Dalton Transactions



PAPER

View Article Online



Cite this: *Dalton Trans.*, 2016, **45**, 11668

Regioisomerism in cationic sulfonyl-substituted $[Ir(C^N)_2(N^N)]^+$ complexes: its influence on photophysical properties and LEC performance†

Cathrin D. Ertl,^a Lidón Gil-Escrig,^b Jesús Cerdá,^b Antonio Pertegás,^b Henk J. Bolink,^b José M. Junquera-Hernández,^b Alessandro Prescimone,^a Markus Neuburger,^a Edwin C. Constable,^a Enrique Ortí*^b and Catherine E. Housecroft*^a

A series of regioisomeric cationic iridium complexes of the type $[Ir(C^N)_2(bpy)][PF_6]$ (bpy = 2,2'-bipyridine) is reported. The complexes contain 2-phenylpyridine-based cyclometallating ligands with a methylsulfonyl group in either the 3-, 4- or 5-position of the phenyl ring. All the complexes have been fully characterized, including their crystal structures. In acetonitrile solution, all the compounds are green emitters with emission maxima between 493 and 517 nm. Whereas substitution meta to the Ir-C bond leads to vibrationally structured emission profiles and photoluminescence quantum yields of 74 and 77%, placing a sulfone substituent in a para position results in a broad, featureless emission band, an enhanced quantum yield of 92% and a shorter excited-state lifetime. These results suggest a larger ligand-centred (³LC) character of the emissive triplet state in the case of meta substitution and a more pronounced charge transfer (CT) character in the case of para substitution. Going from solution to the solid state (powder samples and thin films), the emission maxima are red-shifted for all the complexes, resulting in green-vellow emission. Data obtained from electrochemical measurements and density functional theory calculations parallel the photophysical trends. Light-emitting electrochemical cells (LECs) based on the complexes were fabricated and evaluated. A maximum efficiency of 4.5 lm W⁻¹ at a maximum luminance of 940 cd m⁻² was observed for the LEC with the complex incorporating the sulfone substituent in the 4-position when operated under pulsed current driving conditions.

Received 6th April 2016, Accepted 27th April 2016 DOI: 10.1039/c6dt01325b

www.rsc.org/dalton

Introduction

Ionic transition metal complexes (iTMCs), 1,2 in particular bis-cyclometallated cationic iridium(III) complexes of the type $[Ir(C^N)_2(N^N)]^+$, are promising candidates for application in light-emitting electrochemical cells (LECs) and have therefore been intensely studied as emissive materials. In this class of iridium emitters, high quantum yields are possible with emission colours spanning the whole visible spectrum, combined with high stability of the complexes. 3,4 Emission colour tuning is possible by varying the cyclometallating (C^N) ligands as well as the ancillary (N^N) ligand. The frontier orbitals in

While fluorine substituents have been widely used as electron-withdrawing groups on the cyclometallating ligands, 9-12 the use of sulfone groups has remained limited. Most examples of sulfone-substituted cyclometallating ligands in iridium complexes are those of neutral emitters used in organic light-emitting diodes (OLEDs). 13-19 Recently, we have reported the synthesis and characterization of cationic [Ir(C^N)₂(N^N)]⁺ complexes containing sulfone-substituted phenylpyrazole²⁰ or phenylpyridine ligands,²¹⁻²³ which give promising performances in LECs. In comparison with the archetypal $[Ir(ppy)_2(bpy)][PF_6]^{24}$ (Hppy = 2-phenylpyridine, bpy = 2,2'-bipyridine), replacement of Hppy by 2-(4-methylsulfonylphenyl)pyridine (H1) leads to a 92 nm blue-shift of the maximum in MeCN $[Ir(1)_2(bpy)][PF_6]$ in Scheme 1).²²

these types of complexes are spatially separated, with the HOMO being located mainly on the iridium centre and the C^N ligands and the LUMO on the N^N ligand. $^{5-7}$ An increase in the HOMO-LUMO gap and a resulting blue-shift in the emission maximum are therefore achieved by the combination of electron-withdrawing substituents on the C^N ligands and electron-donating groups on the N^N ligand. 3,4,8

^aDepartment of Chemistry, University of Basel, Spitalstrasse 51, CH-4056 Basel, Switzerland. E-mail: catherine.housecroft@unibas.ch

^bInstituto de Ciencia Molecular, Universidad de Valencia, Catedrático José Beltrán 2, Paterna, E-46980, Spain. E-mail: enrique.orti@uv.es

 $[\]dagger$ Electronic supplementary information (ESI) available: Fig. S1–S4 and Table S1: PL and EL spectra, CVs and singlet TD-DFT calculations. CCDC 1421913–1421915. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/c6dt01325b

Dalton Transactions Paper

Scheme 1 Chemical structures of cyclometallating ligands H1-H3 and complexes $[Ir(C^N)_2(bpy)][PF_6]$ (C^N = [1] to [3]) with methylsulfonyl substituents in different positions of the cyclometallating ligands.

With the aim of achieving blue emission, we recently reported the synthesis of a series of iridium complexes with 2-(4-methylsulfonylphenyl)pyridine (H1) and 2-(3-methylsulfonylphenyl) pyridine (H2) as cyclometallating ligands and electron-rich pyrazolylpyridine as the diimine N^N ligand.²³ Surprisingly, a difference of ~30 nm in the emission maximum was observed on changing the substitution position of the sulfone group.

There have been few investigations on the influence of the substitution position of ligand functionalities on the properties of [Ir(C^N)2(N^N)]+ complexes. Examples include the introduction of diphenylamino, ²⁵ fluoro, ²⁶ methyl, ²⁷ trifluoromethyl, ^{28,29} methylpyridinium, ³⁰ bromo and fluorenyl ³¹ and benzylsulfonyl¹⁵ groups in the 3-, 4- and 5-positions of the cyclometallating phenyl ring. In all of these studies, except for the one investigating bromo and fluorenyl groups,31 it was found that substitution in the 4-position, i.e. para to the Ir-C bond, has the largest influence on emission colour. This observation was supported by density functional theory (DFT) calculations.²⁷ In most of the cases, however, cyclometallating ligands with substituents in the 5-position, meta to the Ir-C bond, have been used. Only a few examples of iridium complexes with substituents in the 4-position of the cyclometallating ligands have been reported, 32,33 and even fewer with substituents in the 3-position. 34

Herein, we report the synthesis of a series of [Ir-(C^N)₂(bpy) [PF₆] complexes with methylsulfonyl groups in the 3-, 4- and 5-positions of the phenyl ring of the cyclometallating ligands (Scheme 1). The effect of the substitution position on the photophysical and electrochemical properties was investigated and further analysed with the help of DFT calculations. The electroluminescence properties of the complexes have been investigated in LEC devices operated under pulsed driving conditions.

Experimental

General

Microwave reactions were carried out in a Biotage Initiator 8 reactor. ¹H and ¹³C{¹H} NMR spectra as well as 2D NMR

spectra were recorded on a Bruker Avance III-500 spectrometer at 295 K; chemical shifts were referenced to residual solvent peaks with δ (TMS) = 0 ppm. For electrospray ionization (ESI) and LC-ESI mass spectra, a Bruker Esquire 3000^{plus} spectrometer and a combination of Shimadzu (LC) and Bruker AmaZon X instruments were used, respectively. FT-IR spectra were recorded on a Perkin Elmer Spectrum Two UATR instrument. Absorption spectra were recorded on an Agilent 8453 spectrophotometer and solution emission spectra on a Shimadzu 5301PC spectrofluorophotometer. Solution and powder photoluminescence quantum yields were recorded on a Hamamatsu absolute PL quantum vield spectrometer, C11347 Quantaurus QY. Emission spectra of powder samples as well as solution and powder excited-state lifetime measurements were carried out using a Hamamatsu Compact Fluorescence Lifetime spectrometer, C11367 Quantaurus Tau. The photoluminescence (PL) properties (spectra and quantum yields) in thin films were measured using a Hamamatsu absolute quantum vield C9920. Electrochemical measurements were performed using cyclic and square wave voltammetry on a CH Instruments 900B potentiostat with glassy carbon working and platinum auxiliary electrodes; a silver wire was used as a pseudo-reference electrode. Dry, purified CH₃CN was used as the solvent and 0.1 M [n Bu₄N][PF₆] as the supporting electrolyte. Ferrocene as an internal reference was added at the end of each experiment.

Synthesis

The synthesis of $[Ir(1)_2(bpy)][PF_6]$, ²² 2-(3-methylsulfonylphenyl) pyridine $(H2)^{23}$ and $[Ir(2)_2Cl]_2^{23}$ has been reported previously. 2-(2-Methylsulfonylphenyl)pyridine (H3) was prepared according to a literature method and ¹H NMR data matched those reported.35,36 Silica was purchased from Fluka (silica gel 60, 0.040-0.063 mm).

[Ir(3)₂Cl]₂. 2-(2-Methylsulfonylphenyl)pyridine (354 mg, 1.52 mmol) was suspended in a mixture of 2-ethoxyethanol and H₂O (3:1, 4 mL) in a microwave vial and purged with N₂. IrCl₃·xH₂O (ca. 82%, 307 mg, 0.842 mmol) was added and the mixture was heated at 110 °C for 1.5 h in a microwave reactor (2 bar). The resulting precipitate was filtered off, washed with H₂O and EtOH, and redissolved in DCM and the solvent was removed under reduced pressure. The orange precipitate formed in the filtrate was filtered off, washed with EtOH, and redissolved with DCM and the solvent removed. Both residues were combined to yield the product as a brownish-orange solid (424 mg, 0.306 mmol, 80.5%). ¹H NMR (500 MHz, CDCl₃) δ /ppm 9.33 (pseudo-dt, J = 8.5, 1.1 Hz, 4H, H^{B3}), 9.14 (ddd, J = 5.8, 1.8, 0.8 Hz, 4H, H^{B6}), 7.92 (ddd, *J* = 8.5, 7.4, 1.7 Hz, 4H, H^{B4}), 7.62 (dd, J = 7.7, 1.1 Hz, 4H, H^{A4}), 6.90 (ddd, J = 7.3, 5.7, 1.4 Hz, 4H, H^{B5}), 6.69 (pseudo-t, J = 7.8 Hz, 4H, H^{A5}), 5.88 (dd, $J = 7.9, 1.1 \text{ Hz}, 4\text{H}, \text{H}^{\text{A5}}), 3.22 \text{ (s, 12H, H}^{\text{Me}}). \text{ IR (solid, } \tilde{\nu}/\text{cm}^{-1})$ 2928 (w), 1604 (w), 1561 (w), 1474 (m), 1422 (w), 1395 (w), 1290 (m), 1274 (m), 1256 (m), 1154 (m), 1129 (s), 1067 (m), 961 (m), 786 (m), 755 (m), 736 (m), 722 (m), 709 (m), 646 (w), 586 (w), 573 (w), 551 (w), 537 (s), 527 (s), 488 (s), 465 (m). LC-ESI-MS m/z 657.1 $[Ir(C^N)_2]^+$ (calc. 657.1), 698.1 $[Ir(C^N)_2(MeCN)]^+$

(calc. 698.1), 739.0 $[Ir(C^N)_2(MeCN)_2]^+$ (calc. 739.1). Found C 40.62, H 3.28, N 4.08; $C_{48}H_{40}Cl_2Ir_2N_4O_8S_4\cdot 2H_2O$ requires C 40.59, H 3.12, N 3.94%.

General procedure for the synthesis of iridium(III) complexes

Iridium dimers and ancillary ligand were suspended in MeOH (15 mL) in a microwave vial and heated at 120 °C for 1 h in a microwave reactor (14 bar). The resulting yellow solution was filtered through cotton and concentrated under reduced pressure. The residue was dissolved in a little MeOH, an excess of solid $\mathrm{NH_4PF_6}$ was added and the resulting suspension was stirred for 5 min at room temperature. The yellow precipitate was filtered off and redissolved in $\mathrm{CH_2Cl_2}$. The solvent was removed under reduced pressure, the crude product was purified by column chromatography (silica) and the solvent was removed under reduced pressure. The residue was dissolved in a little $\mathrm{CH_2Cl_2}$, precipitated with $\mathrm{Et_2O}$ and left in the fridge overnight. The resulting precipitation was filtered off, washed with MeOH and $\mathrm{Et_2O}$ and dried under vacuum.

 $[Ir(2)_2(bpy)][PF_6]$. $[Ir(2)_2Cl]_2$ (101 mg, 0.0730 mmol) and bpy (36.4 mg, 0.233 mmol). Purification by column chromatography (silica, CH₂Cl₂ changing to CH₂Cl₂-4% MeOH) and precipitation from a CH₂Cl₂ solution. [Ir(2)₂(bpy)][PF₆] was isolated as a pale yellow solid (63.2 mg, 0.146 mmol, 45.2%). ¹H NMR (500 MHz, CD₃CN) δ /ppm 8.53 (pseudo-dt, J = 8.3, 1.0 Hz, 2H, H^{E3}), 8.30 (d, J = 2.0 Hz, 2H, H^{A3}), 8.26 (pseudo-dt, J =8.2, 1.0 Hz, 2H, H^{B3}), 8.16 (pseudo-td, J = 8.0, 1.6 Hz, 2H, H^{E4}), 7.96 (pseudo-td, J = 7.9, 1.5 Hz, 2H, H^{B4}), 7.90 (ddd, J = 5.3, 1.5, 0.9 Hz, 2H, H^{E6}), 7.66 (pseudo-dt, J = 5.7, 1.2 Hz, 2H, H^{B6}), 7.52 (ddd, J = 7.7, 5.5, 1.2 Hz, 2H, H^{E5}), 7.39 (dd, J = 8.0, 2.0 Hz, 2H, H^{A5}), 7.17 (ddd, J = 7.4, 5.8, 1.4 Hz, 2H, H^{B5}), 6.54 (d, $J = 8.0 \text{ Hz}, 2H, H^{A6}), 3.04 \text{ (s, 6H, } H^{A4-SO_2Me}). ^{13}C\{^1H\} \text{ NMR}$ (126 MHz, CD₃CN) δ /ppm 166.3 (C^{B2}), 159.7 (C^{A1}), 156.5 (C^{E2}), 151.8 (C^{E6}), 150.6 (C^{B6}), 146.3 (C^{A2}), 140.8 (C^{E4}), 140.3 (C^{B4}), 136.7 (C^{A4}), 133.4 (C^{A6}), 129.6 (C^{E5}), 129.0 (C^{A5}), 125.9 (C^{B5}), 125.8 (CE3), 124.1 (CA3), 122.0 (CB3), 44.6 (CA4-SO₂Me). IR (solid, $\tilde{\nu}/\text{cm}^{-1}$) 3041 (w), 1609 (w), 1579 (w), 1480 (w), 1448 (w), 1426 (w), 1401 (w), 1300 (m), 1246 (w), 1225 (w), 1146 (s), 1096 (w), 1067 (w), 1056 (w), 1032 (m), 960 (m), 840 (s), 784 (m), 760 (s), 734 (m), 703 (w), 640 (w), 594 (m), 557 (s), 524 (m), 485 (m). UV/Vis (CH₃CN, $1.0 \times 10^{-5} \text{ mol dm}^{-3}$) $\lambda/\text{nm} (\varepsilon/\text{dm}^3 \text{ mol}^{-1} \text{ cm}^{-1})$ 253 (62 000), 271 sh (48 000), 313 sh (19 000), 337 sh (9100), 406 sh (2800). Emission (CH₃CN, 1.0×10^{-5} mol dm⁻³, $\lambda_{\rm exc}$ = 271 nm): $\lambda_{\text{em}}^{\text{max}} = 517$ nm. ESI-MS m/z 813.4 $[M - PF_6]^+$ (calc. 813.1). Found C 41.62, H 3.10, N 6.03; C₃₄H₂₈F₆IrN₄O₄PS₂·H₂O requires C 41.84, H 3.10, N 5.74%.

[Ir(3)₂(bpy)][PF₆]. [Ir(3)₂Cl]₂ (114 mg, 0.0823 mmol) and bpy (28.5 mg, 0.182 mmol). Purification by column chromatography (silica, CH₂Cl₂ changing to CH₂Cl₂-2% MeOH) and precipitation from a CH₂Cl₂ solution. [Ir(3)₂(bpy)][PF₆] was isolated as a dark yellow solid (97.5 mg, 0.102 mmol, 61.8%). ¹H NMR (500 MHz, CD₃CN) δ /ppm 9.30 (ddd, J = 8.6, 1.3, 0.8 Hz, 2H, H^{B3}), 8.54 (pseudo-dt, J = 8.3, 1.1 Hz, 2H, H^{E3}), 8.16 (ddd, J = 7.7, 1.5 Hz, 2H, H^{E4}), 7.98 (ddd, J = 8.7, 7.5, 1.7 Hz, 2H, H^{B4}), 7.89 (dd, J = 7.8, 1.2 Hz, 2H, H^{A4}), 7.85 (ddd, J = 5.5, 1.6, 0.7 Hz, 2H, H^{E6}), 7.76 (ddd, J = 5.8, 1.7, 0.7 Hz, 2H, H^{B6}),

7.51 (ddd, J = 7.7, 5.5, 1.2 Hz, 2H, H^{E5}), 7.14 (ddd, J = 7.3, 5.8, 1.3 Hz, 2H, H^{B5}), 7.07 (pseudo-t, J = 7.7 Hz, 2H, H^{A5}), 6.47 (dd, $J = 7.7, 1.2 \text{ Hz}, 2H, H^{A6}, 3.25 \text{ (s, 6H, } H^{A3-SO_2Me}). ^{13}C{^1H} \text{ NMR}$ (126 MHz, CD₃CN) δ /ppm 164.6 (C^{B2}), 156.5 (C^{E2}), 155.6 (C^{A1}), 151.5 (C^{E6}), 151.2 (C^{B6}), 141.1 (C^{A2}), 140.8 (C^{E4}), 140.6 (C^{A3}), 139.9 (C^{B4}), 137.7 (C^{A6}), 130.4 (C^{A5}), 129.6 (C^{E5}), 127.7 (C^{B3}), 126.0 (CB5), 125.9 (CE3), 125.5 (CA4), 43.5 (CA3-SO₂Me). IR (solid, $\tilde{\nu}/\text{cm}^{-1}$) 3124 (w), 3039 (w), 2933 (w), 1609 (w), 1563 (w), 1477 (m), 1449 (w), 1411 (w), 1396 (w), 1308 (m), 1278 (m), 1243 (w), 1200 (w), 1170 (w), 1155 (m), 1131 (m), 1113 (m), 1070 (w), 1045 (w), 1031 (w), 1001 (w), 962 (m), 904 (w), 876 (w), 835 (s), 803 (m), 792 (m), 760 (s), 752 (s), 735 (m), 724 (m), 716 (m), 667 (w), 647 (w), 581 (w), 556 (s), 525 (s), 482 (s). UV/Vis (CH₃CN, $1.0 \times 10^{-5} \text{ mol dm}^{-3}$) $\lambda/\text{nm} (\varepsilon/\text{dm}^3 \text{ mol}^{-1} \text{ cm}^{-1}) 259$ (46 000), 300 sh (28 000), 310 sh (25 000), 360 sh (8000), 397 (5400), 437 sh (3300). Emission (CH₃CN, 1.0×10^{-5} mol dm⁻³, $\lambda_{\rm exc} = 400$ nm): $\lambda_{\rm em}^{\rm max} = 506$, 527 nm. ESI-MS m/z 813.4 [M-zPF₆]⁺ (calc. 813.1). Found C 42.49, H 3.24, N 6.08; C₃₄H₂₈F₆Ir-N₄O₄PS₂ requires C 42.63, H 2.95, N 5.85%.

Crystallography

Single crystal data were collected on a Bruker APEX-II diffractometer; for data reduction, solution and refinement, APEX was used.³⁷ For structural analysis, Mercury v. 3.5.1 was used.³⁸

2{[Ir(1)₂(bpy)][PF₆]}·7CH₂Cl₂. C₇₅H₇₀Cl₁₄F₁₂Ir₂N₈O₈P₂S₄, M = 2510.39, needle, orthorhombic, space group *Pbca*, a = 18.1495(15), b = 22.5027(19), c = 23.189(2) Å, U = 9470.7(8) Å³, Z = 4, $D_c = 1.761$ Mg m⁻³, μ (Mo-Kα) = 3.403 mm⁻¹, T = 123 K. Total 72 763 reflections, 14 677 unique, $R_{\rm int} = 0.045$. Refinement of 14 611 reflections (661 parameters) with $I > 2\sigma(I)$ converged at final $R_1 = 0.0524$ (R_1 all data = 0.0871), w $R_2 = 0.1260$ (w R_2 all data = 0.1539), gof = 1.0026. CCDC 1421913.

2{[Ir(2)₂(bpy)][PF₆]}·5.5H₂O. $C_{68}H_{67}F_{12}Ir_2N_8O_{13.50}P_2S_4$, M=2014.95, yellow block, monoclinic, space group $P2_1/c$, a=20.6753(13), b=20.9141(13), c=18.3265(12) Å, $\beta=104.473(2)^\circ$, U=7673.0(5) Å³, Z=4, $D_c=1.744$ Mg m⁻³, $\mu(\text{Cu-K}\alpha)=8.865$ mm⁻¹, T=123 K. Total 105 992 reflections, 13 988 unique, $R_{\text{int}}=0.041$. Refinement of 13 925 reflections (959 parameters) with $I>2\sigma(I)$ converged at final $R_1=0.0422$ (R_1 all data = 0.0458), w $R_2=0.1120$ (w R_2 all data = 0.1143), gof = 0.9557. CCDC 1421914.

[Ir(3)₂(bpy)][PF₆]. $C_{34}H_{28}F_6$ IrN₄O₄PS₂, M=957.93, yellow needle, monoclinic, space group C2/c, a=15.6899(16), b=28.163(3), c=8.4211(9) Å, $\beta=115.797(3)^\circ$, U=3350.3(4) Å³, Z=4, $D_c=1.899$ Mg m⁻³, μ (Cu–K α) = 10.052 mm⁻¹, T=123 K. Total 18 964 reflections, 3003 unique, $R_{\rm int}=0.029$. Refinement of 3003 reflections (236 parameters) with $I>2\sigma(I)$ converged at final $R_1=0.0222$ (R_1 all data = 0.0223), w $R_2=0.0486$ (w R_2 all data = 0.0486), gof = 1.0000. CCDC 1421915.

Computational details

DFT calculations were carried out with the D.01 revision of the Gaussian 09 program package³⁹ using Becke's three-parameter B3LYP exchange–correlation functional^{40,41} together with the 6-31G** basis set for C, H, N, S and O⁴² and the "double- ζ " quality LANL2DZ basis set for the Ir element.⁴³ An effective

Dalton Transactions

core potential (ECP) replaces the inner core electrons of Ir leaving the outer core $[(5s)^2(5p)^6]$ electrons and the $(5d)^6$ valence electrons of Ir(III). The geometries of the singlet ground state (S₀) and those of the lowest-energy triplet states (T₁ to T₃) were fully optimized without imposing any symmetry restriction. The geometry of the triplets was calculated at the spin-unrestricted UB3LYP level with a spin multiplicity of three. Phosphorescence emission energies were estimated as the vertical difference between the energy of the minimum of the lowest-energy triplet state and the energy of S₀ at the T₁ optimized geometry. The calculation of the energy of S₀ at that geometry was performed as an equilibrium single-point calculation with respect to the solvent reaction field/solute electronic density polarization process. All the calculations were performed in the presence of the solvent (acetonitrile). Solvent effects were considered within the self-consistent reaction field (SCRF) theory using the polarized continuum model (PCM) approach. 44-46 Time-dependent DFT (TD-DFT) calculations of the lowest lying singlets and triplets were performed in the presence of the solvent at the minimum-energy geometry optimized for the ground state.

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. An Ambios XP-1 profilometer was used to determine the film thickness. First, 80 nm of poly(3,4ethylenedioxythiophene):poly(styrenesulfonate) (PEDOT:PSS) (CLEVIOSTM P VP AI 4083, aqueous dispersion, 1.3-1.7% solid content, Heraeus) was coated in order to avoid the formation of pinholes and to improve the reproducibility of the cells. Subsequently, the emitting layer (100 nm) was deposited by spin-coating from a MeCN solution of the emitting compound with the addition of the ionic liquid 1-butyl-3-methylimidazolium hexafluoridophosphate [Bmim][PF₆] (>98.5%, Sigma-Aldrich) in a 4 to 1 molar ratio. 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 luminance versus time by using 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 time-on time and dividing by the total cycle time. The average luminance is directly obtained by taking the average of the obtained photodiode results and correlating it to the value of a luminance meter. The current efficiency is obtained by

dividing the average luminance by the average current density. The electroluminescence (EL) spectra were measured using an Avantes AvaSpec-2048 fiber optic spectrometer during device lifetime measurements.

Results and discussion

Ligand synthesis and characterization

Cyclometallating ligands 2-(4-methylsulfonylphenyl)pyridine (H1) and 2-(3-methylsulfonylphenyl)pyridine (H2)²³ were prepared in three steps starting from fluorophenylboronic acid and 2-bromopyridine. The resulting fluorophenylpyridines were converted into the corresponding methylsulfonylphenylpyridines via nucleophilic aromatic substitution with sodium thiomethoxide and subsequent oxidation with H₂O₂/Na₂WO₄·2H₂O. The synthesis and characterization of these ligands has already been reported.22

2-(2-Methylsulfonylphenyl)pyridine (H3) was synthesized according to a method described in the literature.³⁶ Borylation of 2-phenylpyridine with BBr₃ 35,36 was followed by a coppercatalysed coupling reaction with sodium methanesulfinate. ¹H NMR spectroscopic data of the borylated intermediate and H3 matched those reported. 35,36

Synthesis and characterization of [Ir(C^N)2Cl]2 dimers

Dimers $[Ir(1)_2Cl]_2$ and $[Ir(2)_2Cl]_2$ were prepared from [Ir(cod)Cl]₂ and the corresponding cyclometallating ligand under reflux conditions and their synthesis and characterization have been reported previously. 22,23

For [Ir(3)2Cl]2, however, the microwave-assisted reaction of IrCl₃·xH₂O and H3 in a mixture of 2-ethoxyethanol and water proved to be more successful (Scheme 2). The dimer was isolated as an orange solid by filtration and proved to be sufficiently pure for subsequent reactions, as determined by ¹H NMR spectroscopy.

[Ir(3)₂Cl]₂ was characterized by ¹H NMR and IR spectroscopy and elemental analysis. The base peak at m/z 657.1 in the LC-ESI mass spectrum of a methanolic solution of $[Ir(3)_2Cl]_2$ corresponds to the $[Ir(3)_2]^+$ ion. Further peaks at m/z698.1 and 739.0 were assigned to $[Ir(3)_2(MeCN)]^+$ and [Ir(3)₂(MeCN)₂]⁺, respectively. The acetonitrile originates from

Scheme 2 Synthetic route to [Ir(3)₂Cl]₂. Reaction conditions: (a) IrCl₃·xH₂O, 2-ethoxyethanol/H₂O (3:1), N₂, MW, 1.5 h, 110 °C. Dimer structure including the numbering scheme for NMR spectroscopic assignments.

Dalton Transactions Paper

the eluent of the LC column; this has already been observed for analogous dimers.²²

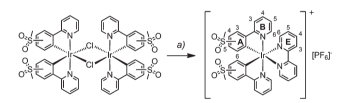
Synthesis and characterization of [Ir(C^N)2(bpy)][PF6] complexes

The synthesis and characterization of [Ir(1)₂(bpy)][PF₆] have been reported previously.²² The complexes [Ir(2)₂(bpy)][PF₆] and [Ir(3)2(bpy)][PF6] were prepared in a similar manner by reactions of [Ir₂(2)₄Cl₂] or [Ir₂(3)₄Cl₂] with 2,2'-bipyridine in MeOH in a microwave reactor (Scheme 3).20 Counterion exchange with NH₄PF₆ yielded [Ir(2)₂(bpy)][PF₆] and [Ir(3)₂(bpy)][PF₆] as yellow solids in moderate yields. The products were characterized by 1H and 13C NMR spectroscopy, electrospray ionization mass spectrometry and elemental

In the ESI mass spectrum of both [Ir(2)₂(bpy)][PF₆] and $[Ir(3)_2(bpy)][PF_6]$, the base peak at m/z 813.4 corresponded to the [Ir(C^N)2(bpy)]+ cation with the typical iridium isotope pattern. 1H and 13C NMR signals were assigned using 2D methods (COSY, HMQC and HMBC). In Fig. 1, the ¹H NMR spectrum of [Ir(3)₂(bpy)][PF₆] in CD₃CN solution is shown as a representative example.

Crystal structures

2{[Ir-Single crystals of $2\{[Ir(1)_2(bpy)][PF_6]\}\cdot 7CH_2Cl_2,$ $(2)_2(bpy)$ [PF₆] $\cdot 5.5H_2O$ and [Ir(3)₂(bpy)] [PF₆] were obtained by layering a CH₂Cl₂ solution with Et₂O, an MeCN solution with



Scheme 3 Synthetic route to iridium complexes [Ir(C^N)₂(bpy)][PF₆] with $C^N = [1]^-$ to $[3]^-$. Reaction conditions: (a) bpy, MeOH, MW, 1 h, 120 °C; then NH₄PF₆. Complex structures including the numbering scheme for NMR spectroscopic assignments.

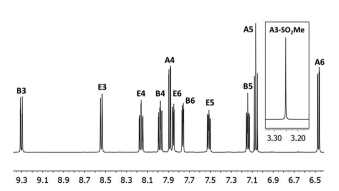


Fig. 1 500 MHz ¹H NMR spectrum of complex [Ir(3)₂(bpy)][PF₆] in CD₃CN with signal assignments. See Scheme 3 for proton labelling. Scale: δ /ppm.

tert-butyl methyl ether (t-BME) and an MeCN solution with Et₂O, respectively. 2{[Ir(1)₂(bpy)][PF₆]}-7CH₂Cl₂ crystallizes in the orthorhombic space group Pbca, whereas 4{[Ir- $(2)_2(bpy)$ [PF₆] $\cdot 11H_2O$ and [Ir(3)₂(bpy)] [PF₆] crystallize in the monoclinic space groups $P2_1/c$ and C2/c, respectively. Structures of the complex cations are shown in Fig. 2-4; selected bond lengths and angles are reported in the figure captions.

In $2\{[Ir(1)_2(bpy)][PF_6]\}\cdot7CH_2Cl_2$, the solvent molecules are heavily disordered and have been modelled over four positions with fractional occupancies. Two independent cations (both are Λ enantiomers) and two ordered $[PF_6]^-$ (one of half occupancy) are present in the asymmetric unit of $2\{[Ir(2)_2(bpy)][PF_6]\}\cdot 5.5H_2O$. In addition, the $[PF_6]^-$ ion containing atom P88 resides on a special position, leading to the ion being shared equally between two unit cells. The asymmetric unit of [Ir(3)2(bpy)][PF6] contains half a cation and half an anion. The second half is generated by a C_2 axis through the central Ir atom, parallel to the b axis of the unit cell.

All the complexes crystallize in achiral space groups, with both enantiomers present in the unit cell.

The bpy ligand is nearly planar in 2{[Ir(1)₂(bpy)][PF₆]}-7CH₂Cl₂, with an angle of 3.0° between the pyridyl ring planes. By moving the sulfone group from the 5- to 4-position, the deviation from planarity becomes larger, with angles between the ring planes of 6.4 and 7.8° in the two independent cations in $2\{[Ir(2)_2(bpy)]\}$ [PF₆]]·5.5H₂O. In [Ir(3)₂(bpy)][PF₆], with the sulfone substituent in the 3-position, the bpy ligand is substantially twisted, with an angle of 13.6° between the pyridine ring planes.

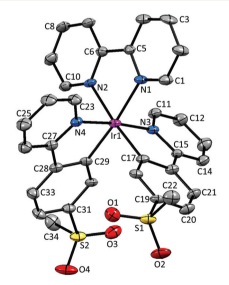


Fig. 2 Structure of the Λ - $[lr(1)_2(bpy)]^+$ cation in $2\{[lr(1)_2(bpy)]$ [PF₆]}-7CH₂Cl₂; ellipsoids are plotted at the 40% probability level and H atoms, [PF₆] counterion and solvent molecules are omitted for clarity. Selected bond lengths and angles: Ir1-N1 = 2.133(5), Ir1-N2 = 2.134(4), Ir1-N3 = 2.044(5), Ir1-N4 = 2.057(5), Ir1-C17 = 2.029(5), Ir1-C29 = 1.029(5)2.009(5), C22-S1 = 1.748(7), S1-O1 = 1.435(5), S1-O2 = 1.440(5), S2-O3= 1.437(6), S2-O4 = 1.451(6), C34-S2 = 1.757(7) Å; N1-Ir1-N2 = 77.53(19), N3-Ir1-C17 = 80.70(19), N4-Ir1-C29 = 80.53(19), N3-Ir1-N4 = 80.70(19)174.41(18), N1-Ir1-C29 = 174.1(2), N2-Ir1-C17 = 174.4(2), O1-S1-O2 = 118.2(3), $O3-S2-O4 = 118.2(4)^\circ$.

C12 C85 CC5

C11 C8 C7

C10 N9 N2 C3

C17 N14 N29

C19 C21 C32 C31 C41

C20 C23 C34 C41

Dalton Transactions Paper

Fig. 3 Structure of one of the independent Λ -[Ir(2)₂(bpy)]⁺ cations in the asymmetric unit in 2{[Ir(2)₂(bpy)][PF₆]}·5.5H₂O; ellipsoids are plotted at the 40% probability level and H atoms, [PF₆]⁻ counterion and solvent molecules are omitted for clarity. Selected bond lengths and angles: Ir1-N2 = 2.134(4), Ir1-N9 = 2.134(4), Ir1-N14 = 2.045(4), Ir1-N29 = 2.045(4), Ir1-C21 = 2.000(4), Ir1-C32 = 2.006(4), S1-C13 = 1.719(10), S1-O27 = 1.454(6), S1-O28 = 1.415(5), S2-O38 = 1.431(4), S2-O39 = 1.431(4), C40-S2 = 1.761(6) Å; N2-Ir1-N9 = 76.72(16), N14-Ir1-C21 = 80.67(16), N29-Ir1-C32 = 80.61(16), N14-Ir1-N29 = 172.70(14), N2-Ir1-C21 = 175.99(16), N9-Ir1-C32 = 173.13(16), O27-S1-O28 = 114.8(4), O38-S2-O39 = 117.5(3)°.

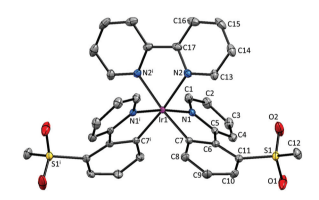


Fig. 4 Structure of the Λ -[Ir(3)₂(bpy)]⁺ cation in [Ir(3)₂(bpy)][PF₆]; ellipsoids are plotted at the 40% probability level and H atoms and [PF₆]⁻ counterion are omitted for clarity. Symmetry code: $i=1-x,y,\frac{1}{2}-z$. Selected bond lengths and angles: Ir1–N1 = 2.036(2), Ir1–N2 = 2.146(2), Ir1–C7 = 2.007(2), S1–O1 = 1.432(2), S1–O2 = 1.433(2), C12–S1 = 1.763(3) Å; N1–Ir1–C7 = 80.08(9), N2–Ir1–N2ⁱ = 76.83(11), N2–Ir1–C7ⁱ = 173.07(9), N1–Ir1–N1ⁱ = 173.67(11), O1–S1–O2 = 118.38(18)°.

The presence of intramolecular $CH_{aryl}\cdots OS$ hydrogen bonds has been observed in a series of alkyl-aryl⁴⁷ and diaryl sulfones, ^{47,48} as well as in ligand H1, ²² dimer $[Ir(1)_2Cl]_2$ ²² and the related complex $[Ir(2)_2(dmpzpy)][PF_6]$ (dmpzpy = 2-(3,5-dimethyl-1*H*-pyrazol-1-yl)pyridine). ²³ The sulfone groups in the $[Ir(1)_2(bpy)]^+$ cation are twisted with respect to the phenyl ring to which they are attached with torsion angles of 14.8° (O3–S2–C31–C30), -34.5° (O4–S2–C31–C32), 20.8° (O1–S1–C19–C18)

and -30.2° (O2–S1–C19–C20). This leads to an optimization of intramolecular CH···O contacts in the range 2.55–2.70 Å. In $[Ir(2)_2(bpy)]^+$, a similar arrangement of the sulfone groups is observed, leading to CH···O contacts ranging from 2.50 to 2.80 Å. In contrast, in $[Ir(3)_2(bpy)]^+$, steric hindrance leads to a different orientation of the SO₂Me group, resulting in interactions between the O atoms of the sulfone group and the adjacent pyridyl ring, with distances of 2.49 Å (O1···HC4) and 2.77 Å (O2···HC4). Due to the steric hindrance, the ppy cyclometallating ligand is severely distorted, giving rise to an angle of 22.2° between the ring planes. In both $[Ir(1)_2(bpy)]^+$ and $[Ir(2)_2(bpy)]^+$, no such strong deviation from planarity is observed and the corresponding angles lie in the range 2.0–6.2°.

Photophysical properties

UV/Vis absorption spectra of the complexes [Ir(C^N)_2(bpy)][PF_6] with C^N = [1]^- to [3]^- in MeCN solution are shown in Fig. 5. All the compounds have relatively similar absorption spectra with maxima in the UV region, between 250 and 260 nm. The bands in the UV region are intense and can be assigned to spin-allowed $\pi \to \pi^*$ transitions centred on the ligands. The less intense bands at higher wavelengths up to 450 nm correspond to metal-to-ligand (¹MLCT) and ligand-to-ligand charge transfer (¹LLCT) transitions.³ The lowest-energy tails of the absorption spectrum of [Ir(3)_2(bpy)][PF_6] extend more in the visible than those of [Ir(1)_2(bpy)][PF_6], and these more than those of [Ir(2)_2(bpy)][PF_6] (Fig. 5).

Photoluminescence spectra shown in Fig. 6 were obtained by excitation of MeCN solutions of compounds $[Ir(C^N)_2(bpy)][PF_6]$ $(C^N = [1]^-$ to $[3]^-$). The emission profiles are independent of the excitation wavelength. Emission occurs from the lowest-lying triplet state (T_1) , which consists of contributions from charge transfer $(^3MLCT$ and $^3LLCT)$ and ligand-centred (^3LC) triplet states in complexes of the type $[Ir(C^N)_2(N^N)]^+$. While $[Ir(2)_2(bpy)][PF_6]$ has a broad and

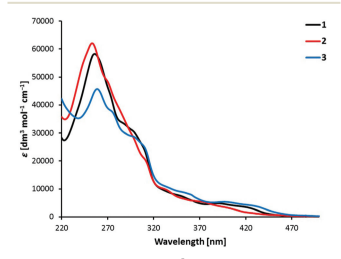


Fig. 5 Absorption spectra of 1×10^{-5} M MeCN solutions of complexes $[Ir(C^N)_2(bpy)][PF_6]$ with $C^N = [1]^-$ to $[3]^-$.

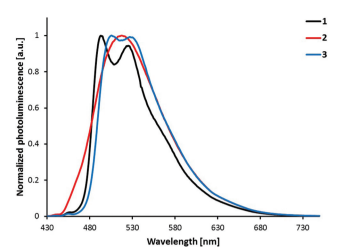


Fig. 6 Photoluminescence spectra of complexes $[Ir(C^N)_2(bpy)][PF_6](C^N = [1]^- \text{ to } [3]^-)$, 1×10^{-5} M in MeCN solution. Excitation wavelength:

unstructured emission profile, indicating a higher charge transfer character, both $[Ir(1)_2(bpy)][PF_6]$ and $[Ir(3)_2(bpy)][PF_6]$ show vibrational structures in their emission profiles, indicating larger 3LC character of the emissive state. This assumption is further supported by the radiative decay rate constants ($k_r = PLQY/\tau$) of the compounds (Table 1). In general, the higher the rate constant, the smaller the contribution of the 3LC state to the emissive triplet state. 3 In this series of complexes, $[Ir(2)_2(bpy)][PF_6]$ has a k_r (7.2 × 10 5 s $^{-1}$) more than double that of the other two complexes (3.2 and 2.6 × 10 5 s $^{-1}$). This observation is therefore in accordance with the supposed higher charge transfer character of the emissive state of $[Ir(2)_2(bpy)][PF_6]$ when compared to complexes $[Ir(1)_2(bpy)][PF_6]$ and $[Ir(3)_2(bpy)][PF_6]$.

All three complexes are green emitters in solution (Fig. 7) and their emission maxima are blue-shifted by 68 to 92 nm when compared to the non-substituted parent complex $[Ir(ppy)_2(bpy)]^+$ (585 nm).²⁴ Looking at the substitution position of the sulfonyl group, complex $[Ir(1)_2(bpy)][PF_6]$ shows the largest blue-shift of the emission maximum, while $[Ir(2)_2(bpy)][PF_6]$ has the smallest. The influence of the sulfone position on the emission maximum therefore follows the

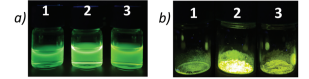


Fig. 7 Photographs of complexes $[Ir(C^N)_2(bpy)][PF_6]$ with $C^N = [1]^-$ to $[3]^-$ under 366 nm light excitation: (a) in MeCN, (b) as powder samples.

trend: 5-position > 3-position > 4-position. However, the different shapes of the emission bands, namely the broad emission of $[Ir(2)_2(bpy)][PF_6]$ with just one maximum, make comparison difficult. In fact, when comparing the emission spectra of $[Ir(2)_2(bpy)][PF_6]$ and $[Ir(3)_2(bpy)][PF_6]$ (Fig. 6), the bands look very similar, with $\lambda_{\rm em}^{\rm max}$ of $[Ir(2)_2(bpy)][PF_6]$ (517 nm) approximately in the middle of the two maxima of $[Ir(3)_2(bpy)][PF_6]$ (506, 527 nm). The emission maxima of $[Ir(1)_2(bpy)][PF_6]$ (493, 525 nm) are definitely shifted to the blue; however, the emission band of $[Ir(2)_2(bpy)][PF_6]$ extends further into the high energy region than those of the other two complexes.

Most of the regioisomeric iridium complexes investigated so far have shown that a substituent in the 4-position has the largest influence on the emission colour. The largest blue-shift has been observed on introducing an electron-withdrawing group in the 4-position 15,26,28,30,33 and the largest red-shift on the introduction of an electron-donating substituent in the 4-position.^{25,27} However, Bronstein et al. have reported a different trend in the emission maximum of their bromo- and fluorenyl-substituted isomers.³¹ For both series, they showed that the largest influence was exerted by a substituent in the 5-position of the phenyl ring. This observation is in accordance with the trend in emission maxima in the present series of regioisomeric complexes. Concerning the nature of the triplet emissive state, the present data can be compared with a series of SF₅-functionalized [Ir(C^N)₂(N^N)]⁺ complexes.³³ In both series, moving the substituent in the phenyl ring of the C^N ligand from the 5- to 4-position (meta to para to the Ir-C bond) leads to a decrease in 3LC and an increase in the CT character of the emissive state, observed by a broadening of

Table 1 Photophysical properties of complexes $[Ir(C^N)_2(bpy)][PF_6]$ ($C^N = [1]^-$ to $[3]^-$) in MeCN solution and as powder samples at room temperature. Quantum yields were measured in de-aerated solution; excited state lifetimes were measured in de-aerated solution under an argon atmosphere. Biexponential fits were used for solid-state lifetime measurements, using the equation $\tau_{ave} = \sum A_i \tau_i / \sum A_i$ (A_i is the pre-exponential factor for the lifetime). $k_r = PLQY/\tau$; $k_{nr} = (1 - PLQY)/\tau$

	MeCN solution				Powder samples				
Complex	$\lambda_{\rm em}^{{ m max}a}$ [nm]	$ au_{1/2}^{\ b} \left[\mu s \right]$	PLQY ^a [%]	$k_{\rm r} \left[10^5 \; {\rm s}^{-1} \right]$	$k_{\rm nr} \left[10^5 \ {\rm s}^{-1} \right]$	$\lambda_{\rm em}^{{ m max}\ c}$ [nm]	$ au_{\mathrm{av}}^{b,d} [\mu \mathrm{s}]$	τ_1 [μ s] (A_1), τ_2 [μ s] (A_2) b,d	PLQY ^a [%]
[Ir(1) ₂ (bpy)][PF ₆] [Ir(2) ₂ (bpy)][PF ₆] [Ir(3) ₂ (bpy)][PF ₆]	493, 525 517 506, 527	2.33 1.28 2.97	74 92 77	3.2 7.2 2.6	1.1 0.63 0.77	535 542 542	0.475 1.19 0.663	0.299 (7035), 1.01 (691) 0.456 (2233), 1.58 (1201) 0.306 (4221), 1.41 (439)	6.6 27 6.1

 $[^]a\lambda_{\rm exc} = 262$ nm for $[{\rm Ir}(1)_2({\rm bpy})][{\rm PF}_6]$, 271 nm for $[{\rm Ir}(2)_2({\rm bpy})][{\rm PF}_6]$ and 261 nm for $[{\rm Ir}(3)_2({\rm bpy})][{\rm PF}_6]$. $^b\lambda_{\rm exc} = 280$ nm. $^c\lambda_{\rm exc} = 340$ nm. d Biexponential fit using the equation $\tau_{\rm av} = \sum A_i \tau_i / \sum A_i$ where A_i is the pre-exponential factor of the lifetime.

the emission spectrum and a shorter excited-state lifetime. Replacement of the bpy ancillary ligand by pyrazolylpyridines in combination with the 4- and 5-substituted sulfonyl cyclometallating ligands gives a different trend, with the *para*-substituted complexes resulting in the largest blue-shift in the emission maximum. ²³ This observation can be explained by a change in the nature of the emissive state, from a larger charge transfer (bpy) to a more ligand-centred character (pzpy), causing a structured emission band and a 54 nm blue-shift. The emission colour is therefore strongly dependent on both cyclometallating and ancillary ligands, making the comparison within the herein presented series difficult due to different contributions of ligand-centred and charge transfer

Dalton Transactions

states to the triplet emissive state.

Photoluminescence quantum yields (PLQY, Table 1) of complexes $[Ir(C^N)_2(bpy)][PF_6]$ with $C^N = [1]^-$ to $[3]^-$ were measured in de-aerated MeCN solution and are relatively high with values between 74 and 92%. While $[Ir(1)_2(bpy)][PF_6]$ (74%) and $[Ir(3)_2(bpy)][PF_6]$ (77%) have similar quantum yields, the PLQY of $[Ir(2)_2(bpy)][PF_6]$ (92%) is significantly higher. Such a trend has also been described by Bronstein $et\ al.;^{31}$ in their series, they observed the highest quantum yield for the complex with a substituent in the 4-position of the cyclometallating ligand. When solutions of compounds $[Ir(C^N)_2(bpy)][PF_6]$ ($C^N = [1]^-$ to $[3]^-$) are not de-aerated, PLQYs are much lower (between 4.5 and 7.0%), indicating strong oxygen quenching.

Excited-state lifetimes in de-aerated MeCN solutions under an argon atmosphere are of the order of a few microseconds, ranging from 1.28 μs for $[Ir(2)_2(bpy)][PF_6]$ to 2.97 μs for $[Ir(3)_2(bpy)][PF_6]$ (Table 1). Again, much shorter excited state lifetimes are observed for the non-de-aerated solutions (0.144 to 0.310 μs), which can be attributed to quenching due to oxygen present in the system.

The photoluminescence spectra shown in Fig. 8 are obtained by the excitation of powder samples of complexes

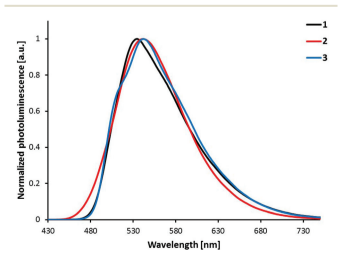


Fig. 8 Photoluminescence spectra of complexes $[lr(C^N)_2(bpy)][PF_6](C^N = [1]^- to [3]^-)$ in the solid state (powder). Excitation wavelength: 340 nm.

 $[Ir(C^N)_2(bpy)][PF_6]$ with $C^N = [1]^-$ to $[3]^-$. The vibrational structure observed in the solution spectra of $[Ir(1)_2(bpy)][PF_6]$ and $[Ir(3)_2(bpy)][PF_6]$ is almost completely lost in the solid state spectra. For all the complexes, powder emission is redshifted compared to solution emission (Fig. 7). The largest red-shift is observed for $[Ir(1)_2(bpy)][PF_6]$ (42 nm), and the smallest for $[Ir(2)_2(bpy)][PF_6]$ (25 nm). In powder samples, the difference between the emission maxima of the complexes is smaller (7 nm) than that in solution (24 nm).

PLQYs of powder samples are significantly lower than solution quantum yields, ranging from 6–7% for $[Ir(1)_2(bpy)][PF_6]$ and $[Ir(3)_2(bpy)][PF_6]$ to 27% for $[Ir(2)_2(bpy)][PF_6]$ (Table 1). As for solution PLQYs, $[Ir(2)_2(bpy)][PF_6]$ has the highest quantum yield. Excited-state lifetimes in powder samples are shorter than those in solution (Table 1); biexponential fits were used for τ of all three complexes. Shorter lifetimes, lower quantum yields and less vibrational structure indicate strong lumine-scence quenching in the solid state due to intermolecular interactions.¹¹

The photoluminescence properties in the amorphous thin film configuration used in LEC devices, where the complex is mixed with the ionic liquid (IL) 1-butyl-3-methylimidazolium hexafluoridophosphate [Bmim][PF₆] in a complex:IL 4:1 molar ratio, were also investigated (Fig. S1†). The band maximum is positioned at 533 nm for $[Ir(1)_2(bpy)][PF_6]$, 548 nm for $[Ir(2)_2(bpy)][PF_6]$ and 539 nm for $[Ir(3)_2(bpy)][PF_6]$. These values are very similar to those observed in powder. However, the photoluminescence intensity is enhanced in the amorphous environment. Whereas the PLQYs of [Ir(1)2(bpy)][PF6] and $[Ir(3)_2(bpy)][PF_6]$ are in the range 11-13%, $[Ir(2)_2(bpy)][PF_6]$ shows a significantly larger PLQY of 45%. These values double the intensity of the photoluminescence emission recorded in powder, which indicates a strong decrease of the luminescence quenching due to the presence of the ionic liquid that decreases the intermolecular interactions between the complexes.

Electrochemical properties

Cyclic voltammetric data in MeCN solution of compounds $[Ir(C^N)_2(bpy)][PF_6]$ with $C^N = [1]^-$ to $[3]^-$ are summarized in Table 2 and cyclic voltammograms are shown in Fig. S2.† In complexes of the type $[Ir(C^N)_2(N^N)][PF_6]$, the first reduction is based on the N^N ligand, while the first oxidation is metal-based with a contribution from the C^N ligands.³² In our series, the first reduction waves are reversible and the poten-

Table 2 Electrochemical data of complexes [Ir(C^N)₂(bpy)][PF₆] (C^N = $[1]^-$ to $[3]^-$) in MeCN solution referenced to Fc/Fc⁺ with 0.1 M [n Bu₄][PF₆] as the supporting electrolyte and a scan rate of 0.1 V s⁻¹ (ir = irreversible, gr = quasi-reversible)

Complex	$E_{1/2}^{\mathrm{ox}}\left[\mathbf{V}\right]$	$E_{1/2}^{\mathrm{red}}\left[\mathbf{V}\right]$	$\Delta E_{1/2}$ [V]
[Ir(1) ₂ (bpy)][PF ₆]	+1.18	$\begin{array}{l} -1.72, -2.16, -2.61^{\mathrm{ir}} \\ -1.67, -2.27^{\mathrm{qr}}, -2.52^{\mathrm{qr}} \\ -1.69, -2.06^{\mathrm{ir}}, -2.16^{\mathrm{ir}}, \\ -2.34^{\mathrm{ir}}, -2.52^{\mathrm{ir}} \end{array}$	2.90
[Ir(2) ₂ (bpy)][PF ₆]	+1.29		2.96
[Ir(3) ₂ (bpy)][PF ₆]	+1.20 ^{qr}		2.89

tials are similar, ranging from -1.67 to -1.72 V (with respect to Fc/Fc⁺). As all three complexes have the same N^N ligand (bpy), this result was expected and is comparable to the reduction potential of [Ir(ppy)₂(bpy)][PF₆] (-1.77 V, in DMF).²⁴ Two to four more reduction waves have been observed for compounds $[Ir(C^N)_2(bpy)][PF_6](C^N = [1]^- \text{ to } [3]^-)$, but were not investigated in detail. A larger influence on the oxidation depending on the substitution position of the sulfone group on the cyclometallating ligand was anticipated and has been observed in our case. Oxidation processes are reversible or quasi-reversible and occur at +1.18, +1.20 and +1.29 V for $[Ir(C^N)_2(bpy)][PF_6]$ with $C^N = [1]^-$, $[3]^-$ and $[2]^-$, respectively. The largest stabilization of the HOMO and the largest influence on the oxidation process are therefore observed for $[Ir(2)_2(bpy)][PF_6]$ with the sulfone substituent in the 4-position. Compared to [Ir(ppy)₂(bpy)][PF₆] (+0.84 V),²⁴ the oxidation waves of all three complexes $[Ir(C^N)_2(bpy)][PF_6](C^N = [1]^-$ to [3] are shifted to more positive potential, consistent with the

Theoretical calculations

blue-shift in the emission maximum.

Paper

To gain a deeper insight into the electrochemical and photophysical properties of complexes $[Ir(C^N)_2(bpy)][PF_6](C^N = [1]^-)$ to $[3]^-)$, a combined DFT/TD-DFT theoretical investigation of their respective cations was undertaken at the B3LYP/(6-31G** + LANL2DZ) level in the presence of the solvent (acetonitrile) (see the Experimental section for full computational details).

The geometry of the complexes in their ground electronic state (S₀) was fully optimized without imposing any symmetry restriction. The calculations reproduce the main trends observed in the experiments. They predict a near octahedral coordination of the Ir metal where the ancillary bpy ligands remain mostly planar, with N-C-C-N dihedral angles of 3.2, 3.5 and 2.8°. In accord with the X-ray structures, the ppy cyclometallating ligands deviate from planarity in passing from $[Ir(1)_2(bpy)]^+$ and $[Ir(2)_2(bpy)]^+$ (inter-ring angle of 0.8–0.9°) to [Ir(3)₂(bpy)]⁺ for which the ring planes form an angle of 12.8°. In $[Ir(1)_2(bpy)]^+$ and $[Ir(2)_2(bpy)]^+$, the SO₂Me groups adopt a conformation in which the methyl is perpendicular to the phenyl ring and the oxygen atoms form CH···O contacts of about 2.60 Å. In $[Ir(3)_2(bpy)]^+$ the SO_2Me groups form closer CH···O contacts of 2.02 and 2.22 Å with adjacent pyridyl and phenyl rings, respectively.

Fig. 9 shows the energy and electron density contours calculated for the highest-occupied (HOMO) and lowest-unoccupied molecular orbitals (LUMO and LUMO+1) of $[Ir(1)_2(bpy)]^+$, $[Ir(2)_2(bpy)]^+$ and $[Ir(3)_2(bpy)]^+$, which are compared with those obtained for the archetypal complex $[Ir(ppy)_2(bpy)]^+$. The topology of the MOs is the same for all three complexes and reproduce that of the MOs of the reference complex $[Ir(ppy)_2(bpy)]^+$. The LUMO+2, which is not displayed in Fig. 9, is close in energy to the LUMO+1 and shows a similar topology.

The introduction of the sulfone groups stabilizes the HOMO by 0.4–0.5 eV because, as expected, this orbital is located on the Ir atom and the phenyl rings of the C^N ligands to which the electron-withdrawing SO₂Me groups are attached

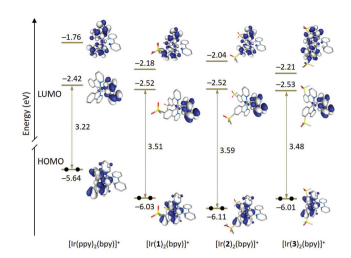


Fig. 9 Schematic representation showing the isovalue contours (± 0.03 a.u.) and energies calculated for the frontier molecular orbitals of $[lr(ppy)_2(bpy)]^+$ and complexes $[lr(C^N)_2(bpy)]^+$ ($C^N = [1]^-$ to $[3]^-$). Hydrogen atoms are omitted.

(Fig. 9). Calculations confirm the larger stabilization of the HOMO by ~ 0.1 eV for complex $[Ir(2)_2(bpy)]^+$ in good agreement with the higher oxidation potential measured for $[Ir(2)_2(bpy)]^+$ (1.29 V) when compared with $[Ir(1)_2(bpy)]^+$ (1.18 V) and $[Ir(3)_2(bpy)]^+$ (1.20 V). The stabilization is larger for $[Ir(2)_2(bpy)]^+$ because the carbon in the 4-position, to which the SO_2Me group is linked, contributes to the HOMO to a higher degree than the carbons in 5- and 3-positions. This enhances the electron-withdrawing effect of the sulfone groups compared with $[Ir(1)_2(bpy)]^+$ and $[Ir(3)_2(bpy)]^+$. A similar result was observed when changing the sulfone position from 5 to 4 in a closely related family of complexes with pyrazolylpyridine N^N ligands.²³

Regarding the unoccupied MOs, the introduction of the sulfone groups especially stabilizes the orbitals localized on the C^N ligands. In this way, the LUMO+1 of [Ir(ppy)₂(bpy)]⁺ (-1.76 eV) decreases in energy by 0.42 and 0.45 eV in passing to $[Ir(1)_2(bpy)]^+$ and $[Ir(3)_2(bpy)]^+$, respectively. The stabilization is smaller for $[Ir(2)_2(bpy)]^+$ (0.28 eV) because of the minor participation of the carbon in the 4-position to which the sulfone group is attached. Similar trends are found for LUMO+2, the partner of LUMO+1 also located on the C^N ligands. The LUMO, which is mainly concentrated on the bpy ligand, undergoes a smaller stabilization of about 0.10 eV for all three complexes (Fig. 9). The energies predicted for the LUMO are in good agreement with similar values recorded for the first reduction potential of $[Ir(C^N)_2(bpy)][PF_6](C^N = [1]^- to [3]^-)$ that are ~0.10 V less negative than that reported for $[Ir(ppy)_2(bpy)]^+$ (-1.77 V) (Table 2).³

The higher HOMO-LUMO gap predicted for $[Ir(2)_2(bpy)]^+$ (3.59 eV) compared with $[Ir(1)_2(bpy)]^+$ (3.51 eV) and $[Ir(2)_2(bpy)]^+$ (3.48 eV) is in accord with the blue shift observed for the lowest energy band of $[Ir(2)_2(bpy)]^+$ in the absorption spectrum (Fig. 5). This band is actually due to a set of electronic transitions involving singlet excited states of

Dalton Transactions Paper

³MLCT/³LLCT nature calculated at ~350 nm, but also to the lower-energy ³LC states located on the sulfone-substituted ppy ligands (Table S1†). The theoretical simulation obtained from the TD-DFT calculation of the singlet excited states (Fig. S6†) correctly predicts the shape of the experimental absorption spectra (Fig. 5).

To investigate the nature of the lowest-energy triplet excited states, a TD-DFT study was first performed for [Ir(C^N)₂(bpy)]⁺ $(C^N = [1]^- \text{ to } [3]^-)$ at the optimized geometry of S_0 . Table 3 compares the excitation energies and electronic nature computed for the three lowest triplet states and includes those obtained for the reference complex $[Ir(ppy)_2(bpy)]^+$. The nature of the three triplets is the same for all four complexes but the energy ordering changes. For $[Ir(ppy)_2(bpy)]^+$, the lowest lying triplet (T_1) results from the HOMO \rightarrow LUMO excitation and therefore implies a charge transfer from the metal and the phenyl rings of the cyclometallating ligands, where the HOMO is located, to the ancillary ligand, where the LUMO resides (see Fig. 9). The T₁ state therefore shows a mixed ³MLCT/³LLCT character. The T_2 and T_3 triplets are well above T_1 by ~ 0.3 eV and mainly imply excitations from the HOMO to the LUMO+1 and LUMO+2 located on the cyclometallating ppy ligands. T₂ and T₃ therefore correspond to ³LC states with some contribution from the metal.

Compared with [Ir(ppy)₂(bpy)]⁺, the energy ordering and electronic nature of the T1 to T3 triplets remain the same for $[Ir(2)_2(bpy)]^+$ but are reversed for $[Ir(1)_2(bpy)]^+$ and $[Ir(3)_2(bpy)]^+$ (Table 3). For the latter, T_1 and T_2 now correspond to the 3LC states and T_3 is the HOMO \rightarrow LUMO 3 MLCT/ 3 LLCT state. The three states are indeed very close in energy (0.06-0.09 eV) for all three complexes. The energy collapse of the three states is due to the larger stabilization of the MOs located on the C^N ligands induced by the sulfone substituents (Fig. 9). As a result, the LUMO+1 and LUMO+2 come closer to the LUMO in $[Ir(C^N)_2(bpy)]^+$ $(C^N = [1]^-$ to $[3]^-)$ than they are in

Table 3 Lowest triplet excited states calculated at the TD-DFT B3LYP/ $(6-31G^{**} + LANL2DZ)$ level for complexes $[Ir(ppy)_2(bpy)]^+$, $[Ir(1)_2(bpy)]^+$, [lr(2)₂(bpy)]⁺ and [lr(3)₂(bpy)]⁺ in acetonitrile solution. Vertical excitation energies (E), dominant monoexcitations with contributions (within parentheses) greater than 20% and description of the excited state are summarized. H and L denote HOMO and LUMO, respectively

Complex	State	$E (eV)/\lambda (nm)$	Monoexcitations	Description
[Ir(ppy) ₂ (bpy)] ⁺	T_1	2.50/496	H → L (98)	³ MLCT/ ³ LLCT
	T_2	2.77/448	$H \rightarrow L+1 (66)$	^{3}LC
	T_3	2.81/441	$H \rightarrow L+2 (54)$	^{3}LC
			$H-1 \to L+1 (26)$	^{3}LC
$[Ir(1)_2(bpy)]^+$	T_1	2.71/457	$H \rightarrow L+1 (70)$	^{3}LC
	T_2	2.76/448	$H \rightarrow L+2 (60)$	^{3}LC
	T_3	2.77/447	$H \rightarrow L (94)$	³ MLCT/ ³ LLCT
$[Ir(2)_2(bpy)]^+$	T_1	2.85/435	$H \rightarrow L (89)$	³ MLCT/ ³ LLCT
	T_2	2.87/432	$H \rightarrow L+1 (57)$	^{3}LC
	T_3	2.91/426	$H \rightarrow L+2 (44)$	^{3}LC
			$H-1 \rightarrow L+1$ (29)	^{3}LC
[Ir(3)2(bpy)] ⁺	T_1	2.67/464	$H \rightarrow L+1 (59)$	^{3}LC
	T_2	2.72/456	$H \rightarrow L+2 (66)$	^{3}LC
	T_3	2.76/449	$H \rightarrow L (77)$	³ MLCT/ ³ LLCT

 $[Ir(ppy)_2(bpy)]^+$, and the T_1 to T_3 states become almost degenerate for the former. The HOMO → LUMO+1, LUMO+2 ³LC triplets are slightly less stable for $[Ir(2)_2(bpy)]^+$ due to the higher stabilization of the HOMO and the lower stabilization of the LUMO+1 (Fig. 9). This combined effect determines that the HOMO-LUMO+1 gap is larger for $[Ir(2)_2(bpy)]^+$ (4.07 eV) than for $[Ir(1)_2(bpy)]^+$ (3.85 eV) and $[Ir(3)_2(bpy)]^+$ (3.80 eV) and, as a consequence, the T_1 state preserves the HOMO \rightarrow LUMO 3 MLCT/ 3 LLCT nature for [Ir(2)₂(bpy)] $^{+}$.

To verify the predicted trends and to obtain additional information about the emitting state, the geometries of the lowest triplet excited states of $[Ir(C^N)_2(bpy)]^+$ (C^N = [1] to [3] and [Ir(ppy)₂(bpy]⁺ were optimized using the spin-unrestricted UB3LYP approach. In all cases, we were able to locate the minimum energy geometries of the three lowest triplet states by carefully selecting the starting point for the optimization process. Fig. 10a summarizes the adiabatic energy difference (ΔE), calculated as the difference between the total energies of So and T1, T2 or T3 at their respective minimumenergy structures, and the emission energy (E_{em}) , estimated as the vertical energy difference between T1 and S0 at the optimized minimum-energy geometry of T1. Fig. 10b shows the unpaired-electron spin density distributions calculated for T₁ to T_3 at their optimized geometries.

The T_1 and T_2 states of $[Ir(1)_2(bpy)]^+$ and $[Ir(3)_2(bpy)]^+$ become degenerate upon full-geometry relaxation and present

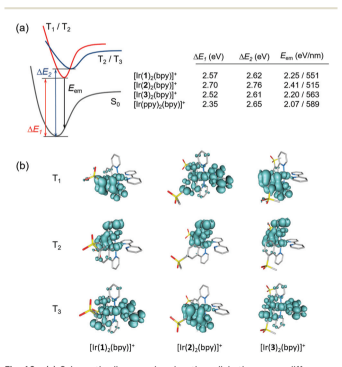


Fig. 10 (a) Schematic diagram showing the adiabatic energy differences $(\Delta E_1, \Delta E_2)$ between S₀ and T₁-T₃ and the emission energy (E_{em}) from T₁ computed for $[lr(C^N)_2(bpy)]^+$ ($C^N = [1]^-$ to $[3]^-$) and $[lr(ppy)_2(bpy)]^+$. The 3 LC triplets $(T_1/T_2 \text{ for } [lr(1)_2(bpy)]^+ \text{ and } [lr(3)_2(bpy)]^+ \text{ and } T_2/T_3 \text{ for }$ $[Ir(2)_2(bpy)]^+$ and $[Ir(ppy)_2(bpy)]^+)$ are degenerate. (b) Unpaired-electron spin density contours (0.002 a.u.) calculated for fully relaxed T_1 , T_2 and T_3 states of $[Ir(C^N)_2(bpy)]^+$ $(C^N = [1]^-$ to $[3]^-$). Hydrogen atoms are omitted.

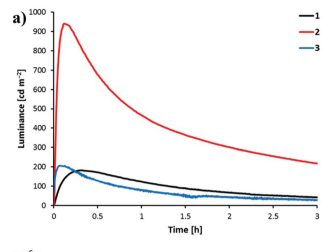
a spin-density distribution mainly centred on one of the cyclometallating ligands (\sim 1.7 unpaired electrons) with a small contribution from the metal (\sim 0.3e) (Fig. 10b). In contrast, $\rm T_3$ features a spin density distribution spreading the ppy-Ir environment and the bpy ligand (Ir \sim 0.5e, C^N ligands \sim 0.5e, N^N ligand \sim 1.0e) that perfectly matches the topology of the HOMO \rightarrow LUMO MLCT/LLCT excitation. Calculations therefore confirm the predominant 3 LC nature of the lowest-energy triplet state of $[\rm Ir(1)_2(bpy)]^+$ and $[\rm Ir(3)_2(bpy)]^+$ in accord with the structured shape of the emission band observed experimentally for these two complexes (Fig. 6).

The T_1 to T_3 states of $[Ir(2)_2(bpy)]^+$ appear at higher energies (Fig. 10a), and their spin-density distributions are reversed with respect to those computed for $[Ir(1)_2(bpy)]^+$ and $[Ir(3)_2(bpy)]^+$ (Fig. 10b). Now, the T₁ triplet corresponds to the ³MLCT/³LLCT state whereas T₂ and T₃ are of ³LC nature. The ³MLCT/³LLCT nature predicted for the lowest-energy triplet is in good agreement with the structureless shape of the emission band registered experimentally for [Ir(2)₂(bpy)]⁺ (Fig. 6). Calculations therefore support the change in the electronic nature of the emitting state from 3LC to 3MLCT/3LLCT in passing from the $[Ir(1)_2(bpy)]^+$ and $[Ir(3)_2(bpy)]^+$ complexes to [Ir(2)₂(bpy)]⁺. The change is due to the different effect the electron-withdrawing sulfone groups have on the energies of the frontier molecular orbitals depending on the phenyl position of the ppy ligands to which they are attached. When introduced in the 4-position, as in [Ir(2)₂(bpy)]⁺, they have a larger stabilization effect on the HOMO and a lower stabilization effect on the LUMO+1 and LUMO+2, and this determines that the emitting triplet corresponds to the HOMO -> LUMO ³MLCT/³LLCT in state as the reference complex $[Ir(ppy)_2(bpy)]^+$. The emission wavelength predicted for [Ir(2)₂(bpy)]⁺ (515 nm) is in good agreement with the $\lambda_{\rm em}^{\rm max}$ measured experimentally in MeCN solution (517 nm).

Electroluminescence

The electroluminescent behaviour of complexes [Ir- $(C^N)_2(bpy)$][PF₆] ($C^N = [1]^-$ to $[3]^-$) was tested by incorporating them into LEC devices. The preparation of the devices consisted in the deposition in air of an 80 nm layer of PEDOT:PSS on top of a patterned indium tin oxide coated glass substrate followed by a 100 nm emitting layer. The emitting layer was prepared by mixing one of the complexes with the ionic liquid [Bmim][PF₆] at a molar ratio of 4:1. After this, the devices were annealed at 100 °C for one hour under an inert atmosphere and the top aluminium contact was deposited by thermal evaporation.

The devices were operated using a block-wave pulsed current driving mode (average current density: 100 A m⁻², frequency: 1000 Hz, duty cycle: 50%). This driving method was selected in order to enhance the device response. Under these conditions, the voltage required to maintain the current density decreases *versus* time due to the formation of p- and n-doped regions, which reduces the resistance of the active layer. The LEC behaviour is depicted in Fig. 11 and the performance parameters are summarized in Table 4. All the LECs prepared



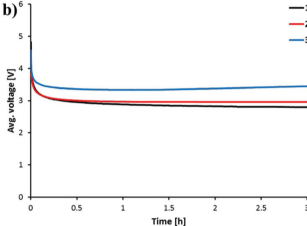


Fig. 11 Luminance (a) and average voltage (b) for ITO/PEDOT:PSS/active layer/Al LEC devices measured by applying a block-wave pulsed current of 100 A $\rm m^{-2}$ at a frequency of 1 kHz and duty cycles of 50%. Active layer: $[Ir(C^N)_2(bpy)][PF_6]$ ($\rm C^N = [1]^-$ to $\rm [3]^-$): $[Bmim][PF_6]$ 4:1 molar ratio.

show a fast decrease of the voltage accompanied by a fast increase of the luminance. A luminance of $100 \text{ cd m}^{-2} (t_{100})$ is reached in 14, 32 and 250 s when complexes $[\text{Ir}(\text{C}^{\,}\text{N})_2(\text{bpy})]$ $[\text{PF}_6]$ with $\text{C}^{\,}\text{N} = [1]^-$, $[2]^-$ and $[3]^-$, respectively, are incorporated into the LEC. Once the maximum luminance (Lum_{max}) is reached, the decay of the luminance leads to short lifetimes (t_2) as has been observed for previously reported LECs based on green emitters under the same driving conditions. Faster LEC shows faster decay of luminance and shorter lifetime. This behaviour can be related with a fast growth of the doped regions, which efficiently quench light-emitting excitons.

Interestingly, the four-times higher PLQY recorded in thin films for $[Ir(2)_2(bpy)][PF_6]$ (45.0%) compared with $[Ir(1)_2(bpy)][PF_6]$ (11.6%) and $[Ir(3)_2(bpy)][PF_6]$ (13.4%) is reflected in the luminance level and the efficiency of the device. The LEC containing $[Ir(2)_2(bpy)][PF_6]$ reaches a Lum_{max} of 940 cd m⁻², whereas LECs incorporating $[Ir(1)_2(bpy)][PF_6]$ and $[Ir(3)_2(bpy)][PF_6]$ reach Lum_{max} values close to 200 cd m⁻². The higher charge transfer character of the emissive state in

Dalton Transactions Paper

Table 4 Performance parameters obtained for ITO/PEDOT:PSS/active layer/Al LEC devices by applying a block-wave pulsed current of 100 A m⁻² at a frequency of 1 kHz and duty cycles of 50%. Active layer: $[lr(C^N)_2(bpy)][PF_6]$ (C^N = [1] to [3] : $[Bmim][PF_6]$ 4:1 molar ratio

Complex	$t_{100}^{a}[s]$	$\operatorname{Lum}_{\operatorname{max}}{}^{b}\left[\operatorname{cd}\operatorname{m}^{-2}\right]$	$t_{1/2}^{c}\left[\mathrm{h}\right]$	Efficacy _{max} d [cd A $^{-1}$]	PCE_{max}^{e} [lm W ⁻¹]	$EQE_{max}^{f}[\%]$
[Ir(1) ₂ (bpy)][PF ₆]	250	182	1.5	1.7	0.9	0.5
$[Ir(2)_2(bpy)][PF_6]$	32	940	1.0	8.8	4.5	2.6
$[Ir(3)_2(bpy)][PF_6]$	14	206	0.7	1.9	0.9	0.6

^a Time to reach 100 cd m⁻². ^b Maximum luminance. ^c Time to reach one-half the maximum luminance. ^d Maximum efficacy. ^e Maximum power conversion efficiency. ^f Maximum external quantum efficiency (EQE).

complex [Ir(2)₂(bpy)][PF₆] therefore leads to a much better LEC performance. LECs with [Ir(2)₂(bpy)][PF₆] show higher efficacy (8.8 cd A^{-1}), power conversion efficiency (PCE = 4.5 lm W^{-1}) and external quantum efficiency (EQE = 2.6%) than those with [Ir(1)₂(bpy)][PF₆] and [Ir(3)₂(bpy)][PF₆] (efficacy < 2 cd A^{-1} , PCE < 1 lm W^{-1} and EQE < 1%) (Table 4). The electroluminescence efficiency obtained for [Ir(2)₂(bpy)][PF₆] is indeed comparable to that obtained for closely related [Ir(msppz)₂(N^N)][PF₆] complexes (msppz = 1-[4-(methylsulfonyl)phenyl]-1*H*-pyrazole) and N^N = bpy-type ligand), which exhibit higher PLQY values in thin films.²⁰ This indicates a better charge-carrier balance in the LECs built with complex [Ir(2)₂(bpy)][PF₆].

The electroluminescence (EL) spectra recorded for the LECs prepared show emission maxima in the 550–560 nm range for all three complexes (Fig. S7†). The emission band is redshifted compared with the photoluminescence spectra in solution, powder and thin film, and all the LEC devices exhibit green colour.

Conclusions

Three regioisomeric iridium(\mathfrak{m}) complexes containing methylsulfonyl-functionalized cyclometallating ligands were prepared to study the effect of the substituent's position on the photophysical, electrochemical and LEC device properties. Structural data for the complexes showed the expected core structures and the influence of the steric hindrance induced by the SO_2Me group in the 3-position of the cyclometallating ligand which leads to ligand distortions.

The complexes are green emitters in solution and green-yellow emitters as powder samples. Vibrationally structured emission bands were observed in MeCN solution for complexes with sulfone groups in the 3- and 5-position of the phenyl ring (*meta* to the Ir–C bond). The two complexes showed similar quantum yields and lifetimes. An enhanced PLQY of 92%, a shorter excited-state lifetime and a broad unstructured emission profile were obtained for the 4-substituted compound (substituent *para* to the Ir–C bond). The lack of a vibrational emission profile suggests a more pronounced charge transfer character of the emissive triplet state compared to the other two complexes. In the solid state (powder), emission maxima are red-shifted, vibrational structure is lost and quantum yields and lifetimes are decreased, indicating excited-state quenching due to intermolecular interactions.

The *para*-substituted complex (4-SO₂Me) again exhibits the highest quantum yields of 27% as powder samples and 45% in thin films. Electrochemical data parallel the photoluminescence trends and show that a methylsulfone substituent in the *para* position to the Ir–C bond has the largest influence on the oxidation potential. DFT calculations rationalize the different effect exerted by the sulfone group depending on the substitution position, and support the experimentally gained results. They confirm the different nature of the emissive triplet state of the *meta*- (³LC) *versus para*-substituted (³MLCT/³LLCT) complexes.

Green electroluminescence with maxima ranging from 550 to 560 nm is observed for LECs with all the complexes. Maximum luminance levels, power conversion efficiencies and EQEs are similar for complexes with the sulfone substituent *meta* to the Ir–C bond. For the complex containing the methylsulfonyl group in the 4-position (para to the Ir–C bond), significantly higher luminance (940 cd m⁻²) and efficiencies (PCE = 4.4 lm W⁻¹ and EQE = 2.6%) are obtained. The increased efficiency correlates with the higher solid state quantum yield. Lifetimes are rather short and comparable for all the devices, ranging from 0.7 to 1.5 h.

We have shown that while the influence on the emission colour is negligible on changing the substitution position of a sulfone group on the cyclometallating ligand, the nature of the emissive triplet state (ligand-centred or charge transfer) is sensitive to the substitution pattern. This leads to significantly enhanced quantum yields in the case of substitution *para* to the Ir–C bond and, as a consequence, superior device performance.

Acknowledgements

We thank the Swiss National Science Foundation (grant number 200020_144500), the Spanish Ministry of Economy and Competitiveness (MINECO) of Spain (MAT2014-55200, CTQ2015-71154-P and Unidad de Excelencia María de Maeztu MDM-2015-0552), the European Research Council (Advanced Grant 267816 LiLo) and the University of Basel for financial support.

Notes and references

J. D. Slinker, D. Bernards, P. L. Houston, H. D. Abruña,
 S. Bernhard and G. G. Malliaras, *Chem. Commun.*, 2003,
 2392–2399.

2 J. D. Slinker, J. Rivnay, J. S. Moskowitz, J. B. Parker, S. Bernhard, H. D. Abruña and G. G. Malliaras, *J. Mater. Chem.*, 2007, 17, 2976–2988.

- 3 R. D. Costa, E. Ortí, H. J. Bolink, F. Monti, G. Accorsi and N. Armaroli, *Angew. Chem., Int. Ed.*, 2012, **51**, 8178–8211.
- 4 T. Hu, L. He, L. Duan and Y. Qiu, *J. Mater. Chem.*, 2012, 22, 4206–4215.
- 5 R. D. Costa, F. Monti, G. Accorsi, A. Barbieri, H. J. Bolink, E. Ortí and N. Armaroli, *Inorg. Chem.*, 2011, **50**, 7229–7238.
- 6 R. D. Costa, E. Ortí, H. J. Bolink, S. Graber, S. Schaffner, M. Neuburger, C. E. Housecroft and E. C. Constable, Adv. Funct. Mater., 2009, 19, 3456–3463.
- 7 P. Pla, J. M. Junquera-Hernández, H. J. Bolink and E. Ortí, *Dalton Trans.*, 2015, 44, 8497–8505.
- 8 M. S. Lowry and S. Bernhard, *Chem. Eur. J.*, 2006, **12**, 7970–7977.
- 9 E. Baranoff, H. J. Bolink, E. C. Constable, M. Delgado, D. Häussinger, C. E. Housecroft, M. K. Nazeeruddin, M. Neuburger, E. Ortí, G. E. Schneider, D. Tordera, R. M. Walliser and J. A. Zampese, *Dalton Trans.*, 2013, 42, 1073–1087.
- F. De Angelis, S. Fantacci, N. Evans, C. Klein,
 S. M. Zakeeruddin, J.-E. Moser, K. Kalyanasundaram,
 H. J. Bolink, M. Grätzel and M. K. Nazeeruddin, *Inorg. Chem.*, 2007, 46, 5989–6001.
- 11 L. He, L. Duan, J. Qiao, R. Wang, P. Wei, L. Wang and Y. Qiu, *Adv. Funct. Mater.*, 2008, **18**, 2123–2131.
- 12 H. J. Bolink, E. Coronado, R. D. Costa, N. Lardiés and E. Ortí, *Inorg. Chem.*, 2008, 47, 9149–9151.
- 13 M. Tavasli, S. Bettington, I. F. Perepichka, A. S. Batsanov, M. R. Bryce, C. Rothe and A. P. Monkman, Eur. J. Inorg. Chem., 2007, 4808–4814.
- 14 G. Zhou, C.-L. Ho, W.-Y. Wong, Q. Wang, D. Ma, L. Wang, Z. Lin, T. B. Marder and A. Beeby, *Adv. Funct. Mater.*, 2008, 18, 499–511.
- 15 R. Ragni, E. Orselli, G. S. Kottas, O. H. Omar, F. Babudri, A. Pedone, F. Naso, G. M. Farinola and L. De Cola, *Chem. Eur. J.*, 2009, **15**, 136–148.
- 16 G. Zhou, X. Yang, W.-Y. Wong, Q. Wang, S. Suo, D. Ma, J. Feng and L. Wang, *ChemPhysChem*, 2011, 12, 2836–2843.
- 17 Y. Hisamatsu and S. Aoki, *Eur. J. Inorg. Chem.*, 2011, 5360–5369.
- 18 C. Fan, Y. Li, C. Yang, H. Wu, J. Qin and Y. Cao, *Chem. Mater.*, 2012, **24**, 4581–4587.
- 19 X. Xu, X. Yang, J. Dang, G. Zhou, Y. Wu, H. Li and W.-Y. Wong, *Chem. Commun.*, 2014, **50**, 2473–2476.
- 20 D. Tordera, A. M. Bünzli, A. Pertegás, J. M. Junquera-Hernández, E. C. Constable, J. A. Zampese, C. E. Housecroft, E. Ortí and H. J. Bolink, *Chem. Eur. J.*, 2013, 19, 8597–8609.
- 21 G. E. Schneider, H. J. Bolink, E. C. Constable, C. D. Ertl, C. E. Housecroft, A. Pertegàs, J. A. Zampese, A. Kanitz, F. Kessler and S. B. Meier, *Dalton Trans.*, 2014, 43, 1961– 1964.
- 22 E. C. Constable, C. D. Ertl, C. E. Housecroft and J. A. Zampese, *Dalton Trans.*, 2014, 43, 5343–5356.

- 23 C. D. Ertl, J. Cerdá, J. M. Junquera-Hernández, A. Pertegás, H. J. Bolink, E. C. Constable, M. Neuburger, E. Ortí and C. E. Housecroft, RSC Adv., 2015, 5, 42815–42827.
- 24 R. D. Costa, E. Ortí, D. Tordera, A. Pertegás, H. J. Bolink, S. Graber, C. E. Housecroft, L. Sachno, M. Neuburger and E. C. Constable, *Adv. Energy Mater.*, 2011, 1, 282–290.
- 25 J. Nishida, H. Echizen, T. Iwata and Y. Yamashita, *Chem. Lett.*, 2005, 34, 1378–1379.
- 26 V. V. Grushin, N. Herron, D. D. LeCloux, W. J. Marshall, V. A. Petrov and Y. Wang, *Chem. Commun.*, 2001, 1494– 1495.
- 27 T.-H. Kwon, H. S. Cho, M. K. Kim, J.-W. Kim, J.-J. Kim, K. H. Lee, S. J. Park, I.-S. Shin, H. Kim, D. M. Shin, Y. K. Chung and J.-I. Hong, *Organometallics*, 2005, 24, 1578–1585.
- 28 Q.-L. Xu, C.-C. Wang, T.-Y. Li, M.-Y. Teng, S. Zhang, Y.-M. Jing, X. Yang, W.-N. Li, C. Lin, Y.-X. Zheng, J.-L. Zuo and X.-Z. You, *Inorg. Chem.*, 2013, 52, 4916–4925.
- 29 G. Sarada, J. Yoon, W. Cho, M. Cho, D. W. Cho, S. O. Kang, Y. Nam, J. Y. Lee and S.-H. Jin, *J. Mater. Chem. C*, 2016, 4, 113–120.
- 30 B. J. Coe, M. Helliwell, J. Raftery, S. Sánchez, M. K. Peers and N. S. Scrutton, *Dalton Trans.*, 2015, 44, 20392–20405.
- 31 H. A. Bronstein, C. E. Finlayson, K. R. Kirov, R. H. Friend and C. K. Williams, *Organometallics*, 2008, 27, 2980–2989.
- 32 D. L. Davies, M. P. Lowe, K. S. Ryder, K. Singh and S. Singh, *Dalton Trans.*, 2011, **40**, 1028–1030.
- 33 N. M. Shavaleev, G. Xie, S. Varghese, D. B. Cordes, A. M. Z. Slawin, C. Momblona, E. Ortí, H. J. Bolink, I. D. W. Samuel and E. Zysman-Colman, *Inorg. Chem.*, 2015, 54, 5907–5914.
- 34 D. Tordera, J. J. Serrano-Pérez, A. Pertegás, E. Ortí, H. J. Bolink, E. Baranoff, M. K. Nazeeruddin and J. Frey, Chem. Mater., 2013, 25, 3391–3397.
- 35 N. Ishida, T. Moriya, T. Goya and M. Murakami, *J. Org. Chem.*, 2010, 75, 8709–8712.
- 36 L. Niu, H. Yang, D. Yang and H. Fu, *Adv. Synth. Catal.*, 2012, **354**, 2211–2217.
- 37 APEX2, version 2 User Manual, M86-E01078, Bruker Analytical X-ray Systems, Inc., Madison, WI, 2006.
- 38 P. W. Betteridge, J. P. Carruthers, R. I. Cooper, K. Prout and D. J. Watkin, *J. Appl. Crystallogr.*, 2003, **36**, 1487.
- M. J. Frisch, G. W. Trucks, H. B. Schlegel, G. E. Scuseria, M. A. Robb, J. R. Cheeseman, G. Scalmani, V. Barone, B. Mennucci, G. A. Petersson, H. Nakatsuji, M. Caricato, X. Li, H. P. Hratchian, A. F. Izmaylov, J. Bloino, G. Zheng, J. L. Sonnenberg, M. Hada, M. Ehara, K. Toyota, R. Fukuda, J. Hasegawa, M. Ishida, T. Nakajima, Y. Honda, O. Kitao, H. Nakai, T. Vreven, J. A. Montgomery Jr., J. E. Peralta, F. Ogliaro, M. Bearpark, J. J. Heyd, E. Brothers, K. N. Kudin, V. N. Staroverov, R. Kobayashi, J. Normand, K. Raghavachari, A. Rendell, J. C. Burant, S. S. Iyengar, J. Tomasi, M. Cossi, N. Rega, N. J. Millam, M. Klene, J. E. Knox, J. B. Cross, V. Bakken, C. Adamo, J. Jaramillo, R. Gomperts, R. E. Stratmann, O. Yazyev, A. J. Austin, R. Cammi, C. Pomelli, J. W. Ochterski, R. L. Martin,

K. Morokuma, V. G. Zakrzewski, G. A. Voth, P. Salvador, J. J. Dannenberg, S. Dapprich, A. D. Daniels, O. Farkas, J. B. Foresman, J. V. Ortiz, J. Cioslowski and D. J. Fox, *Gaussian 09, Revision D.01*, Gaussian, Inc., Wallingford, CT, 2009.

40 A. D. Becke, J. Chem. Phys., 1993, 98, 5648-5652.

Dalton Transactions

- 41 C. Lee, W. Yang and R. G. Parr, *Phys. Rev. B: Condens. Matter*, 1988, 37, 785–789.
- 42 M. M. Francl, W. J. Pietro, W. J. Hehre, J. S. Binkley, M. S. Gordon, D. J. DeFrees and J. A. Pople, *J. Chem. Phys.*, 1982, 77, 3654–3665.
- 43 P. J. Hay and W. R. Wadt, J. Chem. Phys., 1985, 82, 299-310.
- 44 J. Tomasi and M. Persico, Chem. Rev., 1994, 94, 2027-2094.

- 45 C. S. Cramer and D. G. Truhlar, in *Solvent Effects and Chemical Reactivity*, Kluwer, Dordrecht, 1996, pp. 1–80.
- 46 J. Tomasi, B. Mennucci and R. Cammi, *Chem. Rev.*, 2005, **105**, 2999–3093.
- 47 F. A. M. Rudolph, A. L. Fuller, A. M. Z. Slawin, M. Bühl, R. A. Aitken and J. D. Woollins, *J. Chem. Crystallogr.*, 2010, 40, 253–265.
- 48 C. Glidewell, W. T. A. Harrison, J. N. Low, J. G. Sime and J. L. Wardell, *Acta Crystallogr., Sect. B: Struct. Sci.*, 2001, 57, 190–200.
- 49 N. M. Shavaleev, R. Scopelliti, M. Grätzel, M. K. Nazeeruddin, A. Pertegás, C. Roldán-Carmona, D. Tordera and H. J. Bolink, J. Mater. Chem. C, 2013, 1, 2241–2248.