

ABSTRACT: The structure–property relationship study of a series of cationic Ir(III) complexes in the form of [Ir(C^N)2(N^N)]+ [where dBuppy = 4,4′-di-tert-butyl-2,2′-bipyridine and C^N = cyclometallating ligand bearing an electron-removing group (EWG)] at C4 of the phenyl substituent, i.e., –CF3 (1), –OCF3 (2), –SCF3 (3), –SO2CF3 (4)] has been investigated. The physical and optoelectronic properties of the four complexes were comprehensively characterized, including by X-ray diffraction analysis. All the complexes exhibit quasi-reversible dBuppy-based reductions from −1.29 to −1.34 V (vs SCE). The oxidation processes are likewise quasi-reversible (metal + C^N ligand) and are between 1.54 and 1.72 V (vs SCE). The relative oxidation potentials follow a general trend associated with the Hammett parameter (σ) of the EWGs. Surprisingly, complex 4 bearing the strongest EWG does not adhere to the expected Hammett behavior and was found to exhibit red-shifted absorption and emission maxima. Nevertheless, the concept of introducing EWGs was found to be generally useful in blue-shifting the emission maxima of the complexes (λem = 484–545 nm) compared to that of the prototype complex [Ir(ppy)2(dBuppy)]PF6 (where ppy = 2-phenylpyridine) (λem = 591 nm). The complexes were found to be bright emitters at solution concentration at room temperature (ΦPL = 45–66%) with microsecond excited-state lifetimes (τe = 1.14–4.28 μs). The photophysical properties along with density functional theory (DFT) calculations suggest that the emission of these complexes originates from mixed contributions from ligand-centered (LC) transitions and mixed metal-to-ligand and ligand-to-ligand charge transfer (LLCT/MLCT) transitions, depending on the EWG. In complexes 1, 3, and 4 the 3LC character is prominent over the mixed 3CT character, while in complex 2, the mixed 3CT character is much more pronounced, as demonstrated by DFT calculations and the observed positive solvatochromism effect. Due to the quasi-reversible nature of the oxidation and reduction waves, fabrication of light-emitting electrochemical cells (LEECs) using these complexes as emitters was possible with the LEECs showing moderate efficiencies.

INTRODUCTION

Over the past few decades, heteroleptic cationic Ir(III) complexes have garnered widespread interest due to their frequently bright phosphorescence that can be tuned across the visible spectrum through simple ligand modification. Due to facile color tuning, high photoluminescence quantum yield (ΦPL) and short emission lifetimes (τe), iridium complexes are ideal emissive materials for electroluminescent (EL) devices and remain the most popular materials for use in organic light-emitting diodes (OLEDs) and in light-emitting electrochemical cells (LEECs). For lighting and display applications, bright and stable red, green, and blue emitters are all required. Whereas the performance of phosphorescent red and green organometallic emitters is satisfactory, there is presently a dearth of bright and stable blue emitters for OLEDs and a near complete absence of blue emitters for LEECs.

Generally, two strategies are adopted to blue-shift the luminescence of cationic cyclometalated Ir(III) complexes of the form [Ir(C^N)2(N^N)]+: (a) increase the energy of the emissive metal-to-ligand, intraligand or ligand-to-ligand charge transfer state (MLCT, ILCT or LLCT, respectively) by...
introducing electron-donating groups (EDGs) on the ancillary N^N ligand or electron-withdrawing groups (EWGs) on the cyclometallating C^N ligands; and (b) increase the energy of the emissive π−π* ligand-centered states by limiting conjugation within the ligand. With EWGs positioned on the C^N ligands, the HOMO is stabilized more than the LUMO, which translates to an increased band gap and, generally, bluer emission. Currently, it is a near-universal strategy to employ C^N ligands like 2-(4,6-difluorophenyl)-pyridine, dFppy, to obtain blue/blue-green-emitting Ir complexes in solution, a subset of which have been used in LEECs. While the Ir(III) complexes containing fluorinated C^N ligands exhibit sky-blue to blue emission, they are prone to degradation via defluorination. Thus, while the presence of fluorne atoms on the C^N ligands blue-shifts the emission, its deleterious effect on device lifetime mitigates against its inclusion in the emitter design. Apart from fluorination, the other EWGs that have been used to blue-shift the phosphorescence of Ir(III) complexes include the following: sulfonyl (−SO2R), triluoromethyl (−CF3), pentafluoro-λ5-sulfanyl (SF5), and cyclometalated heterocycles such as 2,3′-bipyridinato. Sky-blue and deep-blue emitting cationic Ir(III) complexes bearing biimidazole, bis(NHC), substituted triazole or tetrazole, or pyrazolyl-pyridine as ancillary ligands have also been explored, but challenges still remain regarding efficiencies and stabilities of these emitters in devices, primarily as a consequence of thermal population of metal-centered states (3MC). Thus, there is still a demand for blue-emitting phosphors as emitters in lighting devices.

Among various other EWGs, those that find interest in organometallic chemistry are fluorocarbon groups, given the fact that these groups are bulky, polar, hydrophobic, and chemically inert. They are therefore attractive for reducing interactions between iridium complexes, which could otherwise lead to aggregation-induced broadening and red-shifting of the phosphorescence. This is particularly important in blue electroluminescent devices and in particular LEECs as the light-emitting layer is often a neat film. Tuning of optoelectronic properties in organometallic Ir(III) complexes using bulky fluorocarbon groups is also relatively rare compared to those with other transition metal ions. Thus, stable, bulky, and electron-withdrawing fluorocarbon groups are desirable alternatives to the commonly used C(aryl)−F motif and should serve the similar purpose of fine-tuning of optoelectronic properties of their Ir(III) complexes.

Herein we report a systematic and comparative synthetic, structural, electrochemical, and spectroscopic study of phosphorescent Ir(III) complexes with a series of electron-
withdrawn fluorocarbon groups attached at the 4-position of the C\(^N\) ligands (ligands L1−L4 in Chart 1), including the following: trifluoromethyl (−CF\(_3\)), trifluoromethoxy (−OCF\(_3\)), trifluoromethylsulfonyl (−SCF\(_3\)), and trifluoromethylsulfonyl (−SO\(_2\)CF\(_3\)). These C\(^N\) ligands were used in combination with the electron-donating 4,4\(^\prime\)-di-tert-butyl-2,2\(^\prime\)-bipyridine (dtBubpy) as the ancillary N\(^N\) ligand (complexes 1−4 in Chart 1). The effects of the presence of different EWGs on the optoelectronic properties in conjunction with density functional theory (DFT) calculations are discussed, and the results are compared with six benchmark complexes (complexes R1−R6 in Chart 2) along with the application of the new complexes in LEECs as solid-state lighting devices.

**RESULTS AND DISCUSSION**

**Synthesis.** The syntheses of the C\(^N\) ligands L1−L4 and the heteroleptic iridium complexes 1−4 are shown in Scheme 1. Each of the substituted C\(^N\) ligands was prepared in high yield via Stille palladium-catalyzed cross-coupling reaction in moderate to good yield.\(^5\) Ligands L1 and L2 were synthesized following literature procedure.\(^5\) The tin byproducts were removed by passing the reaction mixture through silica gel and (10 wt %) potassium carbonate.\(^5\) The C\(^N\) ligands, L1−L4, dimers D-L1−D-L4, and cationic complexes 1−4 were characterized by 1H and 19F NMR spectroscopy (see Figures S1−S6 for stacked NMR spectra and Figures S7−S37 for individual NMR spectra), HRMS, melting point determination, and elemental analyses. Complexes 1−4 are air- and moisture-stable solids that are soluble in polar organic solvents including acetonitrile and dichloromethane. The ligands L1−L4, dimers D-L1−D-L4, and cationic complexes 1−4 were characterized by \(^1\)H and \(^19\)F NMR spectroscopy (see Figures S1−S6 for stacked NMR spectra and Figures S7−S37 for individual NMR spectra), HRMS, melting point determination, and elemental analyses. Complexes 1−4 were further characterized by single crystal X-ray structural analysis. Inherent C\(_2\) symmetry in the solution state of complexes 1−4 was confirmed by both \(^1\)H and \(^19\)F NMR

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**Scheme 1. Synthesis of C\(^N\) Ligands, L1−L4, and [Ir(C\(^N\))\(_2\)(dtBubpy)]PF\(_6\) Complexes, 1−4**

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**Figure 1.** Solid-state structures of complexes 1−4. Hydrogen atoms, PF\(_6^-\) counterions, solvent molecules, minor components of disordered molecules, and additional independent molecules are omitted for clarity. Thermal ellipsoids correspond to a 50% probability level.
spectroscopy. The effect of the EWG on the electronics of the phenyl ring is pronounced, evidenced by the gradual and linear downfield shift of the proton ortho to the nitrogen atom of the pyridyl ring of the C^N ligands from complex 2 to 3 to 4 (Figure S5) in accordance with the increasing Hammett constant (σ_m) from −OCF3 (0.38) to −SCF3 (0.40) to −CF3 (0.43) to −SO2CF3 (0.83) as calculated by Hansch et al. (Figure S38). The 19F NMR spectra exhibit the characteristic singlet peak in the range −42 ppm to −80 ppm for a substituted −CF3 group and a doublet (J = 713 Hz at 400 MHz instrument due to ^19P−19F nuclear coupling) at ∼−73 ppm for the six magnetically equivalent fluorine atoms in the PF6 anion (Figure S6). The HRMS analyses of L2–L4 and 1–4 showed the diagnostic peaks of the protonated ligand [M + H]^+ and the cation [Ir(C^N)2(dbubbpy)]^+, respectively (Figures S39–S45).

**Crystal Structures.** Crystals of the complexes, suitable for X-ray analysis, were grown by slow diffusion of diethyl ether into moderately concentrated solutions of the complexes in dichloromethane (1 and 2) and by slow evaporation of mixed solutions of CH2Cl2/heptanes (3) or CH2Cl2/hexanes (4) (Figure 1). Selected crystallographic parameters are tabulated in Table S1. Selected bond distances and angles in comparison to those predicted by DFT calculations are summarized in Table S2. In each of the complexes, the Ir(III) ion exhibits a coordinatively saturated distorted octahedral coordination environment with the two N atoms of the C^N ligands trans to each other, similar to the solid-state structure of the archetypical complex R2. The average Ir−Cppy (2.015 Å) and Ir−Nppy (2.050 Å) bond distances in 1–4 are similar to those in complex R2 (Ir−Cppy: 2.013 Å; Ir−Nppy: 2.045 Å). In each of the four complexes, the Ir−N bond to the ancillary dbubbpy ligand [2.113(10)–2.134(9) Å] is longer than that to the C^N ligands [2.039(4)–2.061(11) Å], and the N_N−Ir−N_N^N bite angle [75.89(15)–76.8(4)^°] is narrower than that of C^N−Ir−N_C^N [80.1(3)–80.6(4)^°]. The DFT calculated bond distances and angles are in line with those observed for the solid-state structures. In all the complexes, the presence of bulky substituents in the cyclometallating ligand and the dbubbpy groups onto the backbone of the bipyridine unit prevents any face-to-face π−π stacking of the complexes.

**Electrochemical Properties.** The electrochemical properties of the ligands and mononuclear complexes have been investigated by both cyclic and differential pulse voltammetry (CV and DPV, respectively) in degassed MeCN, and the first redox potentials, reported with respect to SCE (Fc/Fc^+ = 0.38 V in MeCN), are compiled in Table 1, while the full set of redox potentials are detailed in Table S3, and the CVs along with DPVs are shown in Figure 2. Electrochemistry data of R1, R3, R5, and R6 have been corrected to account for the experimental setup and referencing versus SCE (see Electrochemistry section in the Supporting Information for full details).

At positive potential, complexes 1–4 exhibit a quasi reversible single electron oxidation between 1.54 and 1.72 V. DFT calculations using the B3LYP functional indicate that incorporation of various electron-withdrawing groups results in a stabilization of the HOMOs (HOMO = highest occupied molecular orbital) of complexes 1–4 compared to that of reference complex R1 and the HOMOs of these complexes are almost equally constituted of the metal center as well as the C^N ligands (Figure 3). Thus, and also following literature data of structurally similar cationic Ir(III) complexes, the oxidation potentials of complexes 1–4 are assigned to the removal of an electron from an admixture of the (metal +
C^N)-based orbitals. The lower energies calculated for the HOMOs of 1 (\(E_{\text{HOMO}} = -5.87 \text{ eV}\)), 2 (\(E_{\text{HOMO}} = -5.76 \text{ eV}\)), 3 (\(E_{\text{HOMO}} = -5.83 \text{ eV}\)), and 4 (\(E_{\text{HOMO}} = -6.13 \text{ eV}\)) compared to that of R1 (\(E_{\text{HOMO}} = -5.56 \text{ eV}\)) are in good agreement with the higher anodic potentials measured for complexes 1–4 in comparison to that of R1 (Table 1). The \(-\text{SO}_2\text{CF}_3\) substituted ppy moiety (L4) in complex 4 acts as the strongest electron-withdrawing group (EWG) compared to \(-\text{CF}_3\), \(-\text{OCF}_3\), and \(-\text{SCF}_3\), as demonstrated by the highest oxidation potential of this complex, while the \(-\text{OCF}_3\) substituted ppy moiety (L2) acts as the poorest EWG. This fact is in line with the increasing energy of the HOMO from complexes 4 to 1 to 3 to 2 as calculated by DFT and also in good agreement with the higher Hammett meta-constant (\(\sigma_m\)) of the \(-\text{SO}_2\text{CF}_3\) group (0.83) compared to \(-\text{CF}_3\) (0.43), \(-\text{SCF}_3\) (0.40), and \(-\text{OCF}_3\) (0.38) groups.53 A linear relationship (\(R^2 = 0.93\)) was found between the Hammett constants of the different EWGs of 1–4, those found on the reference complexes, and the oxidation potentials of the corresponding complexes (Figure 4).

Upon scanning to negative potential, several ligand-based reductions are exhibited by complexes 1–4. While for complexes 1, 3, and 4, the first reductions are mono/electronic, for complex 2 this reduction is found to be dielectronic as inferred from the DPVs of these complexes (Figure 2). The reduction profile of 4 was found to be irreversible while the reduction for 1–3 was quasi reversible in nature, thus suggesting an electrochemical instability of 4. DFT calculations point to a LUMO (LUMO = lowest unoccupied molecular orbital) that has predominantly dtypy character. Therefore, the first reduction can unambiguously be assigned to reduction of the dtypy moiety, while LUMO + 1 and LUMO + 2 are localized mainly on the L1–L4 ligands. Thus, in a coarse approximation, the second and third reduction waves may be assigned to reduction of the L1–L4 moieties. Unlike the other EWGs found in 1–3, the strongest \(-\text{SO}_2\text{CF}_3\) EWG group in 4 is prone to undergo irreversible reduction. This second irreversible reduction of 4, presumably localized on the \(-\text{SO}_2\text{CF}_3\) group of L4 as suggested by DFT calculation, occurs very closely to the first quasi reversible reduction, thus contributing to the overall irreversible nature of the reduction profile of complex 4. Such behavior had previously been observed for \text{SF}_5\)-containing cationic iridium complexes.26 The trend in the redox gap (\(\Delta E_{\text{redox}}\)) of complexes 1–4, along with that of R1, satisfactorily matches the trend in the calculated HOMO–LUMO energy differences (Table 1). The EWGs found on the C^N ligands stabilize both the HOMO and the LUMO though the former is affected to a much more appreciable degree than the latter. Thus, the electronics of 1–4 are essentially HOMO-modulated (Figure 3). Due to the strong electron-withdrawing nature of the \(-\text{SO}_2\text{CF}_3\) group, the
first reduction of 4 is anodically shifted by between 20 and 50 mV compared to those of 1–3. The reduction potential of 1, where the −CF₃ substituent is positioned meta to the Ir−Cₚₚₚ bond, is found to be anodically shifted by ∼30 mV compared to those of R₃, where the −CF₃ substituent is positioned para to the Ir−Cₚₚₚ bond. In a comparison of both the oxidation and reduction potentials, the result demonstrates that 1 is harder to oxidize, but easier to reduce compared to R₃. This fact does not follow the trend of Hammett constants (σ) of the electron-withdrawing −CF₃ substituent when regiospecifically positioned (σₘₐₜ = 0.43, σₚₐʳᵃ = 0.54).⁴²

Photophysical Properties. The UV–vis absorption properties of complexes 1–4 have been investigated in MeCN solution at room temperature and the respective spectra are shown in Figure 5; the data are summarized in Table 2 and Table S4. Overlays of each of the experimentally observed UV–vis absorption spectra of the complexes with their predicted transitions at different wavelengths as obtained by singlet TD-DFT calculations are shown in Figure S46. The absorption spectra of all the complexes are characterized by two intense bands between 250 and 300 nm and several lower-intensity bands beyond 300 nm. Predominant spin-allowed singlet 1π → π* ligand-centered (1 LC) transitions centered both on cyclometallating and the ancillary ligands, as predicted by TD-DFT calculations of complexes 1–4 (Tables S5–S8), are observed at the highest-energy band, whereas the other intense peaks at ∼290 nm result mainly from 1LC transitions and along with minor amounts of 1MLCT transitions. Between 300 and 400 nm the electronic transitions constitute 1LC and 1MLCT transitions, with varying contributions of 1MLCT and singlet ligand-to-ligand transition (1LLCT) from one complex to another. For complexes 1, 3, and 4 the electronic transitions between 410–436 nm are assigned as HOMO → LUMO transitions, whereas for complex 2 the similar peak at 400 nm is assigned as a HOMO → LUMO + 1 transition (Tables S5–S8). For 2, the lowest-energy HOMO → LUMO transition is

Figure 5. UV–vis spectra of complexes 1, 2, 3, and 4 recorded in MeCN at 298 K (inset shows magnified spectra from 400 to 525 nm).

Table 2. Relevant Photophysical Data for Complexes 1–4

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<th>compd</th>
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<th>λ_em/nm</th>
<th>τ_e/µs</th>
<th>Φ$_{PL}$/%</th>
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<th>10⁻³ × k_nr/s⁻¹</th>
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<td>DCM</td>
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<tr>
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<td>543</td>
<td>4.28</td>
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<td>R2</td>
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<td>602</td>
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<td>R6</td>
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<td>496</td>
<td>498</td>
<td>2.0</td>
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*In degassed MeCN at room temperature. Steady-state emission spectra were also recorded in degassed DCM. Steady-state emission (in MeCN): λ_em = 360 nm. Time-resolved emission (in MeCN): λ_em = 378 nm. Solution Φ_PL values were measured using quinine sulfate as the external reference (λ_em = 450 nm in MeCN, Φ_c = 54.6% in 0.5 M H₂SO₄ as found in ref 66). From ref 8. Photophysical data in degassed DCM solution (Φ_PL was determined using quinine hemisulfate salt monohydrate (Φ_ref = 54.6% in 0.5 M H₂SO₄) as standard) from ref 58. Photophysical data in degassed MeCN solution (Φ_PL was determined using [Ir(ppy)$_{(2)}$(bpy)]Cl (Φ_ref = 6.22%) as standard) from ref 60. From ref 26.
emission spectra of the EWGs. (band (at 360 nm) in degassed acetonitrile. Emission maxima \(\lambda_{\text{em}}\) found in the literature. This spectral feature is predicted by TD-DFT calculations, and these bands are assigned to a mixture of spin-allowed and spin-forbidden charge transfer transitions (1MLCT, 3MLCT, and 1LLCT) due to strong spin–orbit coupling of the Ir-metal center. Although a blue-shift in the lowest-energy absorption band is expected for complex 4 in the presence of the strongly electron-withdrawing \(-\text{SO}_2\text{CF}_3\) group compared to those of complexes 1, 2, and 3, surprisingly, a strong bathochromic shift is observed for this complex. Contrary to the DFT calculated HOMO–LUMO gap for 1–4 (Figure 3), the lowest-energy absorption maxima of these complexes are red-shifted compared to that of R1, which may suggest an additional stabilization of the LUMOs of the complexes 1–4 compared to the energy of the LUMO of complex R1 due to the strong electron-withdrawing nature of the EWGs.

Figure 6 illustrates the normalized room temperature emission spectra of 1–4 upon photoexcitation into the CT band (at 360 nm) in degassed acetonitrile. Emission maxima \(\lambda_{\text{em}}\), excited-state lifetime \(\tau_{\text{e}}\), and photoluminescence quantum yield \(\Phi_{\text{PL}}\) values along with the low-energy absorption maxima of 1–4 are summarized in Table 2. In MeCN solution, sky-blue to blue-green emission with maxima ranging from 484 to 545 nm is observed for 1–4 (Figure 6a). The emission intensity increases upon degassing with nitrogen, which is a hallmark of phosphorescence. In degassed dilute MeCN solution, 1, 3, and 4 exhibit structured emission that is characteristic of emission origination from a \(^3\)LC state while 2 displays broad and unstructured emission, typical of mixed \(^3\)CT emission. Spin-unrestricted DFT calculations point to a spin density that is more localized on the C\(^{\alpha}\)N ligands and the central Ir(III) ion for 1, 3, and 4 while it is distributed to some extent to the diBupy ligand for 2 (Figure 7). These predictions are consistent with the presence of vibronic structure in both the phosphorescence and low-energy absorbance spectra and relatively long radiative lifetimes present in 1, 3, and 4; calculations likewise predict the mixed CT character of the excited state found in 2.  

Unexpectedly, 4, with the strongest \(-\text{SO}_2\text{CF}_3\) EWG group, \(\sigma_m = 0.83\) exhibits the most red-shifted emission maximum in MeCN whereas 1 with the \(-\text{CF}_3\) \(\sigma_m = 0.43\) exhibits the most blue-shifted emission in MeCN. With the exception of 2, concomitant to the red-shift in the low-energy absorption maxima from 1 to 3 to 4, the emission maxima are also red-shifted (Table 2). The predicted emission maxima, \(E_{\text{AE}} = E(T_1) - E(S_0)\), at the \(T_1\) optimized geometries (adiabatic electronic emission) obtained by DFT calculations for complexes 1, 3, and 4 are at 541, 544, and 588 nm and match closely those observed experimentally and also fall in agreement with the observed trend of red-shifted emission maxima from complex 1 to 3 to 4. By contrast, for complex 2 the predicted emission maximum at 494 nm is blue-shifted. (The predicted emission maxima were calculated with relative errors of 8%, 6%, 7%, and 10% for 1, 2, 3, and 4, respectively, using the equation Error = \| \lambda_{\text{em}}(298 \text{ K}) - E_{\text{AE}}/\lambda_{\text{em}}(298 \text{ K}) \| \) in eV.) The emission profiles of the complexes in degassed DCM essentially mirror those in degassed MeCN with similar emission energies (Figure 6b and Table 2).

Incorporation of electron-withdrawing substituents on the C\(^{\alpha}\)N ligands promotes the expected stabilization of the frontier molecular orbitals (Figure 3) and the blue-shift in the emission observed for complexes 1–4 compared to the reference
complexes R1 and R2 (Table 2). The observed blue-shift in emission maxima (either in MeCN or DCM) of reference complexes R3, R5, and R6 compared to those of 1–3 are in line with the higher Hammett parameter of the regiospecifically positioned EWGs (see Table 1 for Hammett parameters of different EWGs). The F atom in R4 acts as a moderate EWG (m-F, $\sigma_m = 0.34$), and thus, the emission maximum of R4 is red-shifted compared to those of 1–3. While the trend in $\sigma_m$ correlates well with complexes 1–3, this paradigm does not fit with the red-shifted emission maximum of 4 ($m$-$SO_2$CF$_3$, $\sigma_m = 0.83$) compared to the emission maxima observed for R5 and R6.

All the complexes are found to be bright emitters with high $\Phi_{pl}$ values in the range 45–66%. Time-resolved phosphorescence measurements were performed, and the decays were found to be monoexponential, indicating the presence of a single emissive species (Figure S47). The results are shown in Table 2, and the phosphorescence lifetimes are in the range 1.14–4.28 $\mu$s. The observed higher $r_e$ values for 3 and 4 may be attributed to the presence of increased steric shielding around the iridium in these two complexes that inhibits nonradiative intermolecular charge recombination. The calculated radiative, $k_r$, and nonradiative, $k_{nr}$, decay constants, where $k_e = \Phi_{pl}/r_e$ and $k_{nr} = (k_e/\Phi_{pl}) - k_e$, are shown in Table 2 and fall in the ranges (1.28–4.38) $\times 10^4$ and (1.06–4.39) $\times 10^5$ s$^{-1}$. Among 1, 3, and 4 that exhibit greater LC emission, compounds 3 and 4 have lower $k_r$ and much lower $k_{nr}$ than compound 1, leading to higher $\Phi_{pl}$ values for 3 and 4 that are supported by the decrease in nonradiative decay by about 3 times compared to that of 1. Both the radiative and nonradiative decay rates are counterbalanced in the case of complex 2, which has more CT character in its emission.

**Light-Emitting Electrochemical Cells (LEECs).** Single-layer LEECs were fabricated with complexes 1–4 acting as the emitters. The device architecture consisted of an indium tin oxide (ITO) semitransparent anode on which a thin layer of poly[(3,4-ethylenedioxythiophene):polystyrenesulfonate, PEDOT:PSS, was spin-coated in order to facilitate charge injection into the emissive layer. The emissive layer consisted of a 4-to-1 molar ratio of the complex to the ionic liquid (IL) 1-butyl-3-methylimidazolium hexafluorophosphat [Bmim][PF$_6$]. The small amount of IL was added to shorten the turn-on time of the LEEC by increasing the concentration of ionic species and the ionic mobility in the active layer.71 The time-dependence of the luminance and average voltage of the LEECs prepared with complexes 1–4 (for clarity, denoted as LEECs 1–4) as well as the electroluminescence spectra (EL) were evaluated using a pulsed-current driving mode. Pulsed-current compared to fixed current/voltage driving leads to devices with longer lifetimes and faster turn-on times ($t_{on}$).71

Figure 8 shows the luminance and operating voltage versus time of LEECs 1–4 driven at an average current density of 50 (Figure 8a) or 100 A m$^{-2}$ (Figure 8b). The luminance in all LEECs shows the typical behavior under pulsed-current driving: after biasing the device, the luminance increases, and once the maximum luminance is reached, the luminance starts to decay. However, the typical operating voltage behavior (a fast drop toward a steady-state low voltage level) was not observed in all LEECs. At 50 A m$^{-2}$, the operating voltage of LEECs 1–3 presents a fast decrease and reaches a steady-state

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**Table 3. Key Parameters of LEECs 1–4 under Block-Wave Pulsed-Current Driving Mode (Frequency, 1 kHz; Duty Cycle, 50%) at 50 A m$^{-2}$ and 100 A m$^{-2}$**

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<th>Lum$^{an}$/cd m$^{-2}$</th>
<th>Lum$^{b}$/cd m$^{-2}$</th>
<th>$t_{on}$/s</th>
<th>$t_{on}$/min</th>
<th>$t_{off}$/min</th>
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<th>EQE/%</th>
<th>PCE$/\text{lm}$ W$^{-1}$</th>
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**Figure 8. Luminance (solid lines) and average voltage (open symbols) vs time for LEECs 1–4. Block-wave pulsed current driving mode (frequency, 1 kHz; duty cycle, 50%): (a) 50 A m$^{-2}$; (b) 100 A m$^{-2}$.
value at 2.8–3.0 V, indicating no charge injection barrier. However, higher steady-state voltage values are required for LEEC 1 at a current density of 100 A m$^{-2}$ and for LEEC 4 at a current density of 50 A m$^{-2}$ that denotes an issue of a charge injection barrier during operation in these devices. Moreover, LEEC 4 at 100 A m$^{-2}$ has a continuous increase of the operating voltage. An increase in driving voltage is generally associated with device degradation, yet the particular origin of the degradation mechanism is not easily identified. One possible device degradation mechanism can be related to the chemical degradation of the complex. The presence of an electrochemically unstable group in the complex could lead the complex to be electrochemically degraded under LEEC operation. Complex 4 has the strongest EWG group of the series (−SO$_2$CF$_3$), and it is the only complex of the series that has an irreversible wave in the reduction process detected in the cyclic voltammetry (cf. Figure 2). This irreversibility in the reduction process produces an unbalanced amount of charges during LEEC operation causing poor LEEC performance. A similarly poor LEEC performance with lack of electroluminescence was previously reported by us when cationic iridium complexes incorporating strongly electron-withdrawing pentafluorosulfonyl groups (−SF$_5$) on the cyclometalating ligand were used in the emissive layer. These groups were also found to be electrochemically unstable.

For clarity, the device comparisons will be done at 50 A m$^{-2}$, but similar conclusions can be extracted from the data of the respective LEECs driven at 100 A m$^{-2}$. LEEC performance is summarized in Table 3. LEECs 1–3 only require a short turn-on time ($t_{on}$), here defined as the time needed to reach a luminance of 100 cd m$^{-2}$. In all cases $t_{on}$ was determined to be faster than 20 s, and the LEECs reached their maximum luminance after a few minutes. For LEEC 4 $t_{on}$ could not be determined because of the low luminance levels achieved. The maximum luminance reached was 437, 564, 215, and 5 cd m$^{-2}$ for LEECs 1, 2, 3, and 4, respectively. The trend in device lifetime ($t_{1/2}$), defined as the time to reach one-half of the maximum luminance, mirrors that observed for maximum $t_{1/2}$ of 228 min for LEEC 1, 54 and 16 min for LEECs 2 and 3, respectively, and 6 min for LEEC 4.

The current efficiency, external quantum efficiency (EQE), and power conversion efficiency (PCE) were also analyzed for the LEECs. LEEC 1 has the highest values of current efficiency, EQE, and PCE of the series with 8.9 cd A$^{-1}$, 2.7%, and 4.4 lm W$^{-1}$, respectively. However, the current efficiencies are lower compared to the most efficient green-emitting LEEC ($\lambda_{PL(max MeCN)}$ = 512 nm, CIE = 0.299, 0.451) reported until now with 38 cd A$^{-1}$, 14.9%, and 39.8 lm W$^{-1}$ under constant voltage driving conditions. However, further comparison with green-emitter LEECs driven at constant voltage$^{72}$ cannot be considered due to the voltage dependence of the LEECs’ performance under constant voltage. LEECs 2 and 3 are less efficient than LEEC 1 with metrics of 7.2 cd A$^{-1}$, 2.3%, and 3.1 lm W$^{-1}$, and 4.3 cd A$^{-1}$, 1.4%, and 1.9 lm W$^{-1}$, respectively. LEEC 4 has the lowest values of efficiency with 0.1 cd A$^{-1}$, 0.03%, and 0.02 lm W$^{-1}$. Considering ohmic contact, thin-film photoluminescence quantum yields of 15.9%, 32.6%, 15.7%, and 20.1% for complexes 1–4, and a typical outcoupling of 20%, the theoretical maximum external quantum efficiency (EQEmax) for LEECs 1–4 is 3.2%, 6.5%, 3.1%, and 4.0%, respectively, which elucidate low radiative losses for LEEC 1 and moderate losses for LEECs 2–4. Two mechanisms cause rapid radiationless deactivation. On the one hand, the formation and continuous growth of the doped zones during the LEEC operation lead to efficient quenching of the excitons.$^{73,74}$ On the other hand, the quasireversible reduction for complexes 1–3 and the electrochemical instability of complex 4 suggest an unbalanced hole/electron carrier that leads to exciton quenching at one electrode interface.$^{75}$ As mentioned before, the efficiency of the radiative process is directly related to the photoluminescence quantum yield of the material. However, here the trend observed in the efficiency of the device performance due to the different EWGs: $\text{CF}_3 (1) > \text{OCF}_3 (2) > \text{SCF}_3 (3) > \text{SO}_2\text{CF}_3 (4)$, cannot be related to the $\Phi_{TFPL}$ values measured in the thin-film configuration (see Table 4), where all complexes show moderate $\Phi_{TFPL}$ values, with 15.9%, 32.6%, 15.7%, and 20.1% for complexes 1, 2, 3, and 4, respectively.

### Table 4. Electroluminescence (EL) Data for LEECs 1–4 and Thin-Film Photoluminescence (TFPL) Data for Complexes 1–4

<table>
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<th>CIE</th>
<th>$\lambda_{max,TFPL}$/nm</th>
<th>$\Phi_{TFPL}$/%</th>
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<td>493(sh), 524</td>
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<tr>
<td>2</td>
<td>566</td>
<td>(0.44, 0.53)</td>
<td>546</td>
<td>32.6</td>
</tr>
<tr>
<td>3</td>
<td>496(sh), 568</td>
<td>(0.44, 0.53)</td>
<td>498(sh), 561</td>
<td>15.7</td>
</tr>
<tr>
<td>4</td>
<td>566</td>
<td>(0.45, 0.53)</td>
<td>523, 545</td>
<td>20.1</td>
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</table>

$^\lambda_{max} = 320 \text{ nm}.$

It is important to highlight that there is a detrimental effect of the $\text{CF}_3$ substituent position from $para$ to the Ir–C$_{C=N}$ in R$_3$ to the meta Ir–C$_{C=N}$ in 1 at both current densities,$^{58}$ where the maximum luminance decreases from 852 cd m$^{-2}$ (R$_3$) to 427 cd m$^{-2}$ (1) and the external quantum efficiency (EQE) is reduced from 5.4% (R$_3$) to 2.7% (1), both at 50 A m$^{-2}$. LEEC 4 with the strongest EWG, $\text{SO}_2\text{CF}_3$, installed on the C$_{C=N}$ ligands presents the lowest performance of the series, which we attribute exclusively to the electrochemical instability of this group. Through this study it has become evident that the C$_{C=N}$ ligands cannot be too electron-poor as this results in greater electrochemical irreversibility leading detrimentally to significantly poorer LEEC performance.

The electroluminescence (EL) and thin-film photoluminescence (PL) spectra are shown in Figure 9. Thin-film photoluminescence (PL) spectra and thin-film $\Phi_{TFPL}$ were measured using the device composition of the emissive layer.

**Figure 9.** Electroluminescence (EL) spectra (solid line) of LEECs 1–4 and thin-film photoluminescence (PL) spectra (dashed line) of complexes 1–4.
spin-coated on a quartz plate. EL and thin-film PL maximum emission wavelengths as well as Φ<sub>TPl</sub> values are provided in Table 4. All devices present similar EL maxima at 556, 566, 568, and 566 nm for LEECs 1, 2, 3, and 4, respectively. Additionally, LEECs 1 and 3 have a high-energy shoulder at 493 nm (1) and 496 nm (3). The thin-film PL maxima for 1–3 are modestly blue-shifted at 524, 546, 561 nm, respectively. Similar to their EL spectra, the thin-film PL spectra for 1 and 3 present a shoulder at 493 nm (1) and 498 nm (3). The thin-film PL spectrum for 4 presents two maximum emission peaks at 523 and 545 nm. The structured nature of the thin-film PL spectra for 1, 3, and 4 mirrors the structured spectra observed in MeCN solution. Likewise, the broad and unstructured thin-film PL spectrum for 2 has the same form as that observed in MeCN solution. All EL maxima are red-shifted with respect to the thin-film PL maxima and with respect to the PL maxima observed in solution, but the position of the shoulder in complexes 1 and 3 remains at the same wavelength. This red-shifting of the EL and thin-film PL maxima with respect to the PL maxima in solution is due to the aggregation in the solid state.72

The similar EL and PL emission profiles and energies of 3 denote that the nature of the emission is produced from the same excited state. The absence of structured emission in the EL spectrum of 4 and its coincident emission energy and profile with that observed in the EL spectrum of 3 could imply an electrochemical reduction of the −SO₂CF₃ moiety to an −SCF₃ group.

The Commission Internationale de l’Eclairage (CIE) coordinates were determined from the electroluminescence spectra for 1–4 (Table 4). LEEC 1 presents a blue-green emission with CIE coordinates (0.37, 0.55), and LEECs 2–4 have a green emission with similar CIE coordinates (0.44, 0.53) for LEEC 2, (0.44, 0.53) for LEEC 3, and (0.45, 0.53) for LEEC 4.

In summary, four sky-blue to blue-green emitting (λ<sub>em</sub> 484–525 nm) cationic heteroleptic iridium(III) complexes bearing electron-withdrawing fluorocarbon ligands have been synthesized, and their optoelectronic properties were investigated. The complexes exhibit quasi-reversible first oxidation and reduction peaks, thereby rendering them suitable as emitters in LEECs. Surprisingly, despite containing stronger EWGs, the lowest-energy absorption maxima of 1–4 are more red-shifted compared to the reference complexes R1 and R2. While the trends observed in the absorption spectra of 1–4 are not in line with the predicted trend obtained from singlet TD-DFT calculations, the triplet spin density calculations are in agreement with the trend observed for the emission behavior, except for 2. However, 1–4 were shown to exhibit blue-shifted emission maxima compared to those of R1 and R2. Thus, this study demonstrates that the common design paradigm of achieving bluer emission upon introduction of increasingly stronger EWG may not always be applicable. Successful applications of these complexes in LEECs have been achieved, albeit with moderate external quantum efficiencies.

EXPERIMENTAL SECTION

General Synthetic Procedures. Commercial chemicals were used as supplied. All reactions were performed using standard Schlenk techniques under inert (N₂) atmosphere with reagent grade solvents. Flash column chromatography was performed using silica gel (Silia-P from Silicycle, 60 Å, 40–63 μm). Analytical thin layer chromatography (TLC) was performed with silica plates with aluminum backings (250 μm with indicator F–254). Compounds were visualized under UV light. H for (ligands and dimers), 13C, and 19F NMR spectra were recorded on a Bruker Avance spectrometer at 400, 125, and 376 MHz, respectively. 1H NMR spectra for charged complexes were recorded on a Bruker Avance spectrometer at 500 MHz. The following abbreviations have been used for multiplicity assignments: s for singlet, d for doublet, t for triplet, m for multiplet, and br for broad. Deuterated dimethyl sulfoxide (DMSO-δ_6) and deuterated dichloromethane (CDCl₃) were used as the solvents of record. 1H NMR spectra were referenced to the solvent peak. Melting points (MP’s) were recorded using open-ended capillaries on an Electro-thermal melting point apparatus and are uncorrected. High-resolution mass spectra were recorded at the EPSRC UK National Mass Spectrometry Facility at Swansea University on a quadrupole time-of-flight (ESI-Q-TOF), model ABSciex 5600 Triple TOF, in positive electrospray ionization mode, and spectra were recorded using sodium formate solution as the calibrant. Elemental analyses were performed by Mr. Stephen Boyer, London Metropolitan University.

Syntheses of Ligands: L1–L4. 2-(4-(Trifluoromethyl)phenyl)pyridine (L1). The synthesis is a modification to that previously reported.73 4-Bromotrifluoromethylbenzene (1.2 g, 5.3 mmol, 1.18 equiv), 2-(tri-n-butylstannyl)pyridine (85–95%; 1.06 g, 4.5 mmol, 1 equiv), and [Pd(PPh₃)₄] (0.23 g, 0.2 mol % catalyst) were stirred in dry degassed toluene at 120 °C for 48 h to give a yellow solution. The product was purified by column chromatography twice (the reaction mixture was loaded on the column directly): first on a fine mixture of silica and K₂CO₃ (1.5 g, anhydrous) to remove tin byproducts, and then just on silica (20 g). The elution was performed with hexane/dichloromethane (2/1 to 1.5/1 v/v/δ) to afford the compound as a white crystalline solid. Yield: 0.56 g, 56%. 1H NMR (400 MHz, DMSO-δ₆) δ (ppm): 8.73 (ddd, J = 4.79, 1.80, 0.94 Hz, 1H), 8.31 (dd, J = 8.90, 0.86 Hz, 2H), 8.08 (dt, J = 8.0, 1.03 Hz, 1H), 7.92–7.98 (m, 1H), 7.86 (dd, J = 8.81, 0.76 Hz, 2H), 7.45 (ddd, J = 7.48, 4.75, 1.11 Hz, 1H). 19F{1H} NMR (376 MHz, DMSO-δ₆) δ (ppm): −61.03. The characterization matches those previously reported.73 2-(4-(Trifluoromethoxy)phenyl)pyridine (L2). 1-Bromo-4-(trifluoromethoxy)benzene (1.27 g, 5.3 mmol), 2-(tri-n-butylstannyl)pyridine (85–95%; 1.53 g, 4.2 mmol), and [Pd(PPh₃)₄] (0.21 g, 0.12 mol %) were stirred in dry degassed toluene at 120 °C for 48 h to give yellow solution. The product was purified by column chromatography twice (the reaction mixture was loaded on the column directly): first on a fine mixture of silica and K₂CO₃ (1.5 g) to remove tin byproducts, and then just on silica (20 g). The elution was performed with hexane/dichloromethane (2/1 to 1 by volume) to afford the compound as a white crystalline solid. Yield: 0.8 g, 80%. Rf: 0.28 (1:1, v/v dichloromethane/hexanes on silica). Mp: 52–54 °C. 1H NMR (400 MHz, DMSO-δ₆) δ (ppm): 8.67–8.70 (m, 1H), 8.19–8.24 (m, 2H), 8.01 (dt, J = 8.04, 1.03 Hz, 1H), 7.79–7.94 (m, 1H), 7.46–7.50 (m, 2H), 7.39 (ddd, J = 7.44, 4.79, 1.11 Hz, 1H). 13C NMR (125 MHz, DMSO-δ₆) δ (ppm): 154.56, 149.67, 148.97, 137.87, 137.44, 128.49, 123.17, 123.02, 121.19, 120.47, 119.09, and 117.05. 19F{1H} NMR (376 MHz, DMSO-δ₆) δ (ppm): −56.67. HR NMR: MS: [M + 2 H]⁺: calc 240.0631 ([C₂H₆F₃NO]⁺); found 240.0632. Anal. Calcd (C₂H₆F₃NO): C, 60.26; H, 3.37; N, 5.86%. Found: C, 60.38; H, 3.26; N, 5.88%. The characterization matches those previously reported.51 2-(4-(Trifluoromethyl)thio)phenylpyridine (L3). 4-Bromophenyl-trifluoromethyl sulfide (1.22 g, 4.719 mmol), 2-(tri-n-butylstannyl)pyridine (85–95%; 1.43 g, 3.9 mmol), and [Pd(PPh₃)₄] (0.22 g, 0.12 mol %) were stirred in dry degassed toluene (15 mL) at 120 °C for 48 h to give yellow solution. The product was purified by column chromatography twice (the reaction mixture was loaded on the column directly): first on a fine mixture of silica (15 g) and K₂CO₃ (1.5 g, anhydrous) to remove tin byproducts, and then just on silica (20 g). The elution was performed with hexane/dichloromethane (2/1 by volume) to afford the compound as a white crystalline solid. Yield: 0.76 g, 77%. Rf: 0.34 (1:1, v/v dichloromethane/hexanes on silica). Mp: 64–66 °C. 1H NMR (400 MHz, DMSO-δ₆) δ (ppm): 8.72 (ddd, J = 4.7, 1.8, 0.9 Hz, 1H), 8.27–8.21 (m, 2H), 8.06 (dt, J = 8.0, 1.0 Hz, 1H).
1H), 7.94 (td, J = 7.8, 1.8 Hz, 1H), 7.86–7.81 (m, 2H), 7.43 (ddd, J = 7.4, 4.7, 1.1 Hz, 1H). 13C NMR (125 MHz, DMSO-d6) δ (ppm): 154.42, 149.80, 141.37, 136.52, 130.85, 128.40, 127.92, 125.96, 123.59, 123.51, and 120.90. 31P 1H NMR (376 MHz, DMSO-d6) δ (ppm): -41.9. HR APCI MS: [M + H]+ calculated 256.0402 ([(C12H9F3NS)+]; found 256.0402. Anal. Calcd (C 12H8F3NS): C, 25.52; H, 1.26; N, 4.67. Found: C, 25.06; H, 1.16; N, 4.59.

2-(4-(Trifluoromethyl)sulfonyl)phenyl)pyridinato-N,C 2-(μ-chloro)diiridium(III), [Ir(L2)2(Cl)-μ-Cl] (D-L2). [Ir(COD)Cl]2 (250 mg, 0.37 mmol) and ligand L2 (375 mg, 1.57 mmol) were stirred in degassed 2-ethoxyethanol (4 mL) at 130 °C (bath temperature) for 5 h. The mixture briefly turned black on mixing and heating but became a pale red solution by the end of the reaction. The solution was cooled to room temperature. Water (4 mL) was added dropwise to give a precipitate. The solid was filtered and washed with ethanol/water (1/1 v/v) and hexane. The solid was dried under vacuum to afford the compound as a yellow-orange solid. The product was used as a reagent for the next step without further purification. Yield: 0.498 g, 91%. 1H NMR (400 MHz, CDCl3) δ (ppm): 9.19 (ddd, J = 5.8, 1.6, 0.8 Hz, 4H), 8.02 (ddd, J = 8.3, 1.4, 0.7 Hz, 4H), 7.90 (ddd, J = 9.0, 7.5, 1.6 Hz, 4H), 7.61 (dd, J = 8.1, 1.8 Hz, 4H), 7.11 (dd, J = 8.1, 1.8 Hz, 4H), 6.93 (ddd, J = 7.4, 5.8, 1.4 Hz, 4H), 6.09 (dd, J = 1.2 Hz, 4H). 31P 1H NMR (376 MHz, CDCl3) δ (ppm): -42.8.

Tetrakis[2-(4-(trifluoromethyl)phenyl)pyridinato-N,C 2-(μ-chloro)diiridium(III)], [Ir{L4}(μ-Cl)] (D-L4). IrCl3-3H2O (220 mg, 0.62 mmol; iridium(III) chloride hydrate) was dissolved in a degassed mixture of 2-ethoxyethanol (6 mL) and water (2 mL) at 70 °C (bath temperature). Ligand L4 (400 mg, 1.39 mmol) was added as a solid. The mixture was stirred at 120 °C (bath temperature) for 24 h to give a red solution. The solution was cooled to room temperature to give a red suspension. Water (2 mL) was added dropwise to give the precipitate. The solid was filtered and washed with ethanol/water (1/1 v/v) and hexane. The solid was dried under vacuum to afford the compound as an orange solid. The product was used as a reagent for the next step without further purification. Yield: 0.445 g, 90%. 1H NMR (400 MHz, CDCl3) δ (ppm): 9.21 (ddd, J = 5.8, 1.5, 0.8 Hz, 4H), 8.19 (ddd, J = 7.7, 4.7, 0.8 Hz, 4H), 8.07 (ddd, J = 9.1, 7.5, 1.6 Hz, 4H), 7.84 (ddd, J = 8.2, 1.9 Hz, 4H), 7.49 (dd, J = 8.2, 1.9 Hz, 4H), 7.11 (ddy, J = 7.4, 5.8, 1.4 Hz, 4H), 6.38 (dd, J = 1.4 Hz, 4H). 31P 1H NMR (376 MHz, CDCl3) δ (ppm): -79.4.

Syntheses of Caticonic Ir-Complexes of General Molecular Formula [Ir(CN)4(μ-Cl)][PF6]−: 1–4. Bis[2-(4-(trifluoromethyl)phenyl)pyridinato-N,C 2,2'-bipyridinato-N,C 2,2'-dipyridyl)](μ-chloro)hexafluorophosphatetetrakis[2-(4-(trifluoromethyl)phenyl)pyridinato-N,C 2-(μ-chloro)diiridium(III)], [Ir(L1)(μ-Cl)][PF6] (I). Precursor dimer complex D-L1 (200 mg, 0.15 mmol) and 4,4′-diter-butyl-2,2′- bipyridine (0.53 mmol) were stirred in a degassed mixture of dichloromethane (20 mL) and methanol (4 mL) at 40 °C (bath temperature) for 24 h to give a yellow solution. The solution was evaporated. The complex was purified by column chromatography on silica (12 g). The elution was performed with 3% of methanol in dichloromethane to remove the impurities and with 6% of methanol in dichloromethane to give a yellow eluate of the product. The fractions were evaporated to dryness. The product was redissolved in methanol (5 mL) and added to a vigorously stirred aqueous solution (30 mL of NH4PF6 (1.4 g, 8.6 mmol) over 30 min to give a precipitate of the hexafluorophosphate salt. The product was filtered and washed with water and ether/hexane (1/1 v/v). The product was dissolved in 2 mL of dichloromethane and added to 40 mL of vigorously stirred ether. The product separated first as oil, but then it crystallized to a solid. The product was filtered, washed with ether, and dried under vacuum to afford a yellow solid. Yield: 0.221 g, 70%. 1H NMR (400 MHz, CD2Cl2) δ (ppm): 8.32 (d, J = 1.7 Hz, 2H), 8.07 (d, J = 8.0 Hz, 2H), 7.91 (dd, J = 7.6, 1.5 Hz, 2H), 7.86 (d, J = 8.2 Hz, 2H), 7.82 (d, J = 5.8 Hz, 2H), 7.59 (dd, J = 5.6, 0.7 Hz, 2H), 7.47 (dd, J = 5.9, 1.9 Hz, 2H), 7.34 (dd, J = 8.2, 1.1 Hz, 2H), 7.30 (dd, J = 7.3, 5.8, 1.4 Hz, 2H), 6.46 (d, J = 0.8 Hz, 2H), 1.43 (s, 18H).

13C NMR (125 MHz, CD2Cl2) δ (ppm): 166.17, 164.60, 155.42, 150.15, 149.83, 149.00, 147.41, 138.91, 131.45, 131.21, 127.27, 125.88, 124.87, 124.72, 121.34, 120.93, 119.88, 35.69, 29.94. 31P 1H NMR (376 MHz, CDCl3) δ (ppm): -73.3 (d, J = 713 Hz, PF6−), -63.2 (s, CF3). HR NISI MS: [M − PF6−]+ (100%) calcd 905.2627.
(C4H3N4F6Ir); found 905.2601. Anal. Calcld (C42H38N4F6S2Ir): C, 48.05; H, 3.65; N, 5.34%. Found: C, 48.19; H, 3.54; N, 5.35%.

**18H)**. 13C NMR (125 MHz, CD2Cl2) (ddd, J = 713 Hz, PF6−) (100%) calcd 1033.1862 (C42H38N4O4F6S2Ir+); found 1033.1828. Anal. Calcld (C42H38N4O4F12PS2Ir): C, 42.82; H, 3.25; N, 4.76%. Found: C, 42.99; H, 3.19; N, 4.87%.

**X-ray Crystallography.** Single crystals were grown by vapor diffusion of ether into concentrated CH2Cl2 solution (1 and 2) and by slow evaporation of mixed solutions of CH2Cl2/heptanes (3) or CH2Cl2/hexanes (4). Data were collected on a Rigaku FR-X Ultra-high Brilliant Microfocus RA generator/confocal optics and a Rigaku XtaLAB P2000 system, with Mo Kα radiation (λ = 0.71073 Å). Intensity data were collected using o steps accumulating area detector images spanning at least a hemisphere of reciprocal space. All data were corrected for Lorentz polarization effects, and a multisite absorption correction was applied using CrystalClear.5 Structures were solved by Patterson methods (PATTY)7 and refined by full-matrix least-squares against F2 (SHELXL-2013).5 Non-hydrogen atoms were refined anisotropically, and hydrogen atoms were refined using a riding model. All calculations were performed using the CrystalStructure interface.

**Photophysical Measurements.** All samples were prepared in HPLC grade acetonitrile with varying concentrations in the order of micromolar. Absorption spectra were recorded at room temperature using a Shimadzu UV-1800 double beam spectrophotometer. Molar absorptivity determination was verified by linear least-squares fit of values obtained from at least four independent solutions at varying concentrations ranging from 8.62 × 10−5 to 5.48 × 10−4 M.

The sample solutions for the emission spectra were prepared in HPLC grade MeCN and degassed via three freeze–pump–thaw cycles using an in-house designed quartz cuvette. Steady-state and time-resolved emission spectra were recorded at 298 K using an Edinburgh Instruments F980. All samples for steady-state measurements were excited at 360 nm. The excited-state lifetimes of the complexes were obtained by time correlated single photon counting (TCSPC) at an excitation wavelength of 378 nm using an Edinburgh Instruments FL5980 fluorimeter using a pulsed diode laser, and PL emission was detected at the corresponding steady-state emission maximum for each complex. The PL decays were fitted with a single exponential decay function. Emission quantum yields were determined using the optically dilute method.28 A stock solution with absorbance of ca. 0.5 was prepared, and then four dilutions were prepared with dilution factors between 2 and 20 to obtain solutions with absorbances of ca. 0.095, 0.065, 0.05, and 0.018, respectively. The Beer–Lambert law was found to be linear at the concentrations of the solutions. The emission
spectra were then measured after the solutions were rigorously degassed via three freeze–pump–thaw cycles prior to spectrum acquisition. For each sample, linearity between absorption and emission intensity was verified through linear regression analysis, and additional measurements were acquired until the Pearson regression factor (R^2) for the linear fit of the data set surpassed 0.9. Individual relative quantum yield values were calculated for each solution, and the values reported represent the slope value. The equation \( \Phi_i = \Phi_0 \cdot (A_i/A_0) \cdot (I_i/I_0) \cdot (n_i/n_0)^2 \) was used to calculate the relative quantum yield of each of the sample, where \( \Phi_0 \) is the absolute quantum yield of the reference, \( A \) is the absorbance at the excitation wavelength, and \( I \) is the integrated area under the corrected emission curve. The subscripts \( s \) and \( r \) refer to the sample and reference, respectively. A solution of quinine sulfate in 0.5 M H_2SO_4 (\( \Phi_r = 54.6\% \)) was used as the external reference.

**Electrochemistry Measurements.** Cyclic voltammetry (CV) measurements were performed on an electrochemical analyzer potentiostat model 620E from CH Instruments at a sweep rate of 100 mV/s. Differential pulse voltammetry (DPV) was conducted with an increment potential of 0.004 V and a pulse amplitude, width, and period of 50 mV, 0.05 s, and 0.5 s, respectively. Solutions for CV and DPV were prepared in MeCN and degassed with MeCN-saturated nitrogen by bubbling for about 10 min prior to scanning. Tetra(n-butyl)ammoniumhexafluorophosphate (TBAPF_6; ca. 0.1 M in MeCN) was used as the supporting electrolyte. A silver wire was used as the pseudoreference electrode; a glassy carbon electrode was used for the working electrode, and a Pt wire was used as the counter electrode. The redox potentials are reported relative to a saturated calomel electrode (SCE) electrode with a ferrocene/ferrocenium (Fc/Fc^+) redox couple as an internal reference (0.38 V vs SCE).55 Experimental uncertainties are as follows: absorption maxima, \( \pm 2 \) nm; molar absorption coefficient, 10%; redox potentials, \( \pm 10 \) mV; emission maxima, \( \pm 3 \) nm; emission lifetimes, 10%; luminescence quantum yields, \( \pm 5\% \).


**Light-Emitting Electrochemical Cells.** LEECs were fabricated with complexes 1–4 in the following way. The complexes were dissolved in acetonitrile (20 mg mL^-1) with 4-to-1 molar ratio of the ionic liquid (IL) 1-butyl-3-methylimidazolium hexafluorophosphate [Bmim][PF_6] and filtered through syringe filters (0.22 \( \mu \)m pore size). Prior to the deposition of the light-emitting layers, poly(3,4-ethylenedioxythiophene):polystyrenesulfonate (PEDOT:PSS, Clevios P VP AI 4083) was deposited on prepared ITO substrates. The PEDOT:PSS layer (80 nm) was spin-coated at 1000 rpm for 60 s and annealed at 150 \( ^\circ \) C for 15 min. The light-emitting layer was applied by spin-coating the respective filtered solution containing the complex and the IL, using a spin-speed of 1000 rpm for 30 s (100 nm). This was all performed in ambient atmosphere. After the film deposition, the layers were transferred to an inert atmosphere glovebox (<0.1 ppm of O_2 and H_2O, MBraun) and annealed at 100 \( ^\circ \) C during 1 h. The top electrode (70 nm thick film of aluminum) was deposited by moving the films to a vacuum chamber integrated in the inert atmosphere glovebox. The time-dependence of the luminescence and average voltage of the LEECs prepared with complexes 1–4 (for clarity, denoted as LEECs 1–4) as well as the electroluminescence (EL) spectra were evaluated using a pulsed-current driving mode. LEECs 1–4 were operated using a block-wave pulsed-current driving mode (frequency, 1 kHz; duty cycle, 50%). The current density during the pulse was set to 100 or 200 A m^-2; the average current density applied was therefore 50 or 100 A m^-2, respectively.

**Thin-Film Photoluminescence.** The photoluminescence spectra and quantum yields of the thin films deposited on a quartz plate (1 cm^2) were measured in air with a Hamamatsu C9920-02 Absolute PL Quantum Yield Measurement System (\( \lambda_{ex} = 320 \) nm). The system is made up of an excitation light source, consisting of a xenon lamp linked to a monochromator, an integration sphere, and a multichannel spectrometer.

**Inorganic Chemistry**

**ASSOCIATED CONTENT**

* Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.inorg-chem.6b01602. The research data supporting this work can be accessed at: http://dx.doi.org/10.17630/1f19249d-1888-417c-9766-3bfddbb1ec5a.

NMR spectra for all C=N ligands, dimers, and complexes; supplementary crystal structure data for complexes 1–4; supplementary optoelectronic and DFT data (PDF)

Crystallographic details for 1–4, corresponding to CSDC 1490428–1490431 (CIF)

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Notes

The authors declare no competing financial interest.

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