperature for 12 h. After that, the solution was concentrated to dryness and the mixture was heated at 30 °C for 1 h. The resulting crude product was filtered through a small plug of silica gel and was eluted with dichloromethane. The concentrated solvent was evaporated to give a colorless crystalline solid, $[\text{B(C}_6\text{F}_5)_2(pypz)]$ (90 mg, 0.16 mmol, 48%).

Spectral data of $[\text{B(C}_6\text{F}_5)_2(pypz)]$: MS (EI), $m/z$ 557, M+; 538, (M – F)+; 390, (M – C_6F_5)+; $^1H$ NMR (400 MHz, CDCl_3, 294 K): δ 8.59 (dd, $j_{HH} = 6.0$ Hz, $j_{HH} = 1.2$ Hz, 1H, CHpy), 8.28 (dd, $j_{HH} = 8.59$ Hz, $j_{HH} = 1.2$ Hz, 1H, CHpy, 8.28 (dd, $j_{HH} = 8.59$ Hz, $j_{HH} = 1.2$ Hz, 1H, CHpy)


product was then subjected to recrystallization in CH2Cl2 and vacuum and placed in an oven maintaining temperatures of and degassed. This mixture/solution was then sealed under (150 mg, 0.165 mmol) were loaded in a 20 mL Carius tube heated for 3 days, the crude product was purified by sublimation (150 mTorr/165 °C), followed by crystallization from a mixture of CH3Cl/hexane, giving the ruthenium complex [Ru(CO)4(pypz2)] (2) as colorless rectangular crystals (191 mg, 0.33 mmol, 70%).

**Synthesis of Complex 2.** 3-Trifluoromethyl-5-(2-pyridyl)-pyrazole, (pypz)H (620 mg, 2.91 mmol), Ru3(CO)12 (300 mg, 0.47 mmol), and hexane solvent (50 mL) were added to a 160 mL stainless steel autoclave. The autoclave was sealed and slowly brought up to 185 °C for 36 h. After that, the autoclave was cooled, the solvent was evaporated to dryness, and the solid residue was purified by column chromatography on SiO2, eluting with a 1:1 mixture of ethyl acetate and hexane. Only one major component was obtained. Removal of excess solvent produced a light yellow solid, which was purified by sublimation (150 mTorr/165 °C), followed by crystallization from a mixture of CH3Cl/hexane, giving the ruthenium complex [Os(CO)2(pypz3)] (1) as colorless needle crystals (43 mg, 0.043 mmol) with a 39% yield.

**Synthesis of Complex 3.** 3-Trifluoromethyl-5-(2-pyridyl)-pyrazole (220 mg, 1.03 mmol) and finely pulverized Os5(CO)12 (150 mg, 0.165 mmol) were loaded in a 20 mL Carius tube and degassed. This mixture/solution was then sealed under vacuum and placed in an oven maintaining temperatures of 180–185 °C for 2.5 days, during which time its color changed from brown to red. After the reaction was stopped, the tube was cooled and opened, and the contents were dissolved in acetone. The insoluble material was filtered off, the filtrate dried under vacuum, and the residue sublimed (300 mTorr/185 °C). The product was then subjected to recrystallization in CH3Cl2 and hexane, giving [Os5(CO)12(pypz3)] (3) as colorless needle-like crystals (43 mg, 0.043 mmol) with a 39% yield.

**Spectral data of 2:** MS (EI, 70 eV), observed m/z (actual) [assignment] (relative intensity): 582 (582) [M–2CO] (2.88), 562 (526) [M–3CO] (12.5), IR (CH2Cl2): ν(CO), 2043 (s), 1970 (s) cm–1, 1H NMR (500 MHz, d6-acetone, 294 K): δ 8.07–8.03 (m, 4H, Hpy), 7.38 (d, JH= 1H, 2H, Hp), 7.32 (d, JH= 6, 6, and 3 Hz, 2H, Hp), 7.09 (d, JH= 6 and 1H, 2H, Hp). 13C NMR (125 MHz, d6-acetone, 294 K): δ 194.7 (CO), 154.4 (CO), 150.7 (CO), 148.9 (CHpy), 146.7 (q, JCF = 36.6 Hz, CHpy), 141.7 (CNpy), 124.5 (CHpz), 123.1 (q, JCF = 266.3 Hz, CF3), 121.7 (CHpz), 104.3 (CHpz). 19F NMR (470 MHz, d6-acetone, 294 K): δ −84.4 (s, CF3), −109.1 (d, JCF = 276.8 Hz, CF3), −111.4 (d, JCF = 276.8 Hz, CF3). Anal. Calcld for C32H14F6N6O2Os: C, 37.82; N, 10.91; H, 1.31. Found: C, 37.69; N, 12.01; H, 2.08.}

**Synthesis of Complex 3.** In a fashion similar to the synthesis of complex 2, 150 mg of Os5(CO)12 (0.165 mmol) and 240 mg of 3-trifluoromethyl-5-(4-methyl-2-pyridyl)pyrazole, (pypz3)H (1.04 mmol) were used. After being heated for 4 days, the crude product was purified by sublimation (0.24 Torr, 220 °C), followed by recrystallization with CH3Cl2 and hexane, giving a colorless crystalline solid [Os5(CO)12(pypz3)] (5, 34 mg, 0.048 mmol) with a 29% yield.

**Spectral data of 5:** MS (EI, 70 eV), observed m/z (actual) [assignment] (relative intensity): 700 (700) [M+5] (3.33), 644 (M–2CO) (14.9), IR (CH2Cl2): ν(CO), 2041 (s), 1970 (s) cm–1, 1H NMR (400 MHz, d6-acetone, 294 K): δ 8.97 (d, JH= 6.0 Hz, 2H, Hp), 7.95 (s, 2H, Hp), 7.31 (d, JH= 6.0 Hz, 2H, Hp), 7.06 (s, 2H, Hp), 2.58 (s, 6H, Me). 13C NMR (125 MHz, d6-acetone, 294 K): δ 177.8 (CO), 156.2 (CNpy), 153.2 (CNpy), 151.8 (CNpz), 144.0 (q, JCF = 35.4 Hz, Cp), 126.2 (CHpz), 122.3 (q, JCF = 241.8 Hz, CHpy), 122.1 (CHpy), 103.1 (CHpz), 21.2 (Me). 19F NMR (470 MHz, d6-acetone, 294 K): δ −59.8 (s). Anal. Calcld for C22H14F6N6O2Os: C, 37.82; N, 12.03; H, 2.02. Found: C, 37.69; N, 12.01; H, 2.08.

**2.2. Measurements.** Single-crystal X-ray diffraction data were measured on a Nonius Kappa or a Bruker SMART CCD diffractometer using λ(Mo Kα) radiation (λ = 0.7073 Å). The data collection was executed using the SMART program. Cell refinement and data reduction were made with the SAINT program. The structure was determined using the SHELX97 PC program and refined using full-matrix least-squares. All non-hydrogen atoms were refined anisotropically, whereas hydrogen atoms were placed at the calculated positions and included in the final stage of refinements with fixed parameters. Crystallographic refinement parameters of complexes 2 and 3 are summarized in Table 1, and the selective bond distances and angles of these complexes are listed in Tables 2 and 3, respectively.

Steady-state absorption and emission spectra were recorded with a Hitachi (U-3310) spectrophotometer and an Edinburgh (F5920) fluorimeter, respectively. Nanosecond–microsecond lifetime studies were performed with an Edinburgh FL 900 photon-counting system with a hydrogen-filled lamp or a nitrogen lamp as the excitation source. The emission decays were analyzed by the sum of exponential functions, which allows partial elimination of instrument time broadening and thus renders a temporal resolution of ~200 ps. Occasionally, for the long-lived (μs) emission species the lifetime was measured with the laser photolysis technique in which the third or fourth harmonic of an Nd:YAG laser (8 ns, Continuum Surlite II) was used as the excitation source, coupled with a fast response photomultiplier (Hamamatsu model R5509-72) operated at −80 °C. Typically, an average of 512 shots were acquired for each measurement. The transient absorption signal was recorded by a laser flash photolysis system (Edinburgh LP920), in which an Nd:YAG laser (355 nm) pumped optical parametric oscillator and a white-light square pulse were used as pump and probe beams, respectively. The temporal resolution was limited by the excitation pulse duration of ~8 ns.
Table 1. Crystal Data and Structure Refinement Parameters for Complexes 2 and 3

<table>
<thead>
<tr>
<th>Complex</th>
<th>Crystal System</th>
<th>Space Group</th>
<th>Unit Cell Parameters [Å]</th>
<th>Temperature</th>
<th>Data Collection</th>
<th>Refinement</th>
<th>Goodness of Fit</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>Orthorhombic</td>
<td>Pbca</td>
<td>a = 12.9716(7) b = 13.4557(1) c = 17.3441(2)</td>
<td>180 K</td>
<td>4940 (000), 4940 (000), 4940 (000)</td>
<td>R1 = 0.0456, wR2 = 0.1212</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>Monoclinic</td>
<td>P21</td>
<td>a = 2.21 to 27.50° b = 2.21 to 27.50° c = 13.4557(1)</td>
<td>180 K</td>
<td>4940 (000), 4940 (000), 4940 (000)</td>
<td>R1 = 0.0456, wR2 = 0.1212</td>
<td></td>
</tr>
</tbody>
</table>

3. Results

3.1. Preparation and Spectroscopic Characterization. 2-Pyridyl pyrazole, (pypz)H, can readily react with various metal source reagents, affording metal chelate complexes in high yields. The first example is represented by the direct treatment of B(C6F5)3 and (pypz)H at room temperature, giving a tetrahedrally coordinated boron complex, [B(C6F5)3(pypz)] (1) (eq 1).

\[
\text{B(C}_6\text{F}_5\text{)}_3 + \text{(pypz)H} \rightarrow [\text{B(C}_6\text{F}_5\text{)}_3(\text{pypz})] (1) + \text{C}_6\text{F}_5\text{H}
\]

Characterization of this boron complex was achieved using the routine spectroscopic methods as well as the previously determined crystal structural data of the phynyl-substituted boron complex [B(C6H5)3(pypz)]. It was found that the pyridyl nitrogen atom (Npy) and the nearby nitrogen atom of the pyrazole were both coordinated to the boron atom, forming a five-membered \( \text{C}_5\text{N}_2\text{B} \) chelate interaction. A similar bonding arrangement was observed in other tetrahedrally arranged \( \text{BPH}_2 \) complexes with chelating ligands, such as (2-pyridyl)-7-azaindole and (2-pyridyl)-7-indole.13

A similar straightforward route was applied to the synthesis of the transition metal carbonyl complexes involving Ru(II) and Os(II) cations. For the complex [Ru3(CO)6(pypz)2](2), the reaction was best carried out in hexane, using a stainless steel autoclave to raise the reaction temperature to 185 °C. It is possible that the reaction proceeds in multistep processes involving prior cluster fragmentation14 to afford mononuclear intermediates such as Ru(CO)4x 4 ≤ x ≤ 2, together with pyrazole addition, \( \text{H}_2 \) elimination, and CO displacement. The overall transformation is thus depicted as

\[
\text{Ru}_3\text{(CO)}_{12} + 6(\text{pypz})\text{H} \rightarrow \text{3[Ru(CO)}_2\text{(pypz)}_2]\text{(2)} + \text{3H}_2 + 6\text{CO} (2)
\]

The Ru complex 2 was isolated in ~70% yield after conducting column chromatography and recrystallization. The electron impact mass spectrometry revealed molecular ion \( \text{M}^+ \) with the expected composition \( \text{C}_{30}\text{H}_{10}\text{N}_6\text{O}_{2}\text{Ru} \), providing evidence for the incorporation of two pyridyl pyrazolate ligands. For the respective reaction with the heavy congener Os3(CO)12, due to its higher thermal stability and lower reactivity, the condensation reaction was best carried out using a technique of direct solid-state pyrolysis and employing a much longer reaction time of ~2.5 days. The osmium analogue complex [Os3(CO)6(pypz)2] (3) was then obtained in 39% yield, after vacuum sublimation and recrystallization from a mixture of CH2Cl2 and hexane at room temperature.

Os₃(CO)₁₂ + 6(pypz)H →
3[Os(CO)₂(pypz)₂] (3) + 3H₂ + 6CO (3)

For Os complex 3, ¹H, ¹³C, and ¹⁹F NMR studies confirmed the presence of 2-pyridyl pyrazolate and CO ligands in an expected ratio of 2:2 within the ligand sphere. An extension of preparing analogues of this Os complex is available by the variation of substituents on the 2-pyridyl pyrazolate ligand. This modification gave two additional derivative complexes, [Os(CO)₂(pypz₂)] (4) and [Os(CO)₂(pypz₃)] (5), pypz₂ = 3-pentafluoroethyl-5-(2-pyridyl) pyrazolate and pypz₃ = 3-trifluoromethyl-5-(4-methyl-2-pyridyl) pyrazolate. Their corresponding analytical and spectroscopic data as well as detailed spectral assignments are shown in the Experimental section.

All three Os(II) complexes 3–5 show excellent thermal stability at temperatures ~200 °C as well as good photochemical stability upon exposure to intensive UV irradiation. Another unique feature is that the introduction of a stronger electron-withdrawing substituent, such as CF₃ or C₂F₅, at the C(3) position of the pyrazole segment is indispensable to the successful preparation of the more robust third-row Os metal complexes. In this study, numerous attempts have been made to synthesize other [Os(CO)₂(pypz)₂] analogues based on methyl, tert-butyl, and phenyl substitution at the C(3) position, but unfortunately all failed, demonstrating the feasibility of synthesizing 3–5 mainly due to the greater reactivity of perfluoroalkyl-substituted pyrazole ligands.

3.2. Structural Characterization. Single-crystal X-ray diffraction studies were carried out to resolve the exact molecular structures. As depicted in Figure 2, the structure of 2 shows an octahedral coordination around the Ru atom. The 2-pyridyl pyrazolate ligands form five-membered chelate ring structures with the pyrazolate nitrogen atoms N(2) and N(5) located at trans positions, while the cis carbonyl fragments occupy the sites trans to the 2-pyridyl nitrogen atoms N(1) and N(4), which are also arranged in the cis (N,N) fashion. The coordination sequence of the pairs of donor centers CO, N(py), and N(pz) determines the observed cis–cis–trans conformation. The corresponding bond length and bond angle data are listed in Table 2. Metal–ligand bond angles found in this complex are within the range expected for typical octahedral Ru(II) complexes. The major deviation from a perfect octahedral coordination is caused by the smaller bite angles observed for the 2-pyridyl pyrazolate chelating interactions (∠N(1)–Ru–N(2) = 77.5(1)° and ∠N(4)–Ru–N(5) = 77.1(1)°), while the overall ligand arrangement in complex 2 resembles that of the structurally characterized Ru(II) complexes such as [Ru(CO)₂(hfac)₂]₁⁵ [Ru(CO)₂(1,2-naphthaquinone-1-oximato)₂]₁⁶ and [Ru(CO)₂(2-pyridylcarboxylato)₂]¹⁷ possessing a pair of cis carbonyl ligands and the donor atoms of the bidentate chelates located in the remainder of the coordination sites. Moreover, the pyrazolate nitrogen atoms exhibit a much stronger donor interaction with the Ru(II) center than the dative interaction from the 2-pyridyl nitrogen atoms. Evidence for this viewpoint is provided by the significantly shorter Ru–N(pz) bond distances (Ru–N(2) = 2.058(3) Å and Ru–N(5) = 2.047(3) Å) versus those of the Ru–N(py) distances (Ru–N(1) = 2.136(3) Å and Ru–N(4) = 2.140(3) Å). This deviation could result from a combination of (i) the stronger donor character of the pyrazolate nitrogen atoms due to its −1 charge nature and (ii) the relatively stronger trans influence exerted by the carbonyl ligands, which significantly weaken the N(py)–Ru dative interaction. In good agreement with this postulation, the long Ru–N(py) distances in complex 2 compare well with those observed in the related Ru(II) metal complexes such as [Ru(CO)(py)(TTP)] (2.193(4) Å), TTP = tetraphenylporphinate and [Ru(bpy)(CO)₂Cl₂] (2.10(1) Å),¹⁸ showing the pyridine group located trans to the strong N-accepting carbonyl ligand.

A perspective view of the Os complex 3 together with the atomic numbering scheme is illustrated in Figure 3, and selected bond lengths and angles are listed in Table 3. To our surprise, the coordination environment around the Os atom, which shows an ideal C₂ symmetry, displays a notable distinction from that established for the previous Ru analogue 2. This is revealed by the detection of pyrazolate nitrogen atoms N(1) and N(1A) being located at the position trans to the cis-oriented carbonyl ligands. Although this subtle change of ligand orientation on the ligand sphere has had almost no effect on the chelate bite angle, cf. ∠N(1)–Os(1)–N(2) = 77.3(1)°, the nitrogen to osmium dative bonding interaction has turned substantially stronger, as is evident from the observation of unusually short Os–N distances (Os(1)–N(1) = 2.013(3) Å and Os(1)–N(2) = 2.087(3) Å). The latter is very close to the Os–N(bpy) distances (2.036–2.076 Å) observed in the related Os(II) complexes such as [Os(bpy)₂(Cl)(NMe)₃]⁺[PF₆]⁻ and [Os(bpy)₃]⁺[PF₆]₂¹⁹. The results may be rationalized by the fact that the third-row transition metal elements tend to have stronger ligand-to-metal interactions compared to the first-row transition metals.
pared with their second-row congeners. In addition, the –1 charged pyrazolate nitrogen atoms are located at the positions trans to the good π-acceptor CO ligand, which can effectively remove the excess charge density that may accumulate at the metal center. As a result, the N(pz)–Os dative interaction is strengthened, resulting in a reduction of bond distances. IR ν(CO) spectral studies of Os complex 3 in CH₂Cl₂ solution revealed two CO stretching bands at 2043 and 1973 cm⁻¹, which are at lower frequencies than those (2076 and 2017 cm⁻¹) observed for the respective Ru complex 2. The results are in good agreement with the delineation, suggesting the occurrence of a stronger π-acceptor interaction within the Os complexes. For comparison, a reverse trend was observed in their parent carbonyl complexes, in which the ν(CO) stretching bands of Os₃(CO)₁₂ occurred at least 4–8 cm⁻¹ higher in frequency than those of the Ru₃(CO)₁₂ counterpart.²⁰

3.3. Photophysical Properties. Figure 4 shows UV–visible absorption spectra of the 2-pyridyl pyrazolate complexes in acetonitrile. For metal complexes 2 and 3, the dominant absorption bands in the 245–316 nm region were ascribed to the ligand-centered π→π* transitions. These spectral assignments were based on the absorption spectra of the related boron complex 1, as well as the corresponding free ligands (shown later in Figure 6). Attempts have also been made to examine any lower energy absorption at >360 nm, presumably associated with typical MLCT transitions. However, no absorption bands could be resolved in the region of 380–700 nm, suggesting that all MLCT transitions in complexes 2 and 3 are hidden in the UV region of the strong intra-ligand π→π* transitions (vide infra). Absorption spectra similar to those for complex 3 were observed for Os(II) complexes 4 and 5, and their relevant photophysical data are listed in Table 4.

Taking complex 3 as a prototype, a typical emission spectrum of 5-(2-pyridyl) pyrazolate Os complexes is depicted in Figure 4. Despite the diffusive S₀ → S₁ (π→π*) absorption band, the corresponding emission spec-

Figure 3. ORTEP diagram of complex 3 with thermal ellipsoids shown at 30% probability; the fluorine atoms of the CF₃ substituent on the pyrazolate fragments were removed for clarity.

Figure 4. Room-temperature UV–vis absorption and emission spectra of the pypz complexes 1 (–), 2 (–Δ–), and 3 (–○–) in CH₃CN, in which the emission of 2 is not detectable. Inset: Relaxation dynamics of emission for 3 in room-temperature degassed CH₃CN.

Figure 5. Temporal evolution of transient absorption spectra for complex 3 at a delay time of (a) 0.36, (b) 6, (c) 11.7, (d) 17.4, (e) 28.8, and (f) 40 μs in degassed CH₃CN; (g) transient absorption spectrum of 3 acquired at a delay time of 2 μs in aerated CH₃CN. Inset: Decay profile of the transient signal at 550 nm (in degassed CH₃CN). λ₀ex: 355 nm.

trum exhibits a distinct vibronic feature with peak maxima at 430, 457, and ~480 nm in CH₃CN. The luminescence intensity is linearly proportional to the increase of concentrations, eliminating the emissions associated with any high-order aggregation. The entire emission band originating from a common ground-state species is ascertained by the same fluorescence excitation spectra throughout the monitored wavelengths of 420–600 nm. The excitation spectra, within the range of experimental error, are also effectively identical to the absorption spectrum, indicating that the entire emission results from a common Franck–Condon excited state. From comparison of the corresponding absorption and emission spectra, we can make several remarks regarding complex 3. First of all, there is a significantly large energy gap of ~4000 cm⁻¹ for the 0–0 vibronic onsets between absorption and emission spectra. In comparison, complex 1 exhibits an emission band

Figure 6. (a) Room-temperature UV–vis absorption (---) and fluorescence (○○○) spectra of pypz anionic species (pypzK) in CH$_3$CN containing 1 M KOH. (b) Phosphorescence spectrum of pypz anion (--) in the 77 K solid (CH$_3$CN) matrix acquired at a delay time of 1 μs. * denotes Raleigh scattering.

Table 4. Photophysical Properties of Complex 1–5 in Room-Temperature CH$_3$CN

<table>
<thead>
<tr>
<th>complex</th>
<th>absorption (S$_0$–S$_1$)</th>
<th>emission</th>
<th>Q.Y.</th>
<th>$\tau$decay</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>319 (ε = 14 250)$^b$</td>
<td>380</td>
<td>0.88</td>
<td>6.4 ns</td>
</tr>
<tr>
<td>2</td>
<td>300 (ε = 14 000)</td>
<td>462</td>
<td>50.5 μs$^c$</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>311 (ε = 16 500)</td>
<td>430</td>
<td>0.14</td>
<td>18.5 μs</td>
</tr>
<tr>
<td>4</td>
<td>314 (ε = 16 120)</td>
<td>430</td>
<td>8.69 $\cdot$ 10$^{-2}$</td>
<td>13.4 μs</td>
</tr>
<tr>
<td>5</td>
<td>306 (ε = 17 200)</td>
<td>428</td>
<td>4.06 $\cdot$ 10$^{-2}$</td>
<td>6.3 μs</td>
</tr>
</tbody>
</table>

$^a$ All samples were degassed via three freeze–pump–thaw cycles. $^b$ Units in M$^{-1}$ cm$^{-1}$. $^c$ Data were obtained at 77 K.

maximized at 380 nm, in which the 0–0 onset overlaps with that of the S$_0$–S$_1$ (π–π$^*$) absorption band. Second, the Stokes shift, defined as the peak-to-peak frequency between absorption and emission, is as large as 9000 cm$^{-1}$. Since the emission peak wavelength revealed a slightly hypsochromic shift from cyclohexane (460 nm) to acetonitrile (457 nm), the anomalously large Stokes shifted emission resulting from the solvent dipolar relaxation also can be discarded. Alternatively, these observations, in combination with the relaxation dynamics, O$_2$ quenching effect, and transient absorption spectra elaborated as follows, lead us to propose that the emission, more plausibly, originates from a triplet manifold.

The inset of Figure 4 shows the relaxation dynamics of emission for complex 3 in room-temperature degassed CH$_3$CN. The logarithm plot of the time-dependent emission intensity is apparently linear and thus can be well fitted straightforwardly by the first-order relaxation dynamics. A best linear least-squares fit gave the lifetime of complex 3 as 18.3 μs. The same method was applied for complexes 4 and 5 with corresponding lifetimes of 13.4 and 6.3 μs, respectively (see Table 4). Upon aeration of the solution, the lifetime of complex 3 was drastically reduced to 310 ns, accompanied by the decrease of the steady-state emission intensity (not shown here). Taking an O$_2$ concentration of 1.9 $\times$ 10$^{-3}$ M in the aerated CH$_3$CN solvent,$^{21}$ a quenching rate constant of $\sim$1.7 $\times$ 10$^9$ M$^{-1}$ s$^{-1}$ was thus deduced, which is nearly 1/9 of the diffusion-controlled rate of 1.8 $\times$ 10$^{10}$ M$^{-1}$ s$^{-1}$ calculated from the Stokes–Einstein equation$^{22}$ in CH$_3$CN. The result is consistent with the O$_2$ quenching triplet state according to the theory of electron-exchange type energy transfer expressed as

$$T + ^3O_2 \rightarrow S_0 + ^1O_2$$

The overall spin must be conserved upon forming a collisional complex. Accordingly, the possibility of each collision generating $^1$O$_2$ is statistically 1/9.

Another key support for the triplet-state population was rendered by the transient absorption experiment. Figure 5 shows the temporal spectral evolution of the transient absorption for complex 3, consisting of a well-resolved band maximized at 520 nm and the other one at <370 nm that was irresolvable due to spectral interference with the ground-state absorption. As shown in the inset of Figure 5, a plot of the transient absorbance versus the delay time reveals a biexponential feature. In other words, the logarithm plot of the transient absorbance does not reveal a straight line (not shown here) and thus cannot be simply fitted by first-order decay kinetics. With an increase in the laser intensity the non-single-exponential decay behavior became more obvious, indicating that the bimolecular quenching process, i.e., the triplet–triplet annihilation, plays a role in the relaxation dynamics. In a degassed solution where O$_2$ has been completely removed, the decay pathways of a triplet-state species incorporating triplet–triplet annihilation can be expressed as

$$T \rightarrow S_0$$

$$T + T \rightarrow S_1 + S_0$$

By solving the associated kinetic equations, the time-dependent triplet-state concentration [T] is depicted as

$$[T] = \frac{k_T[T]_0}{(k_T + k_{TT}[T]_0)e^{k_T[0]} - k_T[0]}$$

(3)

where [T]$_0$ is the concentration of the triplet-state species prepared right after the laser excitation. Under the experimental condition of 1 cm excitation path length, the time-dependent transient absorbance $\Delta A$ should theoretically be proportional to [T] by $\Delta A = \epsilon_{TT}[T]$, where $\epsilon_{TT}$ is the extinction coefficient of the triplet–triplet absorption. Letting $b = k_{TT}/\epsilon_{TT}$, eq 3 can thus be rewritten as

$$\Delta A = \frac{k_T\Delta A_0}{(k_T + b\Delta A_0)e^{k_T} - b\Delta A_0}$$

(4)

where $\Delta A_0$ is the signal intensity right after the laser pulse. The decay of the transient absorbance depicted in the inset of Figure 5 was well fitted by eq 4, and the

(22) For example, see: Birks, J. B. Photochemistry of Aromatic Molecules; Wiley: New York, 1970; p 313.
lifetime was deduced to be 17.2 μs, which, within the range of experimental error, is identical to the one (18.3 μs) derived from the emission relaxation dynamics. Note that due to the rather small triplet-state population, the triplet–triplet annihilation is negligible in the emission study using the time-correlated single-photon-counting technique. The transient absorption originating from the study using the time-correlated single-photon-counting that due to the rather small triplet-state population, the range of experimental error, is identical to the one (18.3 μs) (see Figure 5g). Similar emission properties such as spectral features, long lifespan (>μs), and O₂ quenching dynamics were observed for complexes 4 and 5, and the corresponding photophysical data are listed in Table 4.

Upon unambiguously determining the dominant emission in Os(II) complexes 3–5 to be of phosphorescent nature, our next task focuses on the origin of the phosphorescence. The appearance of phosphorescence with vibronic progression has provided a clue to the 3π−π* configuration, which plausibly originates from the ligand-center chromophore. To validate this assignment, we have carried out a series of comparative studies focusing on the photophysics of the ligand 5-(2-pyridyl) pyrazole (pypzH₂). Due to the deprotonation of pypzH in forming the corresponding Os(II) complex, a fair comparison should be with its anion species. Figure 6a shows the room-temperature absorption and emission spectra of the pypz anion (pypzK), in which the room-temperature emission consists of a unique, normal Stokes-shifted fluorescence maximized at 372 nm (τ₁ ~ 438 ns). No longer wavelength emission ascribed to phosphorescence could be resolved in room-temperature CH₃CN. Figure 6b shows the emission of pypz²⁻ anion in a 77 K solid matrix (CH₃CN) acquired after a delay time of 1 μs to eliminate the fluorescence interference. Apparently, a structured phosphorescence maximized at ~460 nm was resolved, of which the spectral feature resembles that observed in complex 3. This result unambiguously indicates that the room-temperature phosphorescence in the title Os complexes originates from the ligated pypz chromophore. Note that the main difference in phosphorescence properties between the free pypz anion and complex 3 lies in the τp > 1 s phosphorescence lifetime (at 77 K) in the pypz anion, which is longer than that in complex 3 by ~5 orders of magnitude, manifesting the unusually strong spin–orbit coupling in complex 3. Similar phosphorescence properties can be ascribed to compounds 4 and 5. However, due to the relatively small radiative decay rate, the radiationless decay pathway induced by methyl (or CF₃) torsional motion may play a key role in the quenching of phosphorescence. Accordingly, among compounds 3–5 the smallest phosphorescence quantum yield for complex 5 can be rationalized because of its bearing two methyl-like rotors (–CH₃ and –CF₃). Likewise, the phosphorescence quantum yield of complex 4 with C₂F₅ substituent, being smaller than that of complex 3 with CF₃, is plausibly due to the more torsional degrees of freedom in C₂F₅.

Another key comparison was made via the study of complex 2, containing a Ru(II) metal ion. Intriguingly, under the detection limit of ~10⁻⁵ quantum efficiency no detectable emission can be resolved in room-temperature, degassed CH₃CN solution. The transient absorption measurement was also performed for Ru complex 2. Under the same experimental conditions, a transient absorbance of ~1/10 that of Os complex 3 with similar spectral features (λₘₐₓ ~ 520 nm, not shown here) was obtained for Ru complex 2. Upon cooling to 77 K, a weak, structural phosphorescence maximized at ~460 nm (τ₁ ~ 50 μs) was resolved, and it clearly possesses a pypz 3π−π* character, confirming the deactivation of the initial 1π−π* excited states to the lowest 3π−π* triplet state via intersystem crossing. However, as the ligand field strength associated with the second-row Ru(II) atom is much weaker than that of the third-row Os(II) analogues, the d–d levels of the Ru complexes would be significantly lowered and become thermally accessible from the 2π−π* level, as the latter is somewhat independent of the variation of the central metal atom. In such cases, the 3LC (π−π*) state is expected to internally relax to the d–d states through a thermally activated process, followed by a rapid, dominant radiationless pathway to give negligible room-temperature phosphorescence. In contrast, due to the greater ligand field strength, the d–d levels in Os complexes are thermally inaccessible from the 3LC (π−π*) state, so the competing radiationless deactivation no longer plays a major role in quenching the 3LC (π−π*) state, resulting in a strong intra-ligand 3π−π* phosphorescence in the room-temperature solution phase.

Discussion

Based on the above results, the photophysical properties of these new series of [MC(O)₂(pypz)₂] complexes may be rationalized through their structural uniqueness. The strong π-accepting ancillary CO ligands are capable of removing the electron density from the cationic metal center and hence lead to a higher oxidation potential for the current M(II) metal center as well as the destabilization of the M(III) metal center. This electron-withdrawing characteristic, being much greater than those observed for the phosphine and arsine chelates, would substantially increase the relative energy of the typical low-lying metal-to-ligand (Os(dz)−(π*) charge transfer transition and make the pypz ligand-centered π−π* energy level the lowest singlet excited state. Evidence has been found from the electronic absorption spectroscopy, in which the MLCT transitions were obscured and possibly hidden inside the strong ligand-centered π−π* transition. It should be noted that the relatively different ligand-versus-CO orientation, to a certain extent, seems to influence the spectral properties as well. For example, Os(II) complex 3 shows a bathochromic shift of the absorption spectrum and relatively higher peak extinction coefficient (see Table 4) than that of Ru complex 2, consistent with different extents of the N(pz)−metalative interaction. Upon electronic excitation, the strong spin–orbit coupling expected for Os(II) complexes 3–5 would lead to an efficient intersystem crossing from the singlet excited state to the triplet manifold of the ligands. The

fast rate of intersystem crossing is supported by the lack of the 2-pyridyl pyrazolate fluorescence in the region of 370 nm, as well as the system-response-limit rise time for the phosphorescence (<200 ps). The strong spin–orbit coupling, to a great extent, also removes the spin-forbidden nature of the \( T_1 \rightarrow S_0 \) radiative relaxation. As a result, a dominant blue-green phosphorescent emission can be clearly observed in the room-temperature solution phase. For complex 3, the phosphorescence yield (\( \Phi_p \)) was measured to be 0.14 in the degassed CH\(_3\)CN solvent. Assuming unitary efficiency for the intersystem crossing and neglecting the emission loss due to the triplet–triplet annihilation, the \( T_1 \rightarrow S_0 \) radiative decay rate constant \( k_r \) can be derived from a relationship expressed as \( k_r = \Phi_p k_{\text{obs}} \), where \( k_{\text{obs}} \) was measured to be \( 5.46 \times 10^4 \) s\(^{-1}\). Accordingly, a \( k_r \) value of \( 7.64 \times 10^3 \) s\(^{-1}\) was deduced. In comparison to the \(<1.0 \text{ s}^{-1}\) phosphorescence radiative decay rate of the corresponding free ligand anion (i.e., pypz\(^{-}\) anion), the vast enhancement of the \( T_1 \rightarrow S_0 \) radiative decay rate through the spin–orbit coupling in complexes 3–5 is supportive. On the other hand, as indicated by the lack of fluorescence and small \( T_1 \rightarrow T_n \) transient absorbance, the rapid excited-state relaxation in complex 2 seems to be dominated by two additional factors. First, the small difference in energy between the \( 3\pi\pi^* \) and the metal-centered \( d-d \) states would promote facile thermal deactivation through population of the \( d-d \)-states, followed by radiationless, vibrational relaxation. Second, the enhanced nonradiative decay observed in complex 2 may also be rationalized via a ligand orientation effect. This is evidenced by the fact that the spatial separation between the cis pyridyl fragments in complex 2, estimated by the center-to-center distance between two pyridyl ring systems, is approximately 4.76 Å. This would then promote the inter-ligand electron transfer and give a fast nonradiative decay process. In contrast, complex 3 shows pyridyl sites located at the trans disposition and having a much longer intramolecular spatial separation of 6.9 Å, so that the inter-ligand through-space interaction is less effective than that expected in complex 2. Further support of this viewpoint is given by a recent study on the luminescent properties of a series of tris-cyclometalated Ir(III) complexes, which have shown an approximate 10-fold reduction of the emission quantum efficiency for the meridional isomers with respect to the alternative facial isomers.\(^{(25)}\) It is suggested that the weaker Ir–C bonds observed in the meridional isomers would provide a much greater bond distortion in the excited state, and this could be the more likely reason for the effective quenching of the phosphorescent emission.

4. Conclusion

In conclusion, we report the syntheses and characterization of a new family of thermal and photochemical robust Os(II) and Ru(II) metal complexes. While the respective boron complex 1 exhibits strong fluorescent emission derived from the singlet \( \pi-\pi^* \) chromophore, our results clearly demonstrate for the first time that Os(II) complexes 3–5 are capable of displaying a high-energy ligand-centered phosphorescence in the room-temperature solution phase, despite the fact that Os(II) complexes typically exhibit lower energy MLCT emission in the regions of green, red, and even infrared. The presence of strong \( \pi \)-accepting ancillary CO ligands as well as the Os(II) heavy atom effect makes the as-synthesized pyridyl pyrazolate complexes unique in their photophysical properties. Taking into account knowledge that current interest in the blue- and red-color emitter has recently been focusing on the Ir(III) type of \( \pi-\pi^* \) or MLCT phosphorescence, future applications of the title Os(II) complexes and/or relevant derivatives may be motivated by the strategy of using these complexes in the fabrication of organic light-emitting diodes (OLEDs). The strong metal–ligand bonding interaction and high phosphorescence quantum efficiencies with ca. microsecond lifetimes may improve OLED performance. Focus on this approach is currently in progress.

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Supporting Information Available: X-ray crystallographic file (CIF and PDF) for complexes 2 and 3. This material is available free of charge via the Internet at http://pubs.acs.org.

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