[Ir(C^N)2(N^N)]⁺ emitters containing a naphthalene unit within a linker between the two cyclometallating ligands†

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The synthesis of four cyclometallated [Ir(C^N)2(N^N)][PF6] compounds in which N^N is a substituted 2,2'-bipyrordin (bpy) ligand and the naphthyl-centred ligand 2,7-bis(2-(4-(pyridin-2-yl)phenoxy)ethoxy)naphthalene provides the two cyclometallating C^N units is reported. The iridium(III) complexes have been characterized by 1H and 13C NMR spectroscopies, mass spectrometry and elemental analysis, and their electrochemical and photophysical properties are described. Comparisons are made with a model [Ir(ppy)2(N^N)][PF6] compound (Hppy = 2-phenylpyridine). The complexes containing the naphthyl-unit exhibit similar absorption spectra and excitation at 280 nm leads to an orange emission. The incorporation of the naphthalene unit does not lead to a desirable blue contribution to the emission. Density functional theory calculations were performed to investigate the geometries of the complexes in their ground and first triplet excited states, as well as the energies and compositions of the highest-occupied and lowest unoccupied molecular orbital (HOMO and LUMO) manifolds. Trends in the HOMO–LUMO gaps agree with those observed electrochemically. The energy difference between the LUMO and the lowest unoccupied MO located on the naphthyl unit (LUMO+7) is large enough to explain why there is no contribution from the naphthyl-centred triplet excited state to the phosphorescence emission. Single excited states were also investigated. Light-emitting electrochemical cells (LECs) using the [Ir(C^N)2(N^N)][PF6] and [Ir(ppy)2(N^N)][PF6] complexes in the emissive layer were made and evaluated. The presence of the naphthyl-bridge between the cyclometallating units does not significantly alter the device response.

Introduction

Light-emitting electrochemical cells (LECs) are promising electroluminescent devices for highly efficient and low-cost applications in ultrathin and flexible lighting. The charged active layer sandwiched between two electrodes is either a conjugated light-emitting polymer1 or an ionic transition-metal complex (iTMC).2 The ionic nature of the active material allows the charged species to migrate towards the electrodes when a bias is applied, forming doped zones and thus lowering the injection barrier facilitating efficient electron and hole injection.3 Due to their particular stability, extremely high efficiencies and ability to tune the emission color, iTMCs incorporating iridium(III) (Ir-iTMCs) are by far the most versatile active materials used in iTMC-LECs.4,5 Ir-iTMCs are typically of the form [Ir(C^N)2(N^N)]⁺, where C^N is a cyclometallating ligand and N^N is an N,N-chelate. Since the high spin–orbit coupling of Ir-iTMCs permits intersystem-crossing from singlet to triplet states, iridium(III)-based materials achieve spin-forbidden phosphorescence emissions approaching photoluminescence quantum yields of 100%.6,7 After Slinker et al.8 reported the first Ir-iTMC-based LEC in 2004, a broad range of stable and efficient Ir-iTMC emitters has been developed, covering the whole visible range.4–7,9 A major challenge that remains is to attain white light emission.

One approach to white-emitting LECs is to combine blue-green with red-emitting complexes in the active layer of the device or to mix three iTMCs of different colours (blue, red and orange).10–13 Other strategies (not uniquely based on...
Electrospray ionization (ESI) mass spectra were recorded on a Bruker esquire 3000plus instrument. Quantum yields in CH$_2$Cl$_2$ solution and powder were measured using a Hamamatsu absolute photoluminescence (PL) quantum yield spectrometer C11347 Quantaaurus-QY. Emission lifetimes and powder emission spectra were measured with a Hamamatsu Compact Fluorescence lifetime Spectrometer C11367 Quantaaurus-Tau, using an LED light source with $\lambda_{\text{exc}} = 280$ nm. Quantum yields and PL emission spectra in thin films were recorded using a Hamamatsu absolute quantum yield C9920. The preparation of the thin film samples consisted of deposition on a quartz plate (1 cm$^2$) of the complex with addition of the ionic liquid 1-butyl-3-methylimidazolium hexafluorophosphate [Bmim][PF$_6$]. These samples were excited using a light source with $\lambda_{\text{exc}} = 320$ nm at room temperature under ambient conditions.

2,7-Bis[2-(2-hydroxyethoxy)ethoxy]naphthalene,$^{20}$ ligands 2-5$^{1,2,3}$ and [Ir(ppy)$_2$(4)][PF$_6$]$^{23}$ were prepared by standard procedures. Silica gel 60 was bought from Fluka.

**Compound H$_2$I**

A grey suspension of NaH (232 mg, 5.80 mmol) and 2,7-bis [2-(2-hydroxyethoxy)ethoxy]naphthalene (650 mg, 1.93 mmol) in dry DMF (30 mL) was rigorously stirred for 15 min before 2-(4-fluorophenyl)pyridine (1.00 g, 5.80 mmol) was added. The reaction mixture was heated under N$_2$ at 120 °C for 24 h. After cooling to room temperature, the mixture was poured into water. A white precipitate was formed, which was separated by filtration and was washed with water. The solid was added to EtOH (300 mL) and the mixture boiled (15 min) before being filtered. The filtrate was evaporated to dryness to give H$_2$I as a white solid (924 mg, 1.44 mmol, 74.6%). M.p. 129.1 °C. $^1$H NMR (500 MHz, CDCl$_3$) $\delta$ ppm 6.85 (ddd, $J = 4.8, 1.8, 1.0$ Hz, 2H, H$_B$), 7.92 (m, 4H, H$_A$), 7.70 (ddd, $J = 8.0, 7.4, 1.8$ Hz, 2H, H$_B^6$), 7.67–7.62 (overlapping m, 4H, H$_A^{3-5}$), 7.17 (ddd, $J = 7.3, 4.8, 1.2$ Hz, 2H, H$_B^5$), 7.06–6.99 (overlapping m, 8H, H$_A^{3-5}$), 4.82–4.21 (m, 8H, H$_A^{10-12}$), 4.02–3.96 (m, 8H, H$_B^{10-12}$).

$^{13}$C NMR (126 MHz, CDCl$_3$) $\delta$ ppm 159.8 (C$_A$), 157.4 (C$_B$), 157.2 (C$^{2i}$), 149.7 (C$^{2i}$), 136.8 (C$^{2i}$), 135.9 (C$_{N8}$), 132.4 (C$_{N8}$), 129.3 (C$_{N8}$), 128.3 (C$^{A2}$), 124.6 (C$_{N4}$), 121.6 (C$_{B5}$), 120.0 (C$_{B5}$), 116.6 (C$_{N3}$), 115.0 (C$_{A3}$), 106.4 (C$_{N1}$), 70.1 (C$^6$), 70.1 (C$^3$), 67.7 (C$^4$), 67.6 (C$^5$). UV-Vis $\lambda$/nm (ε/L mol$^{-1}$ cm$^{-1}$) (CH$_2$Cl$_2$, 1.00 × 10$^{-5}$ mol L$^{-1}$) 237 (78 000), 266 (33 000), 285 (38 000), 325 (6000), 350 sh (2000). Emission (CH$_2$Cl$_2$, 9.50 × 10$^{-4}$ mol L$^{-1}$, $\lambda_{\text{exc}} = 237$ nm) $\lambda_{\text{em}} = 331, 344, 410$ nm. ESI-MS m/z 643.6 [M + H$^+$] (base peak, calc. 643.3).

Found C 74.44, H 5.91, N 4.58; C$_{40}$H$_{38}$N$_2$O$_6$ requires C 74.75, H 5.96, N 4.36%.

**Experimental**

**General**

$^1$H, $^{13}$C and $^{31}$P NMR spectra were recorded at room temperature using a Bruker Avance III-600, III-500 or III-400 NMR spectrometer. $^1$H and $^{13}$C NMR chemical shifts were referenced to residual solvent peaks with respect to $\delta$(TMS) = 0 ppm and $^{31}$P NMR chemical shifts with respect to $\delta$(85% aqueous H$_3$PO$_4$) = 0 ppm. Solution absorption and emission spectra were measured using an Agilent 8453 spectrophotometer and a Shimadzu RF-5301PC spectrofluorometer, respectively.
filtered and evaporated to dryness. The solid products were combined. The product was purified by column chromatography (silica, CH₂Cl₂ changing to CH₂Cl₂ : MeOH 100:1) and [Ir(1)[μ-Cl₂]] was isolated as a yellow solid (90.0 mg, 0.052 mmol, 33.4%). ¹H NMR (500 MHz, CD₂Cl₂) δ/ppm 9.11 (dd, J = 5.8, 0.9 Hz, 2H, H⁵⁶), 7.71–7.63 (overlapping m, 4H, H⁴⁴/N⁴⁴), 7.56 (d, J = 7.9 Hz, 2H, H³¹), 7.20 (d, J = 8.7 Hz, 2H, H²), 6.97 (dd, J = 8.8, 2.5 Hz, 2H, H²), 6.90 (d, J = 2.4 Hz, 2H, H²), 6.71 (dd, J = 7.3, 5.8, 1.4 Hz, 2H, H⁵⁵), 6.33 (dd, J = 8.5, 2.5 Hz, 2H, H⁴¹), 5.32 (d, J = 2.6 Hz, 2H, H⁶²), 4.04–3.94 (overlapping m, 4H, H⁴⁺¹⁺), 3.87 (m, 2H, H⁴⁺¹⁺), 3.81–3.73 (overlapping m, 4H, H²⁺²⁻/d⁻d⁻), 3.65 (m, 2H, H²⁺²⁻/b⁻b⁻⁻), 3.61–3.57 (overlapping m, 4H, H²⁺²⁻), ¹³C NMR (126 MHz, CD₂Cl₂) δ/ppm 168.1 (C¹), 159.4 (C²), 159.2 (C³), 157.9 (C⁴), 150.3 (C⁵), 149.3 (C⁶), 139.8 (C⁶), 138.3 (C⁷), 138.3 (C⁸), 136.4(C⁹a), 135.4 (C⁹b), 130.1 (C¹⁰), 129.3 (C¹¹), 128.8 (C¹²), 127.9 (C¹³), 125.9 (C¹⁴), 125.0 (C¹⁵), 124.7 (C¹⁶), 121.2 (C¹⁷), 119.0 (C¹⁸), 117.3 (C¹⁹), 116.2 (C²⁰), 109.4 (C²¹), 107.8 (C²²), 70.6 (C³³⁻⁻/c⁻c⁻), 69.8 (C³⁴⁻⁻/c⁻c⁻), 67.3 (C³⁵⁻⁻/d⁻d⁻), UV-Vis J/nm (e/L mol⁻¹ cm⁻¹) (CH₂Cl₂, 1.00 × 10⁻⁵ mol L⁻¹) 237 (102 000), 280 (44 000), 312 (36 000), 325 (33 000), 350 sh (17 000), 400 sh (6000). ESI-MS m/z 1141.3 [M – PF₆]⁻ (base peak, calc. 1141.4). Found C 57.75, H 4.61, N 4.56; C₉₆H₃₂F₂Ir₄N₈O₈P requires C 57.89, H 4.07, N 4.36.

[Ir(1)(MeOH)₂][PF₆]⁻

A yellow suspension of [Ir₂(1)(μ-Cl₂)] (188 mg, 0.108 mmol) and AgPF₆ (59.9 mg, 0.237 mmol) in MeOH (10 ml) were stirred at room temperature for 1.5 h. The reaction mixture was filtered over celite, washed with MeOH and evaporated to dryness. The solid products were used in the next step without further purification. 

[Ir(1)(2)][PF₆]⁻ was used in the next step without further purification.

Dalton Transactions

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The method was as for [Ir(1)]2[PF6] starting with [Ir(1) (MeOH)]2[PF6] (100 mg, 0.096 mmol), 4 (40.8 mg, 0.097 mmol) and NH4PF6 (156 mg, 0.960 mmol) in MeOH (20 mL). Column chromatography used silica eluting with CH2Cl2 changing to CH3Cl2: MeOH 0.5 and finally CH3Cl2: MeOH 100:1. [Ir(1)][PF6] was isolated as an orange solid (101 mg, 0.072 mmol, 75%). 1H NMR (500 MHz, CD2Cl2) δ/ppm 8.15–8.07 (overlapping m, 4H, H6E+/E+) 7.69 (ddd, J = 8.4, 7.5, 1.5 Hz, 2H, H1B4), 7.66 (dJ = 8.6 Hz, 1H, H4N) 7.32 (d, J = 8.1 Hz, 2H, Hd/dHd), 7.25 (d, J = 2.0 Hz, 2H, H5E), 7.01 (ddd, J = 7.3, 1.3 Hz, 2H, H5E), 6.98–6.91 (overlapping m, 6H, H2G4+N1+N3), 6.77–6.69 (overlapping m, 6H, H1A3t+G3), 6.58 (br, 4H, H2G2), 6.00 (ddd, J = 8.6, 2.5 Hz, 2H, H4E), 4.65 (d, J = 2.5 Hz, 2H, H6H), 4.08–3.96 (m, 4H, H4), 3.80–3.74 (m, 2H, H5b/b′), 3.74–3.69 (m, 2H, HDd/d), 3.69–3.63 (m, 2H, H5b/b′), 3.59–3.53 (m, 2H, HDd/d), 3.53–3.45 (m, 4H, H7), 1.40 (s, 18H, H2Bu-ring E). 13C NMR (126 MHz, CD2Cl2) δ/ppm 168.1 (C2F5), 165.3 (C2F3), 164.7 (C4F8), 160.0 (C4F8), 159.2 (C5F7), 157.9 (C1F7), 150.2 (C6F7), 149.5 (C1F7), 138.5 (C1F7), 138.2 (C1F6), 136.4 (C1F7), 129.3 (C1F4), 128.8 (C1F4), 128.0 (C1F3), 127.8 (C1F3), 126.8 (C1F3), 126.0 (C1F3), 124.9 (C1F4), 121.6 (C1F4), 121.0 (C1F4), 118.8 (C1F4), 117.3 (C1F3), 116.2 (C1F4), 109.2 (C1F4), 107.8 (C1F3), 107.6 (C1F2), 106.8 (C1F2), 69.7 (C1F2), 67.9 (C1F2), 67.3 (C1F2), 36.0 (Cquat-Bu-ring F), 30.5 (Cquat-Bu-ring F). UV-Vis 2/nm (e/L mol−1 cm−1) (CH2Cl2, 1.00 × 10−5 mol L−1) 236 (110 000), 280 (52 000), 311 (37 000), 326 (28 000), 350 sh (16 000), 400 sh (7000). ESI-MS m/z 1254.0 [M – PF6]+ (base peak, calc. 1253.5). Found C 60.12, H 4.90, N 4.01%.

The method was as for [Ir(1)][PF6] starting with [Ir(1) (MeOH)][PF6] (59 mmol, 2.00 × 10−5 mol, 3 (17.9 mg, 0.057 mmol) and NH4PF6 (92.3 mg, 0.566 mmol) in MeOH (10 mL). Column chromatography used silica eluting with CH2Cl2 changing to CH3Cl2: MeOH 0.5 and finally CH3Cl2: MeOH 100:1. [Ir(1)][PF6] was isolated as an orange solid (101 mg, 0.072 mmol, 75%). 1H NMR (500 MHz, CD2Cl2) δ/ppm 8.15–8.07 (overlapping m, 4H, H6E+/E+) 7.69 (ddd, J = 8.4, 7.5, 1.5 Hz, 2H, H1B4), 7.66 (dJ = 8.6 Hz, 1H, H4N) 7.32 (d, J = 8.1 Hz, 2H, Hd/dHd), 7.25 (d, J = 2.0 Hz, 2H, H5E), 7.01 (ddd, J = 7.3, 1.3 Hz, 2H, H5E), 6.98–6.91 (overlapping m, 6H, H2G4+N1+N3), 6.77–6.69 (overlapping m, 6H, H1A3t+G3), 6.58 (br, 4H, H2G2), 6.00 (ddd, J = 8.6, 2.5 Hz, 2H, H4E), 4.65 (d, J = 2.5 Hz, 2H, H6H), 4.08–3.96 (m, 4H, H4), 3.80–3.74 (m, 2H, H5b/b′), 3.74–3.69 (m, 2H, HDd/d), 3.69–3.63 (m, 2H, H5b/b′), 3.59–3.53 (m, 2H, HDd/d), 3.53–3.45 (m, 4H, H7), 1.40 (s, 18H, H2Bu-ring E). UV-Vis ε/L mol−1 cm−1 (CH2Cl2, 1.00 × 10−5 mol L−1) 236 (110 000), 280 (52 000), 311 (37 000), 326 (28 000), 350 sh (16 000), 400 sh (7000). ESI-MS m/z 1254.0 [M – PF6]+ (base peak, calc. 1253.5). Found C 60.12, H 4.90, N 4.01%.

Dispersion-corrected density functional calculations (DFT-D) were carried out with the D.01 revision of the Gaussian 09 program package.24 The Becke’s three-parameter B3LYP exchange–correlation functional25,26 was used, together with the 6-31G** basis set for C, H, N and O,27 and the “double-ζ” quality LANL2DZ basis set for the Ir atom.28 Relativistic effects were accounted for by means of an effective core potential (ECP), which was used to replace the inner core electrons of Ir. Intramolecular non-covalent interactions are expected to play a relevant role in the studied systems. Consequently, to get a better description of their molecular geometry, the Grimmes D3 dispersion term with Becke-Johnson damping was added to the B3LYP functional (B3LYP-D3).29,30 The geometries of both the singlet ground electronic state (S0) and the lowest-energy triplet state (T1) were fully optimized. No symmetry restrictions were imposed. The geometry of T1 was calculated at the spin-unrestricted UB3LYP-D3 level using a spin multiplicity of three. All the calculations were performed in the presence of the solvent (CH2Cl2). Solvent effects were considered within the self-consistent reaction field (SCRF) theory using the polarized continuum model (PCM) approach.31–33 Time-dependent DFT (TD-DFT)14–16 calculations of the lowest-lying 30 singlet excited states and the lowest-lying 10 triplets of all the complexes were performed in the presence of the solvent at the minimum-energy geometry optimized for the ground state. The geometry of the first naphthyl-centred triplet excited state of each system was first optimized at TD-DFT level, and then reoptimized at the UB3LYP-D3 level to compare with the results obtained for T1 from DFT calculations.

Device preparation

LECs were prepared on top of a patterned indium tin oxide (ITO, 15 Ω per square) coated glass substrate (http://www.naranjosubstrates.com) previously cleaned as follows: (a) sonication with soap, (b) deionized water, (c) isopropanol and (d) UV-O3 lamp for 20 min. The thickness of the films was determined with an Ambios XP-1 profilometer. Prior to the deposition of the emitting layer, 80 nm of poly(3,4-ethylenedioxythiophene):poly(styrenesulfonate) (PEDOT: PSS) (CLEVIOS™ P VP AI 4083, aqueous dispersion, 1.3–1.7%)
solid content, Heraeus) was coated in order to increase the reproducibility of the cells. The emitting layer (100 nm) was prepared by spin-coating of a MeCN solution consisting of the iTMC with the addition of the ionic liquid (IL) 1-butyl-3-methylimidazolium hexafluorophosphate [Bmim][PF6] (>98.5%, Sigma-Aldrich) in a 4:1 molar ratio (iTMC:IL). The devices were then transferred to an inert atmosphere glovebox (<0.1 ppm O2 and H2O, MBraun), where a layer (70 nm) of aluminium (the top electrode) was thermally evaporated onto the devices using an Edwards Auto500 evaporator integrated in the inert atmosphere glovebox. The area of the device was 6.5 mm². The devices were not encapsulated and were characterized inside the glovebox at room temperature.

Device characterization

The device lifetime was measured by applying a pulsed current and monitoring the voltage and the luminance versus time by a True Colour Sensor MAZeT (MTCSICT Sensor) with a Botest OLT OLED Lifetime-Test System. The average current density is determined by multiplying the peak current density by the average luminance by the average current density. The average luminance by the average current density. The luminance is directly obtained by taking the average of the time-on time and dividing by the total cycle time. The average current density by multiplying the peak current density by the average luminance is determined by multiplying the peak current density by the average luminance.

Results and discussion

Synthesis and characterization of ligand H21 and solvento precursor [Ir(1)(MeOH)2][PF6]

Scheme 1 summarises the synthetic route to the bis(cyclometallating) ligand H21. The precursor 2,7-bis[2-(2-hydroxyethoxy)ethoxy]naphthalene was prepared following published procedures. The choice of 2-(4-fluorophenyl)pyridine as substrate and the reaction conditions for nucleophilic substitution with 2,7-bis[2-(2-hydroxyethoxy)ethoxy]naphthalene were chosen based on those described by Clavier et al. for the formation of 8-(tert-butylthio)-2-methylquinoline. Compound H21 was isolated in 74.6% yield. The base peak in the electrospray mass spectrum (m/z 643.6) arose from the [M + H]+ ion, and elemental analysis was in accord with the expected composition. The ¹H and ¹³C NMR spectra were assigned by 2D methods (COSY, NOESY, HMQC and HMBC) and the spectra were consistent with the symmetrical structure shown in Scheme 1 for H21.

The conventional methodology for preparing complexes of the type [Ir(C-N)2(N=N)2][PF6] is treatment of a [Ir(C-N)2μ-Cl]2 dimer with two equivalents of an N=N ligand followed by anion exchange. However, the unfavourable effect that residual chloride ion has on the performance of LEC devices has led us to develop an alternative strategy in which the chloride dimer is converted to an intermediate solvento complex by treatment of the dimer with AgPF6 in methanol (Scheme 2). The dimer [Ir2(1)2μ-Cl]2 was prepared from H21 and IrCl3·xH2O under dilute conditions to ensure that the two C=N coordination sites of H21 underwent cyclometallation at the same iridium(m) centre. The dimer was then treated with AgPF6 in MeOH to give the solvento complex [Ir(1)(MeOH)2][PF6] which was used for the final reaction step without further purification. Both [Ir2(1)2μ-Cl]2 and [Ir(1)(MeOH)2][PF6] were characterized by ESI MS and NMR spectroscopy. The base peak envelope in the mass spectrum of each complex corresponded to the [Ir(1)+]⁺ cation. Evidence for the coordinated MeOH in [Ir(1)(MeOH)2][PF6] came from δ¹H and δ¹³C NMR signals at 3.34 and 49.9 ppm, respectively. These values compare with δ¹H = 3.35 ppm and δ¹³C = 49.9 ppm in [Ir(Phppy)2(2,7-dimethoxy-9H-carbazole)][PF6] and [Ir(Phphpy)(MeOH)][PF6]. Going from H21 to the coordinated ligand in [Ir2(1)2μ-Cl]2 and [Ir(1)(MeOH)2][PF6] results in a loss in rotational freedom within the polyethyleneoxy-chain, with the consequence that each

![Scheme 2](image-url)
Synthesis and characterization of [Ir(1)(N^N)][PF_6] complexes

The complexes [Ir(1)(N^N)][PF_6] with N^N = 2, 3, 4 or 5 were prepared by reaction of [Ir(1)(MeOH)]_2[PF_6] with the N^N ligand in MeOH at reflux in the presence of NH_4PF_6 (Scheme 2). After work up, [Ir(1)(2)][PF_6], [Ir(1)(3)][PF_6], [Ir(1)(4)][PF_6] and [Ir(1)(5)][PF_6] were isolated in 44.1–75.0% yields. The base peak in the electrospray mass spectrum of each complex corresponded to [M – PF_6]^+ and the isotope pattern of the peak envelope was as predicted. The \(^1\)H and \(^{13}\)C NMR spectra of the complexes were assigned by using COSY, NOESY, HMOC and HMBC experiments. The cations [Ir(1)(N^N)][PF_6] were isolated in 75.0% yields.

The aromatic and alkyl regions of the 500 MHz \(^1\)H NMR spectrum of \([\text{Ir}(\text{ppy})_2(\text{N}^\text{N})]\)[PF_6] (Fig. 2) is consistent with each CH₂ group in the coordinated ligand exhibiting a diastereotopic pair of protons. The \(^1\)H NMR spectrum of [Ir(1)(4)][PF_6] is similar to that of [Ir(1)(2)][PF_6], but with the absence of the signal for H^4d and the addition of a singlet at \(\delta = 1.40\) ppm arising from the tBu substituent in ligand 4. The low frequency shift of the signal for H^4e (\(\delta = 4.62\) ppm in [Ir(1)(2)][PF_6]) and \(\delta = 4.65\) ppm in [Ir(1)(4)][PF_6] is due to a combination of H^4e being sandwiched in a V-shaped cavity between a phenyl and pyridine ring (as observed in related [Ir(C^N)(N^N)]^+ complexes\(^{33,41}\) and from its proximity to one or more oxygen atoms of the polyethyleneoxy-chain. Unfortunately, no X-ray quality single crystals of any of the four complexes could be grown. In Fig. 2, the broadened signal at \(\delta = 6.55\) ppm for protons H^G2 (the ortho-protons on the 6- and 6′-phenyl substituents of ligand 2) is consistent with hindered rotation of the phenyl rings as observed in other [Ir(C^N)(N^N)]^+ complexes with 6-phenyl-2,2′-bipyridine or 6,6′-diphenyl-2,2′-bipyridine N^N ligands.\(^{44,45}\) A similarly broadened resonance for HG2 appears at \(\delta = 6.58\) ppm in the \(^1\)H NMR spectrum of [Ir(1)(4)][PF_6].

On going from [Ir(1)(2)]^+ and [Ir(1)(4)]^+ to [Ir(1)(3)]^+ and [Ir(1)(5)]^+, the complex is desymmetrized. This is apparent from the \(^1\)H NMR spectrum for [Ir(1)(3)][PF_6] (Fig. 3) compared to that of [Ir(1)(2)][PF_6] (Fig. 2). Signals for both H^G2 (\(\delta = 6.57\) ppm) and H^G3 (\(\delta = 6.76\) ppm) are broadened in the \(^1\)H NMR spectrum of [Ir(1)(3)][PF_6] (Fig. 3), and similarly broadened signals are observed for H^G2 (\(\delta = 6.58\) ppm) and H^G3 (\(\delta = 6.76\) ppm) in [Ir(1)(5)][PF_6]. As in [Ir(1)(2)][PF_6] and [Ir(1)(4)][PF_6], this indicates hindered rotation of the 6-phenyl substituent.

Electrochemistry

The electrochemical behaviour of the [Ir(1)(N^N)][PF_6] complexes was investigated using cyclic voltammetry and data are shown in Table 1. A representative cyclic voltammogram (CV) is depicted in Fig. 4. Each complex exhibits an irreversible oxidation and a quasi-reversible reduction process, which are compared to the reference compound [Ir(ppy)_2(4)][PF_6] (Hppy = 2-phenylpyridine). The value of E^ox for each is only marginally affected by the introduction of electron-releasing phenyl and/or tert-butyl groups on the ancillary ligands (e.g. from +0.80 to +0.76 V on going from [Ir(1)(3)][PF_6] to [Ir(1)(4)][PF_6]) leading to a slightly destabilized HOMO. These findings are consistent with the oxidation process being centred on the iridium and C^N ligand, as discussed previously.\(^{42}\)

The reduction processes centred on the N^N ligand are similar for [Ir(1)(4)][PF_6] and [Ir(1)(5)][PF_6] (−1.96 V and
Photophysical properties

The solution absorption spectra of \([\text{Ir(1)}(\text{N}^\text{N})\text{][PF}_6]\) with \(\text{N}^\text{N} = 2-5\) are compared in Fig. 5; the complex \([\text{Ir(ppy)}_2(4)\text{][PF}_6]\) (which has previously been reported)\(^{23}\) is used as a reference compound, differing from \([\text{Ir(1)}(4)\text{][PF}_6]\) only by the introduction of the naphthyl-containing linker between the cyclometalating \(\text{C}^\text{N}\) ligands. The highest-energy absorption band (236 or 237 nm) in each of the spectra of \([\text{Ir}(\text{ppy})_2(\text{4})\text{][PF}_6]\) as compared in Fig. 5; the complex \([\text{Ir(ppy)}_2(4)\text{][PF}_6]\) only by the introduction of tert-butyl groups are introduced into the \(\text{C}^\text{N}\) domain and there is a significant shift of \(E_{\text{irr}}^\text{red}\) to more negative potentials. This is consistent with the LUMO of the complex being localized on the \(\text{N}^\text{N}\) ligand as discussed below and in agreement with, for example, \([\text{Ir(ppy)}_2(2)\text{][PF}_6]\).\(^{43}\)

### Table 1

<table>
<thead>
<tr>
<th>Compound</th>
<th>(E_{\text{irr}}^\text{red}/V)</th>
<th>(E_{\text{irr}}^\text{red}/V)</th>
<th>(\Delta E_{\text{irr}}^\text{red}/V)</th>
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<tr>
<td>([\text{Ir(1)}(2)\text{][PF}_6])</td>
<td>+0.82(^{\text{irr}})</td>
<td>−1.81(^{\text{irr}})</td>
<td>2.63</td>
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<tr>
<td>([\text{Ir(1)}(3)\text{][PF}_6])</td>
<td>+0.80(^{\text{irr}})</td>
<td>−1.84(^{\text{irr}})</td>
<td>2.64</td>
</tr>
<tr>
<td>([\text{Ir(1)}(4)\text{][PF}_6])</td>
<td>+0.76(^{\text{irr}})</td>
<td>−1.96(^{\text{irr}})</td>
<td>2.72</td>
</tr>
<tr>
<td>([\text{Ir(1)}(5)\text{][PF}_6])</td>
<td>+0.77(^{\text{irr}})</td>
<td>−1.92(^{\text{irr}})</td>
<td>2.69</td>
</tr>
<tr>
<td>([\text{Ir(ppy)}_2(4)\text{][PF}_6])</td>
<td>+0.87(^{\text{irr}})</td>
<td>−1.94(^{\text{irr}})</td>
<td>2.81</td>
</tr>
</tbody>
</table>

−1.92 V, respectively) compared to −1.94 V for \([\text{Ir(ppy)}_2(4)\text{][PF}_6]\). On going from \([\text{Ir(1)}(2)\text{][PF}_6]\) to \([\text{Ir(1)}(4)\text{][PF}_6]\) and from \([\text{Ir(1)}(3)\text{][PF}_6]\) to \([\text{Ir(1)}(5)\text{][PF}_6]\), electron-releasing tert-butyl groups are introduced into the \(\text{N}^\text{N}\) domain and there is a significant shift of \(E_{\text{irr}}^\text{red}\) to more negative potentials. This is consistent with the LUMO of the complex being localized on the \(\text{N}^\text{N}\) ligand as discussed below and in agreement with, for example, \([\text{Ir(ppy)}_2(2)\text{][PF}_6]\).\(^{43}\)

The solution absorption spectra of \([\text{Ir(1)}(\text{N}^\text{N})\text{][PF}_6]\) with \(\text{N}^\text{N} = 2-5\) are compared in Fig. 5; the complex \([\text{Ir(pppy)}_2(4)\text{][PF}_6]\) (which has previously been reported)\(^{23}\) is used as a reference compound, differing from \([\text{Ir(1)}(4)\text{][PF}_6]\) only by the introduction of the naphthyl-containing linker between the cyclometalating \(\text{C}^\text{N}\) ligands. The highest-energy absorption band (236 or 237 nm) in each of the spectra of the \([\text{Ir(1)}(\text{N}^\text{N})\text{][PF}_6]\) complexes is assigned to naphthyl-centred \(\pi \rightarrow \pi^*\) transitions; this band is absent in the spectrum of \([\text{Ir(pppy)}_2(4)\text{][PF}_6]\).\(^{23}\) The profile of the absorption bands above −270 nm is similar for the five complexes, consistent with the absorptions between 270 and 375 nm arising from \(\pi \rightarrow \pi^*\) transitions centred on the ppy, 6-Phpppy or 6,6′-Phpppy units. The weakest absorptions at −400−425 nm are assigned to metal-to-ligand charge transfer (MLCT) transitions.

Upon excitation at 280 nm, the complexes all exhibit orange emission. The solution emission spectra are shown in Fig. 6 and emission maxima, quantum yields and lifetimes are given in Tables 2 and 3. Introducing the electron-releasing tert-butyl substituents results in the expected blue-shift in \(\lambda_{\text{em}}^\text{max}\) (23 nm on going from \([\text{Ir(1)}(3)\text{][PF}_6]\) to \([\text{Ir(1)}(5)\text{][PF}_6]\), and 26 nm from \([\text{Ir(1)}(2)\text{][PF}_6]\) to \([\text{Ir(1)}(4)\text{][PF}_6]\). Introducing the naphthyl-containing linker between the two ppy moieties has little effect on \(\lambda_{\text{em}}^\text{max}\) (558 nm in model compound \([\text{Ir(pppy)}_2(4)\text{][PF}_6]\) and 564 nm in \([\text{Ir(1)}(4)\text{][PF}_6]\) consistent with the emission originating from a mixture of MLCT and LLCT transitions\(^{6}\) involving the \(\text{Ir(n)}\) metal centre and the ppy units of the \(\text{C}^\text{N}\) ligands in the HOMO and the \(\text{N}^\text{N}\) ligand in the LUMO as discussed below. This is consistent with the electrochemical data (Table 1). In degassed solutions, the \([\text{Ir(1)}(\text{N}^\text{N})\text{][PF}_6]\) orange emissions exhibit photoluminescence quantum yields (PLQYs) in the range 6−18% (Table 2) with lifetimes of between 135 and 370 ns (Table 3). Fig. 7 depicts the emission spectra of solid-state samples of the complexes. For \([\text{Ir(1)}(2)\text{][PF}_6]\) and
Table 2 Emission maxima\(^a\) and quantum yields for \([\text{Ir}(\text{N}^\text{N})]\)[PF\(_6\)] complexes in solution and as powder samples

<table>
<thead>
<tr>
<th>Complex cation</th>
<th>CH(_2)Cl(_2) solution</th>
<th>Powder</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>(\lambda_{\text{em}}^\text{max}) [nm]</td>
<td>PLQY(^b) [%]</td>
</tr>
<tr>
<td>([\text{Ir}(2)])(^+)</td>
<td>434, 590</td>
<td>16</td>
</tr>
<tr>
<td>([\text{Ir}(3)])(^+)</td>
<td>361, 601</td>
<td>6</td>
</tr>
<tr>
<td>([\text{Ir}(4)])(^+)</td>
<td>425, 564</td>
<td>9</td>
</tr>
<tr>
<td>([\text{Ir}(5)])(^+)</td>
<td>435, 578</td>
<td>18</td>
</tr>
<tr>
<td>(<a href="4">\text{Ir}(ppy)</a>])(^+)</td>
<td>420, 558(^\dagger)</td>
<td>1</td>
</tr>
</tbody>
</table>

\(^a\) \(\lambda_{\text{exc}} = 280\) nm. \(^b\) Argon degassed. \(^\dagger\) A value of 555 nm has been reported for a CH\(_2\)Cl\(_2\) solution of \([\text{Ir}(ppy)](4)]\)[PF\(_6\)].\(^23\)

\([\text{Ir}(1)(3)][PF\(_6\)]\) going from solution to solid leads to a blue-shift in \(\lambda_{\text{em}}^\text{max}\), whereas for the complexes containing the tert-butyl substituents, \(\lambda_{\text{em}}^\text{max}\) is little affected. Both the PLQY and emission lifetimes are enhanced from solution to solid state (Tables 2 and 3). We note that the solution PLQY recorded for the reference complex \([\text{Ir}(ppy)](4)]\)[PF\(_6\)] of 1% (Table 2) is consistent with the 0.07% previously reported.\(^23\)

The emission spectra of the \([\text{Ir}(\text{N}^\text{N})]\)[PF\(_6\)] complexes as amorphous thin films with the composition used in LEC devices were also recorded (Fig. S1\(^\dagger\)). The spectra are similar and follow the same trends discussed above for the powder spectra with the emission maxima slightly red-shifted ([Ir(1)(2)]\)[PF\(_6\)]: 583 nm, [Ir(1)(3)]\)[PF\(_6\)]: 589 nm, [Ir(1)(4)]\)[PF\(_6\)]: 574 nm, [Ir(1)(5)]\)[PF\(_6\)]: 579 nm. The PLQY values (see Table 4) obtained in thin film are comparable to those determined in powder (21.9% for [Ir(1)(5)]\)[PF\(_6\)], 17.5% for [Ir(1)(4)]\)[PF\(_6\)] and 15.5% for [Ir(1)(3)]\)[PF\(_6\)] except for [Ir(1)(2)]\)[PF\(_6\)] which shows a significantly lower PLQY of 22.6% compared to 35% in the powder (Table 2).

A comparison of Fig. 6 and 7 reveals that each complex in solution, most noticeably \([\text{Ir}(1)(4)]\)[PF\(_6\)], exhibits a second emission at higher energies than the dominant band, and that this emission band is absent in the powder samples. Since the aim of introducing the naphthyl domain into the \([\text{Ir}(\text{C}^\text{N})\text{N}^\text{N}])\) complex was to generate a dual emitter, the appearance of the second band, albeit weak, demanded further investigation. The absence of the band in the solid state samples suggested that the high-energy emission might arise from dissociated ligand. Fig. 8 shows an overlay of the normalized emission spectra of \([\text{Ir}(1)(4)]\)[PF\(_6\)], \([\text{Ir}(ppy)](4)](4)]\)[PF\(_6\)], \(\text{H}_{\text{2}}\text{1}\) and \(\text{4}\). Both \([\text{Ir}(1)(4)]\)[PF\(_6\)] and \([\text{Ir}(ppy)](4)](4)]\)[PF\(_6\)] exhibit a band with \(\lambda_{\text{em}}^\text{max}\) ~ 420 nm (Table 2), suggesting that the origin of the emission is not the naphthyl domain. Note that the previously reported emission spectrum of \([\text{Ir}(ppy)](4)](4)]\)[PF\(_6\)] did not extend to below 450 nm.\(^23\) Model compounds HOnaphth (HOnaphth = 2,7-bis [2-(2-hydroxyethoxy)ethoxy]naphthalene) and \(\text{H}_{\text{2}}\text{1}\) both show a structured emission at 331 and 344 nm typical of naphthalene;\(^46\) in addition \(\text{H}_{\text{2}}\text{1}\) shows an emission at 410 nm. Unexpectedly, excitation of a CH\(_2\)Cl\(_2\) solution of ligand \(\text{4}\) gives rise to an extremely similar PL spectrum as that of \(\text{H}_{\text{2}}\text{1}\) and HOnaphth (Fig. 8). Repeated measurements for \(\text{4}\) and comparisons with the emission spectrum of \(\text{5}\) and with the reported emission spectrum of 4,4’-di-tert-butyl-2,2’-bipyridine (\(\lambda_{\text{em}}^\text{max}\) = 357 nm)\(^47\) validated the result.

Excitation spectra of \([\text{Ir}(1)(4)]\)[PF\(_6\)] and \([\text{Ir}(ppy)](4)](4)]\)[PF\(_6\)] (Fig. S2\(^\dagger\)) reveal the origins of the 564 and 425 nm emissions in \([\text{Ir}(1)(4)]\)[PF\(_6\)] and the 558 and 420 nm bands in \([\text{Ir}(ppy)](4)](4)]\)[PF\(_6\)]. The low-energy MLCT emissions for both complexes arise from absorptions over the whole region from 230 to 500 nm, and the profiles of the excitation spectra are

Table 3 Emission lifetimes for \([\text{Ir}(\text{N}^\text{N})\text{N}][PF\(_6\)]\) complexes in solution and as powder samples\(^a\)

<table>
<thead>
<tr>
<th>Complex cation</th>
<th>CH(_2)Cl(_2) solution</th>
<th>Powder</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>(\tau_{\text{av}}^a) [ns]</td>
<td>(\tau_1) [ns] (A(_1))</td>
</tr>
<tr>
<td>([\text{Ir}(1)(2)])(^+)</td>
<td>587</td>
<td>381</td>
</tr>
<tr>
<td>([\text{Ir}(1)(3)])(^+)</td>
<td>601</td>
<td>159</td>
</tr>
<tr>
<td>([\text{Ir}(1)(4)])(^+)</td>
<td>564</td>
<td>144</td>
</tr>
<tr>
<td>([\text{Ir}(1)(5)])(^+)</td>
<td>420</td>
<td>4</td>
</tr>
</tbody>
</table>

\(^a\) \(\lambda_{\text{exc}} = 280\) nm; lifetimes were measured using a Hamamatsu QuantaTaurus-Tau (see Experimental section) and some values of \(\lambda_{\text{em}}^\text{max}\) differ slightly from those in Table 2 where spectra were recorded on a Shimadzu RF-5301PC spectrofluorometer. \(^b\) Biexponential fit using the equation \(\tau_{\text{av}} = \sum A_i\tau_i/\sum A_i\) where \(A_i\) is the pre-exponential factor of the lifetime.
similar to the absorption spectra shown in Fig. 5, but, as expected, lack the naphthyl band at 237 nm. The 425 or 420 nm emission results from two absorption bands at ~258 and ~358 nm (Fig. 9, black and cyan spectra). The excitation spectra for [Ir(1)(4)][PF₆] and [Ir(ppy)₂(4)][PF₆] resemble that of 4 (Fig. 9). In contrast, the excitation spectrum of H₂1 for the emission at 410 nm (Fig. 9, green spectrum) is broader. The similarity between the excitation spectra of [Ir(1)(4)][PF₆], [Ir(ppy)₂(4)][PF₆] and free ligand 4 indicate that the high-energy emission around 420 nm arises from a fluorescent excited state of coordinated ligand 4. Although an extremely low concentration impurity could be responsible for such observations, the method of measuring and the reproducibility of recorded solutions of independently synthesized compounds makes this scenario unlikely.

The PLQY increases from 1% to 9% on going from [Ir(ppy)₂(4)][PF₆] to [Ir(1)(4)][PF₆], but this is not necessarily a consequence of partial energy transfer from the naphthyl unit (PLQY = 15% for H₂1) to the Ir(n) coordination sphere. Excited-state lifetime measurements of [Ir(1)(4)][PF₆] (τₑ = 144 ns for the 564 nm and τₑ = 4 ns for the 420 nm emission band, Table 3) corroborate the fact that the higher-energy emission arises from a fluorescent excited state being in the same range as for H₂1 (τₑ = 6 ns for the 420 nm emission band).

Theoretical calculations

To gain a better understanding of the electrochemical and photophysical properties of complexes [Ir(N^N)]⁺, (N^N = 2–5), a combined DFT/TD-DFT theoretical investigation was undertaken at the B3LYP-D3/(6-31G** + LANL2DZ) level in the presence of the solvent (CH₂Cl₂) (see the Experimental section for full computational details). Calculations were also performed for the reference complex [Ir(ppy)₂(4)]⁺ for comparison purposes.

The geometry of the complexes in their ground electronic state (S₀) was optimized without imposing any symmetry restriction. The calculated geometries reproduce the trends observed typically on this type of complexes, showing a distorted octahedral coordination of the iridium atom. Fig. 10 displays the minimum-energy optimized structures of complexes [Ir(1)(2)]⁺ and [Ir(1)(3)]⁺ as representative examples. Table S1† summarizes the values of the geometrical parameters defining the iridium coordination sphere and of selected distances and dihedral angles. For all the complexes, the pendant phenyl rings introduced as R and R₁ substituents in the N^N ligand (Scheme 2) present intracation face-to-face π-stacking interactions with the phenyl rings of the closest ppy in the cyclometallating ligand. The calculated centroid–centroid distances between rings A and G in Scheme 2 range from 3.40 to 3.56 Å, in good agreement with the X-ray values reported for complexes [Ir(ppy)₂(2)]⁺ and [Ir(ppy)₂(3)]⁺. The steric hindrance produced by the phenyl substituents induces a twisting between the rings of the bpy ligand. The twisting angle is higher for [Ir(1)(2)]⁺ (32.8°), [Ir(1)(4)]⁺ (38.3°) and [Ir(ppy)₂(4)]⁺ (37.1°) than for [Ir(1)(3)]⁺ (21.3°) and [Ir(1)(5)]⁺.

![Fig. 8](image1.png)

**Fig. 8** Normalized solution emission spectra of [Ir(1)(4)][PF₆], [Ir(ppy)₂(4)][PF₆], H₂1, HOnaph and 4 in CH₂Cl₂ (~1.00 × 10⁻⁵ mol L⁻¹). Compounds were excited at 280 nm.

![Fig. 9](image2.png)

**Fig. 9** Normalized excitation spectra for CH₂Cl₂ solutions (1.00 × 10⁻⁵ M) of [Ir(1)(4)][PF₆], [Ir(ppy)₂(4)][PF₆], H₂1 and 4 each at a fixed emission wavelength of 420 nm.

![Fig. 10](image3.png)

**Fig. 10** Minimum-energy optimized structures calculated at the B3LYP-D3/(6-31G** + LANL2DZ) level for [Ir(1)(2)]⁺ and [Ir(1)(3)]⁺. Face-to-face interactions between pendant phenyl rings attached to the bpy ligand and phenyl rings of the ppy ligands are emphasized. Hydrogen atoms have been omitted.
(24.9°) because in the former two pendant phenyl rings are introduced in R and R₁ positions. For all the [Ir(1)](N’N’)] complexes, the cyclometallating ligand 1 remains in an expanded disposition (Fig. 10), featuring a distance from the centroid of the naphthalene group to the iridium atom of 10.6–10.8 Å.

The geometry of the complexes in their first triplet excited state (T₁) was also optimized using the spin-unrestricted UB3LYP-D3/(6-31G**+LANL2DZ) approach. It is worth to note that the face-to-face intramolecular interactions observed in the ground state are preserved in T₁, and that the naphthalene-iridium distance remains almost constant. The most important changes are found for the bpy ligand that becomes more planar in T₁ (twisting angles in the 13–18° range, Table S1, ESIF). This points to a higher electron density on the bpy ligand in T₁, which would stabilize the planar form as it favors the electronic delocalization.

Fig. 11 displays the isovalue contours calculated for the highest-occupied (HOMO) and lowest-unoccupied (LUMO) molecular orbitals of complex [Ir(1)(2)]²⁺ as a representative example. Orbitals HOMO−1 and LUMO+7, which respectively correspond to the highest-occupied and lowest-unoccupied molecular orbitals centred on the naphthalene group of ligand 1, are also displayed. The topology of the corresponding MOs of all the [Ir(1)(N’N’)] complexes fully reproduces that of the selected example. The table inserted in Fig. 11 summarizes the MO energies calculated for the [Ir(1)(N’N’)] complexes and compares them with those obtained for the HOMO and LUMO of [Ir(ppy)₂(4)]²⁺. As it is usually found for ppy-based cyclometallated Ir-TMC complexes, the HOMO results from a mixture of s₆ orbitals of Ir(III) and phenyl π orbitals, with some contribution from the pyridine rings, of the cyclometallating ligand, whereas the LUMO is located over the bpy of the ancillary ligand. The similar energy values estimated for the HOMO of [Ir(1)(N’N’)] are in good agreement with the experimental Eₓ values (Table 1), which do not change greatly along the series.

The introduction of electron-releasing tert-butylic groups in the N’N’ ligand slightly destabilizes the HOMO of [Ir(1)(4)]²⁺ and [Ir(1)(5)]²⁺, in good agreement with the slightly lower Eₓ values recorded for these complexes. The effect of the substituents is higher in the LUMO, centred on the ancillary ligand. It becomes stabilized along the series [Ir(1)(4)]²⁺ (−2.30 eV) > [Ir(ppy)₂(4)]²⁺ (−2.34 eV) > [Ir(1)(5)]²⁺ (−2.41 eV) > [Ir(1)(2)]²⁺ (−2.47 eV) > [Ir(1)(3)]²⁺ (−2.56 eV), in quite good correlation with the less negative E₁/₂ values recorded along this series (Table 1).

The HOMO–LUMO gaps calculated for [Ir(1)(2)]²⁺ (3.20 eV) and [Ir(1)(3)]²⁺ (3.13 eV) are lower than those computed for [Ir(1)(4)]²⁺ (3.33 eV) and [Ir(1)(5)]²⁺ (3.24 eV) in good accord with the trends observed for the electrochemical gap ΔEᶜₑ. If light emission in these complexes originates from a state described by the HOMO → LUMO excitation, MO calculations predict that [Ir(1)(4)]²⁺ would emit more in the blue that the rest of the complexes and close to the emission of [Ir(ppy)₂(4)]²⁺. This is in fact the trend observed in solution for the ƛₑ values (Table 2). The significant energy difference between the LUMO and the LUMO+7 centred on the naphthalene moiety leads us to expect that states associated to this moiety will appear at high energies and will not contribute to the emission.

The nature of the low-lying triplet states was first studied by performing TD-DFT calculations at the optimized geometry of the ground state (S₀). The vertical excitation energies and electronic descriptions computed for the lowest-lying triplet states of the [Ir(1)(N’N’)] complexes and those of [Ir(ppy)₂(4)]²⁺ are given in Table S2.† All the complexes present a first triplet excited state (T₁) mainly defined by the HOMO → LUMO excitation, which implies an electron transfer from the Ir-ppy environment, where the HOMO is localized, to the bpy ligand, where the LUMO resides (see Fig. 11). The T₁ state therefore shows a mixed 3MLCT/3LLCT character, and the calculated excitation energies ([Ir(ppy)₂(4)]²⁺: 2.56 eV > [Ir(1)(4)]²⁺: 2.55 eV > [Ir(1)(5)]²⁺: 2.53 eV > [Ir(1)(2)]²⁺: 2.48 eV > [Ir(1)(3)]²⁺: 2.43 eV) follow the trend expected on the basis of the HOMO–LUMO gap. The T₁ state was further examined by optimizing its structure at the spin-unrestricted UB3LYP-D3 level as described above. The unpaired-electron spin density distribution calculated for this state (Fig. 12a) matches the topology of the HOMO → LUMO excitation and therefore corroborates the electron transfer from the Ir-ppy environment to the N’N’ ligand, thus confirming the 3MLCT/3LLCT character of T₁. The electronic nature predicted for T₁ is in good agreement with the broad unstructured shape of the emission bands observed experimentally (Fig. 6 and 7).

Back to the TD-DFT calculation of triplet states, the first excited state involving the naphthyl group of ligand 1 is found 0.3–0.4 eV above T₁ (Table S2†). As was to be expected, it is mainly described by the HOMO−1 → LUMO+7 excitation, and its vertical excitation energy from S₀ remains almost constant (2.85–2.86 eV) for all the [Ir(1)(N’N’)] complexes, as the structural differences between these complexes concern the N’N’ ligand from which the naphthyl is far apart. The spin-density distribution calculated for this state is shown in Fig. 12b and clearly reflects the localization of the electronic transition on
the naphthyl unit. After full-geometry relaxation at the UB3LYP-D3 level, the naphthyl-centred triplet state is calculated 0.38, 0.41, 0.28 and 0.30 eV above the HOMO → LUMO $T_1$ triplet for $[\text{Ir}(1)(2)]^+$, $[\text{Ir}(1)(3)]^+$, $[\text{Ir}(1)(4)]^+$ and $[\text{Ir}(1)(5)]^+$, respectively. This energy difference is large enough to discard the contribution of the naphthyl-centred triplet excited state to the phosphorescence emission, which is predicted to occur from the lowest-energy, HOMO → LUMO, 3MLCT/3LLCT $T_1$ state.

TD-DFT calculations were also performed for singlet excited states to investigate the nature of the bands appearing in the experimental absorption spectra (Fig. 5). Singlet excited states with mainly MLCT character and small oscillator strengths are found in the 400 nm region, followed by ligand-centred (LC) $\pi \rightarrow \pi^*$ states involving both the ppy and bpy moieties with some MLCT/LLCT character between 400 and 300 nm. No excited state implying the naphthyl moiety of ligand 1 is found below 300 nm. According to the theoretical results, it is unlikely that emission, fluorescent or phosphorescent, takes place from naphthyl-centred excited states.

Electroluminescence devices

The electroluminescence properties of complexes $[\text{Ir}(1)(N^N)]\text{[PF}_6\text{]}$ with $N^N = 2$–5 and complex $[\text{Ir}(\text{ppy})(4)]\text{[PF}_6\text{]}$ were investigated by using them as emitters in LECs. From now on, LECs containing $[\text{Ir}(1)(2)]\text{[PF}_6\text{]}$, $[\text{Ir}(1)(3)]\text{[PF}_6\text{]}$, $[\text{Ir}(1)(4)]\text{[PF}_6\text{]}$, $[\text{Ir}(1)(5)]\text{[PF}_6\text{]}$ and $[\text{Ir}(\text{ppy})(4)]\text{[PF}_6\text{]}$ are referred as LECs A–E, respectively (Table 4). The LECs were prepared on ITO-patterned glass substrates in a double-layer structure. They consist of a PEDOT:PSS layer (80 nm) and the electroluminescent active layer (100 nm) sandwiched between two electrodes. Aluminium was thermally evaporated as the top cathode (70 nm). The LECs were characterized under inert conditions using a pulsed current driving of 25 A m$^{-2}$ (average current density), a frequency of 1000 Hz and duty cycles of 50%. This operation method provides faster response and longer lifetimes than constant voltage operation. In LECs, the initial limited injection is assisted by the ion accumulation at the electrode interface and the formation of electrochemical p- and n-doped regions. Therefore, the behaviour of LECs is mainly governed by the ion mobility in the amorphous thin film. Even though iTMCs are intrinsically ionic, additional ions are usually added in the active layer to speed up the device response and to balance the carriers injected. In this work, the ionic liquid (IL) 1-butyl-3-imidazolium hexafluoridophosphate $[\text{Bmim}]\text{[PF}_6\text{]}$ was added to the active layer at a molar iTMC : IL ratio of 4 : 1.

Fig. 13 displays the luminance versus time curves for LECs A–E, and Table 4 collects the parameters that summarize the

![Fig. 13](image_url)
device performances. The turn-on time (t\text{max}) is defined as the time to reach the maximum luminance (L\text{max}) and the device lifetime (t\text{1/2}) is the time to reach one-half of L\text{max} after this value is achieved. LEC devices can be divided in two groups with LECs D and E, which incorporate complexes [Ir(1)(5)][PF\text{6}] and [Ir(ppy)(4)][PF\text{6}], showing shorter t\text{max} and faster luminance decays than LECs A-C (Table 4). LECs D and E show t\text{max} and t\text{1/2} below 3.5 and 35 hours, respectively, whereas devices A-C have longer turn-on times above 45 hours and a much more stable behaviour with t\text{1/2} values ranging from 581 to 785 hours. In contrast, LECs D and E achieve higher luminance values above 100 cd m\text{-2} (133 and 146 cd m\text{-2}, respectively) compared with LECs A-C, which show maximum luminances of 91, 67 and 54 cd m\text{-2}, respectively. The maximum values obtained for the efficacy, the power conversion efficiency (PCE) and the external quantum efficiency (EQE) of LEC D (5.4 cd A\text{-1}, 2.2 lm W\text{-1} and 2.1%, respectively), are slightly lower than those found for LEC E (6.4 cd A\text{-1}, 3.7 lm W\text{-1} and 2.6%), and are similar to those reported for other orange-emitting Ir-TMC-LECUs under pulsed current operation.\textsuperscript{49,53} The efficiency, PCE and EQE values found for LECs A-C are smaller than those obtained for D, and decrease in passing from A to B and to C (Table 4). This trend is in agreement with the trend of the PLQY for the anthracene thin film (device environment) of the complexes [Ir(1)(2)][PF\text{6}], [Ir(1)(3)][PF\text{6}] and [Ir(1)(4)][PF\text{6}].

The main differentiator between the complexes [Ir(1)(5)][PF\text{6}] and [Ir(ppy)(4)][PF\text{6}] used to prepare LECs D and E is the naphthyl-bridged functionalization of the ppy ligands in the case of LEC D. Yet both the device responses t\text{max} and t\text{1/2} are comparable for these two devices, which indicates that the ppy functionalization does not influence the ionic movement in the films. Therefore, the longer t\text{max} and t\text{1/2} found for LECs A-C should not be ascribed to the large size of ligand 1. Moreover, the [Ir(1)(2)][PF\text{6}] complex, used in the fabrication of LEC A, exhibits the highest PLQY in thin film (22.6%), whereas the device efficiency is superior for D incorporating a complex with a slightly lower PLQY (21.9%). Considering these PLQY values and a typical outcoupling of 20%, the theoretical maximum EQEs predicted for LECs A and D when all injected electrons and holes combine have very similar values of 4.5% and 4.4%, respectively. As the EQE achieved for D (2.1%) is slightly closer to the theoretical value than that obtained for A (1.8%), we can hypothesize that the lower exciton-quenching for D could be the result of a better-balanced carrier injection due to the faster response. Hence, for the devices with slower response (devices A-C), lower efficiencies were achieved. Comparable characteristics have been observed for similar orange complexes reported in LECs.\textsuperscript{43}

The electroluminescence (EL) spectrum of all the LECs was registered during the operation of the devices (Fig. S3\textsuperscript{†}). All the LECs show orange electroluminescence with maxima in the 580–590 nm range but for LEC C ([Ir(1)(4)][PF\text{6}]) which emits at 575 nm. The EL spectra are similar to the PL spectra recorded in powder (Fig. 7) and in thin film (Fig. S1\textsuperscript{†}).

Conclusions

We have prepared and characterized a series of cyclometalated [Ir(C\textsuperscript{3}N\textsubscript{2})(N\textsuperscript{3}N)][PF\text{6}] compounds in which the two cyclometalating C\textsuperscript{3}N units are connected by a naphthyl-containing linker. The N\textsuperscript{3}N ligand (2–5) is a 2,2′-bipyrindine functionalized with phenyl and tert-butyl groups. The electrochemical and photophysical properties were compared with those of [Ir(ppy)(4)][PF\text{6}]. The complexes containing the naphthyl-unit exhibit similar absorption spectra, which differ from that of [Ir(ppy)(4)][PF\text{6}] only in the presence of an intense absorption at ~236 nm arising from naphthyl-centred π → π* transitions. Excitation at 280 nm leads to an orange emission for solutions of each complex, and going from solution to powder or thin film leads to little change or to a small blue shift in the emission. The incorporation of the naphthalene unit does not lead to a desirable blue contribution to the emission, and DFT/ TD-DFT calculations were performed to understand this observation. The energy difference between the LUMO and the lowest-unoccupied MO centred on the naphthyl moiety (LUMO+7) is large enough to explain why there is no contribution from the naphthyl-centred triplet excited state to the phosphorescence emission. Singlet excited states were also investigated. LECs using the [Ir(1)(N\textsuperscript{3}N)][PF\text{6}] and [Ir(ppy)(4)][PF\text{6}] complexes in the emissive layer led to long living devices with modest turn-on times. The presence of the naphthyl-bridge between the cyclometallating units does not significantly alter the device response, indicating that it does not play a significant role in the ionic transport.

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Notes and references

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