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When two are better than one: bright phosphorescence from non-stereogenic dinuclear iridium(III) complexes†

Ruth E. Daniels,^a Stacey Culham,^a Michael Hunter,^a Marcus C. Durrant,^a
Michael R. Probert,^b William Clegg,^b J. A. Gareth Williams^{*c} and
Valery N. Kozhevnikov^{*a}

A new family of eight dinuclear iridium(III) complexes has been prepared, featuring 4,6-diarylpyrimidines L^Y as bis- $N^A C$ -coordinating bridging ligands. The metal ions are also coordinated by a terminal $N^A C^A N$ -cyclometallating ligand L^X based on 1,3-di(2-pyridyl)benzene, and by a monodentate chloride or cyanide. The general formula of the compounds is $\{IrL^XZ\}_2L^Y$ ($Z = Cl$ or CN). The family comprises examples with three different L^X ligands and five different diarylpyrimidines L^Y , of which four are diphenylpyrimidines and one is a dithienylpyrimidine. The requisite proligands have been synthesised *via* standard cross-coupling methodology. The synthesis of the complexes involves a two-step procedure, in which L^XH is reacted with $IrCl_3 \cdot 3H_2O$ to form dinuclear complexes of the form $\{IrL^XCl(\mu-Cl)\}_2$, followed by treatment with the diarylpyrimidine L^YH_2 . Crucially, each complex is formed as a single compound only: the strong *trans* influence of the metallated rings dictates the relative disposition of the ligands, whilst the use of symmetrically substituted tridentate ligands eliminates the possibility of Λ and Δ enantiomers that are obtained when bis-bidentate units are linked through bridging ligands. The crystal structure of one member of the family has been obtained using a synchrotron X-ray source. All of the complexes are very brightly luminescent, with emission maxima in solution varying over the range 517–572 nm, according to the identity of the ligands. The highest-energy emitter is the cyanide derivative whilst the lowest is the complex with the dithienylpyrimidine. The trends in both the absorption and emission energies as a function of ligand substituent have been rationalised accurately with the aid of TD-DFT calculations. The lowest-excited singlet and triplet levels correlate with the trend in the HOMO–LUMO gap. All the complexes have quantum yields that are close to unity and phosphorescence lifetimes – of the order of 500 ns – that are unusually short for complexes of such brightness. These impressive properties stem from an unusually high rate of radiative decay, possibly due to spin–orbit coupling pathways being facilitated by the second metal ion, and to low non-radiative decay rates that may be related to the rigidity of the dinuclear scaffold.

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Introduction

The metal centre is an important component in luminescent cyclometallated iridium complexes, normally playing a key role in determining the nature of the luminescence. The high

degree of covalency of the Ir–C bond typically leads to efficient spin–orbit coupling (SOC), facilitated by the heavy metal, such that the formally forbidden transitions of intersystem crossing and phosphorescence are enhanced.¹ This is an important feature if the complex is to be used as a phosphor in organic light-emitting diodes for example, because it allows both triplet and singlet electro-excitation pathways to contribute to the emission, making the devices more efficient.² Meanwhile, in bio-imaging, bright phosphorescence opens up time-resolved detection techniques.^{3,4} The past 15 years have seen increasingly rapid advances in both fields and a large number of such complexes have been thoroughly investigated.⁵

The vast majority of examples to date contain one metal centre. Incorporating two or more metal centres in one luminophore can be an additional tool for influencing photo-

^aDepartment of Applied Sciences, Northumbria University, Newcastle upon Tyne, NE1 8ST, UK. E-mail: valery.kozhevnikov@northumbria.ac.uk

^bSchool of Chemistry, Newcastle University, Newcastle upon Tyne, NE1 7RU, UK

^cDepartment of Chemistry, Durham University, Durham, DH1 3LE, UK

E-mail: j.a.g.williams@durham.ac.uk

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physical properties, particularly if they are in close proximity.⁶ Of particular interest is the possibility that the presence of a second metal might increase the radiative rate constant of phosphorescence, k_r , compared to a mononuclear complex, by introducing a larger SOC effect than that associated with a single metal centre. Such a notion clearly assumes some synergy between the metal units. However, SOC pathways are complex and poorly understood even in mononuclear complexes,^{1a} and so a largely empirical approach based on experimental data is currently required in order to probe whether such effects are at play in multinuclear systems. The possibility of promoting k_r is of particular significance to the design of efficient red emitters, owing to the natural tendency of k_r to be lower for low-energy emitters (according to the Einstein coefficient for spontaneous emission which depends on ν^3) and to the normally faster rate of competitive non-radiative decay that low-energy excited states are subject to, in line with the energy-gap law.^{2e}

The synthesis of multinuclear complexes as single pure isomers is generally not easy. The D_3 or C_2 local symmetry of Ir(III) complexes containing bidentate ligands {e.g. those based on $\text{Ir}(\text{N}^{\wedge}\text{C})_3$ or $[\text{Ir}(\text{N}^{\wedge}\text{C})_2(\text{L}^{\wedge}\text{X})]^+$ structures} means that such complexes are chiral and are formed as a racemic mixture of Λ and Δ enantiomers. Since enantiomers have identical [linear] optical properties, this is generally of little consequence for mononuclear complexes,⁷ but when two or more such metal centres are incorporated into one molecule, a mixture of diastereoisomers necessarily forms (e.g. $\Lambda\Lambda/\Delta\Delta + \Lambda\Delta$ in the case of two metal centres). Diastereoisomers have different properties and need to be separated, a process that can often be exceptionally laborious and time-consuming. For example, Tsuboyama and co-workers described the synthesis of a red-emitting dinuclear iridium complex, related to the archetypal complex $\text{Ir}(\text{ppy})_3$, but featuring a bridging 1,4-di(2-pyridyl)benzene ligand (dpb).⁸ The desired di-iridium complex $\{\text{Ir}(\text{ppy})_2\}(\text{dpb})$ could be isolated in only a 3% yield after column chromatography. In many other instances of multinuclear complexes, it has proved impossible to isolate all the isomers in pure form.⁹

One way to potentially alleviate this problem is to make use of tridentate ligands. For example, the achirality of bis-terpyridyl metal complexes (C_{2v} symmetry) compared to the chirality of D_3 tris-bipyridyl analogues has driven extensive research into the supramolecular chemistry of terpyridyl complexes since the 1980s.¹⁰ In the field of cyclometallated iridium(III) chemistry, the use of tridentate ligands has lagged far behind that of bidentate ligands, although examples of both charge-neutral and cationic complexes have been reported.^{11–13} In particular, achiral mononuclear Ir(III) complexes of the form $\text{Ir}(\text{N}^{\wedge}\text{C}^{\wedge}\text{N})(\text{N}^{\wedge}\text{C})\text{X}$ – comprising derivatives of the tridentate $\text{N}^{\wedge}\text{C}^{\wedge}\text{N}$ -cyclometallating ligand 1,3-bis(2-pyridyl) benzene in conjunction with bidentate $\text{N}^{\wedge}\text{C}$ -cyclometallating 2-phenylpyridines and monodentate halide or acetylide ligands X – have been found to display excellent luminescent properties, with luminescence quantum yields approaching unity in some case. Some have been successfully incorporated into OLEDs, showing good performance.^{12d,e}

We reasoned that such use of the $\text{N}^{\wedge}\text{C}^{\wedge}\text{N}$ motif, in conjunction with a 4,6-diarylpyrimidine as a bis- $\text{N}^{\wedge}\text{C}$ -coordinating bridging unit, could provide an attractive route to achiral dinuclear complexes, with none of the problems associated with the formation of additional isomers. Our prototypical dinuclear complex of this new class is $\{\text{Ir}(\text{L}^{\wedge}\text{Cl})_2\}\text{L}^1$ (Fig. 1). We have also prepared a series of seven other such complexes, which highlight the structural diversity offered by the approach in terms of variation of the $\text{N}^{\wedge}\text{C}^{\wedge}\text{N}$ ligand (labelled alphabetically L^{A} to L^{G}), the bis- $\text{N}^{\wedge}\text{C}$ coordinating ligand (labelled numerically L^1 to L^5), and metathesis of the monodentate ligand. Note that the approach requires that the $\text{N}^{\wedge}\text{C}^{\wedge}\text{N}$ ligand be symmetrically substituted, i.e., featuring a C_2 axis or mirror plane through the central ring; the use of asymmetrically-substituted ligands would re-introduce chirality.¹⁴ The structures of the tri- and bidentate ligands used are shown in Fig. 1a, with the corresponding complexes in Fig. 1b. The synthesis and characterisation of these new luminophores are reported here, together with a study of their impressive luminescence properties interpreted with the aid of TD-DFT calculations.

Results and discussion

Synthesis

Our objectives were to explore the luminescence properties of this new family of complexes, examining the influence of the substitution pattern in the bridging bis- $\text{N}^{\wedge}\text{C}$ ligand and, independently, in the $\text{N}^{\wedge}\text{C}^{\wedge}\text{N}$ ligand. Thus, we prepared a series of five 4,6-diaryl-substituted pyrimidines $\text{L}^1\text{H}_2\text{--L}^5\text{H}_2$, with varying degrees of electron-withdrawing or donating power, ranging from difluorophenyl through to 2-thienyl (Fig. 1a). These proligands were readily accessible using Suzuki–Miyaura cross-coupling reactions between 4,6-dichloropyrimidine and the corresponding arylboronic acids (Scheme 1a). We have previously used similar pyrimidine-based ligands, together with their pyrazine analogues, to access a range of dinuclear Pt(II) complexes with widely-tuneable emission.¹⁵ Details of all new syntheses and the characterisation of new compounds are provided in the ESI.†

A series of three $\text{N}^{\wedge}\text{C}^{\wedge}\text{N}$ proligands $\text{H}^{\text{A}}\text{L--H}^{\text{L}}\text{C}$ were also synthesised, each comprising lateral 4-*tert*-butylpyridine groups, but in which the central ring was varied (Fig. 1a). We chose to include fluorine atoms in the central ring, *ortho* to the pyridyl rings, since competitive metallation of iridium at these positions has been shown to occur in the absence of substituents, giving $\text{N}^{\wedge}\text{C}$ -coordinated products as opposed to the desired $\text{N}^{\wedge}\text{C}^{\wedge}\text{N}$ systems.^{16,17} The fluorine atoms also facilitate the synthesis of these ligands, by *ortho*-directing the lithiation reactions involved in the synthesis (Scheme 1b). Thus, $\text{L}^{\text{A}}\text{H}$ was prepared from intermediate 2 (whose synthesis we described previously in the preparation of bis- $\text{N}^{\wedge}\text{C}^{\wedge}\text{N}$ -coordinating ligands^{6h}) by conversion to the boronic acid 3 through lithiation and reaction with tri-*iso*-propylborate, followed by Suzuki–Miyaura cross-coupling of 3 with 4-*tert*-butyl-2-chloropyridine.



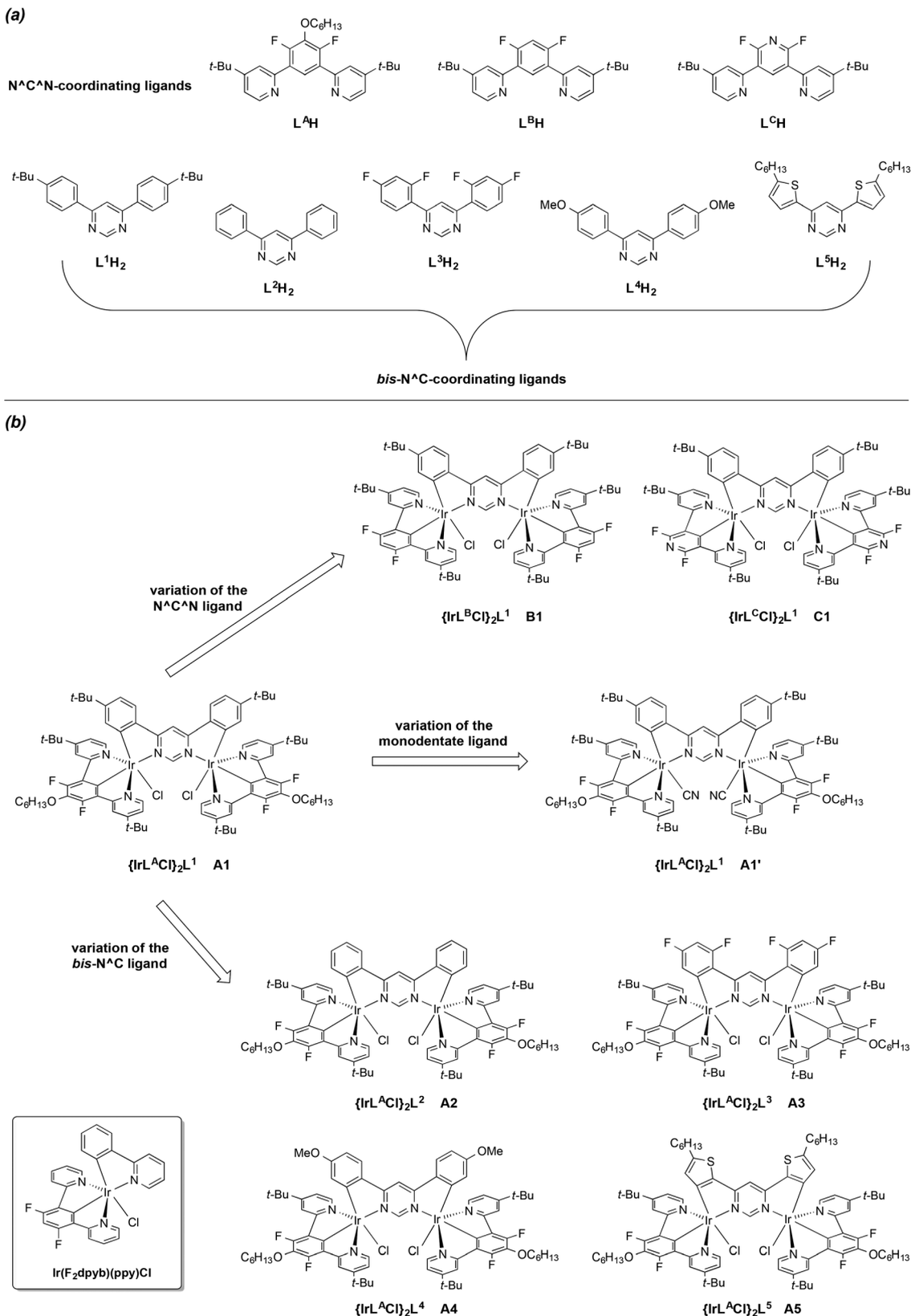


Fig. 1 Structural formulae of the ligands (a) and complexes (b) investigated in this work. The structure of Ir(F₂dpyb)(ppy)Cl, used as a model mono-nuclear complex for comparison,^{12e} is shown in the boxed inset at the bottom left.



for related mononuclear complexes (Scheme 1c).^{12b,e} In the first step, reaction of the tridentate proligands with hydrated iridium(III) chloride in a mixture of ethoxyethanol and water generated the dichloro-bridged dimers $[\text{Ir}(\text{L}^{\text{A-C}})\text{Cl}(\mu\text{-Cl})]_2$. The presence of the *tert*-butyl substituents increases the solubility of these products compared to the related complexes of 1,3-dipyridyl-4,6-difluorobenzene, which are insoluble in all common solvents.^{12e} Indeed, the solubility of all three dimers was sufficiently high to allow them to be characterised fully by ¹H, ¹⁹F and ¹³C-NMR spectroscopy in CDCl₃ solution (see ESI†).

In the second step, the dichloro-bridged dimers were reacted with the bis-N[^]C proligands L¹H₂-L⁵H₂ under reflux in toluene, in the presence of silver triflate as a chloride scavenger. After the reaction, the mixture was treated with excess HCl (2 M, aq.) which ensures that the monodentate ligands in the desired products are exclusively Cl.

The identities of the resulting dinuclear complexes were confirmed by ¹H, ¹⁹F and ¹³C NMR spectroscopy and by mass spectrometry. It is important to stress that, due to the very strong *trans*-influence of the formally anionic, metallated carbon atoms (see also Section 2 below), the product which forms in each case is exclusively that in which the pyrimidine ring is disposed *trans* to the central aryl ring of the N[^]C[^]N ligand. There is no evidence for the formation of isomers in which the metallated rings of the N[^]C and N[^]C[^]N units are *trans* to one another. In other words, the product in each case is uniquely that in which the pyrimidine nitrogen atom lies in the same plane as the N[^]C[^]N ligand. Isomeric products in which one or both of the metallated (*i.e.*, C-coordinating) rings of the bis-N[^]C ligand occupy such a position are not formed. This observation is in line with earlier studies of related mononuclear systems.^{12b-e,17} Thus, in contrast to most other dinuclear complexes studied to date,^{6,8,9} *only one stereoisomer is formed* for these complexes. The most characteristic features in the ¹H NMR spectra are (i) a singlet with a very high chemical shift δ in the region 11–12 ppm, assigned to H² of the pyrimidine ring, and (ii) a low-frequency multiplet in the aromatic region, δ 5.6–6.8 ppm, assigned to the protons *ortho* to the metallated carbon in the aryl ring of the bridging ligand. For ease of discussion in subsequent sections and representation in the figures that follow, the dinuclear complexes $\{\text{Ir}(\text{L}^{\text{XCl}})_2\}\text{L}^{\text{Y}}$ (X = A, B or C; Y = 1 to 5) will be referred to as **Xy**; *e.g.* $\{\text{Ir}(\text{L}^{\text{A}}\text{Cl})_2\}\text{L}^1$ is **A1**, $\{\text{Ir}(\text{L}^{\text{B}}\text{Cl})_2\}\text{L}^2$ is **B2** *etc.*

A monodentate chloro ligand completes the coordination sphere of each Ir(III) centre in the dimers. Metathesis of this ligand with other monodentate ligands could provide access to a range of other derivatives, potentially allowing further tuning of the properties.¹⁷ We have demonstrated the ease with which such metathesis can be accomplished by converting $\{\text{Ir}(\text{L}^{\text{A}}\text{Cl})_2\}\text{L}^1$ (**A1**) to its cyano derivative $\{\text{Ir}(\text{L}^{\text{A}}\text{CN})_2\}\text{L}^1$ (which we shall refer to as **A1'**), simply by treatment with excess KCN.

Crystallography

Crystals of $\{\text{Ir}(\text{L}^{\text{A}}\text{Cl})_2\}\text{L}^3$ (**A3**) suitable for X-ray diffraction analysis were obtained by slow evaporation of a solution of the complex in CH₂Cl₂/MeOH (3 : 1 v/v). The molecular structure is

shown in Fig. 2, with key bond lengths and angles involving the metal listed in Table 1. The molecule has a crystallographic mirror plane, passing through atoms C38 and C39. Two-fold disorder was modelled for atoms C28–C30 (and their H atoms) of the hexyl chain (minor disorder component not shown). Displacement parameters, short contacts, and residual electron density peaks suggest further disorder of this chain and of the *t*-butyl groups, but this was not modelled.

It should be noted that the structure contains highly disordered and unidentified solvent in large void volumes (shown in yellow in the *a*-axis projection in Fig. 3) between the molecules of the dinuclear complex. This was dealt with by the SQUEEZE procedure of the program PLATON.¹⁹ Calculations based on unit cell contents do not include this solvent. Full

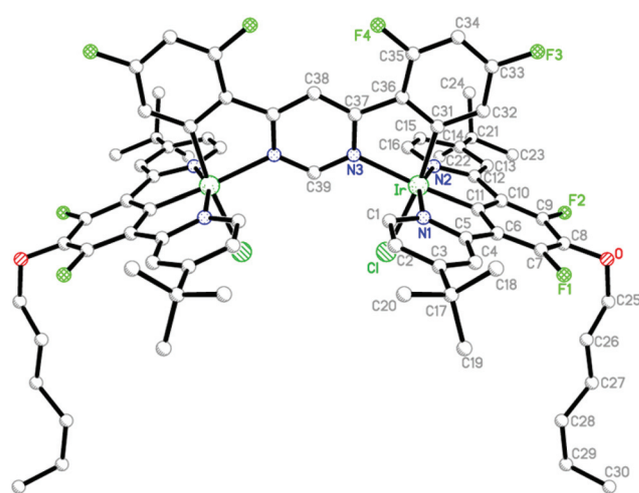


Fig. 2 Molecular structure of the dinuclear complex $\{\text{Ir}(\text{L}^{\text{A}}\text{Cl})_2\}\text{L}^3$ (**A3**) in the crystal. Hydrogen atoms and minor disorder components have been omitted for clarity.

Table 1 Selected bond lengths (Å) and angles (°) for complex **A3** determined by X-ray crystallography, and the corresponding values in the energy-minimised structure obtained by DFT calculations

Bond length (Å) or angle (°)	X-ray data	DFT data ^a
Ir–N1	2.049(5)	2.090 [2.085]
Ir–N2	2.048(5)	2.081 [2.085]
Ir–N3	2.157(5)	2.208 [2.215]
Ir–C11	1.926(7)	1.936 [1.936]
Ir–C31	1.996(5)	2.024 [2.024]
Ir–Cl	2.4406(15)	2.483 [2.479]
N1–Ir–N2	160.4(2)	159.9 [159.9]
N1–Ir–C11	80.3(2)	79.9 [80.0]
N2–Ir–C11	80.1(2)	80.1 [80.0]
N3–Ir–C11	173.4(2)	176.1 [176.5]
N3–Ir–C31	78.9(2)	78.3 [78.5]
N3–Ir–Cl	93.66(12)	92.2 [92.7]
C31–Ir–Cl	172.6(2)	170.1 [171.1]
C11–Ir–C31	95.3(3)	97.9 [98.0]

^a DFT-derived values in square parentheses are for the geometry with imposed mirror symmetry about the central ligand (see text).



details of the structure are provided in the ESI,[†] and the structure has been deposited at the CCDC, reference no: 1415318.

By comparing with structurally related mononuclear complexes based on terpyridine rather than an N[∧]C[∧]N-coordinating ligand, the structure of **A3** provides evidence for the strong *trans* influence associated with the central metallated ring of the N[∧]C[∧]N ligand alluded to in section 2. For example, the Ir–N3 bond length of 2.157(5) Å is elongated compared to the Ir–N bond (of N[∧]C[∧]N ligand) of [Ir(tpy)(MeOmpy)Cl]⁺, for which the value is 2.068(2) Å,^{13g} and likewise the equatorial Ir–N(bpy) bond of [Ir(tpy)(bpy)Cl]²⁺, for which the bond length is 2.081(1) Å (Fig. 4).²⁰ In these compounds, the Ir–N bonds are *trans* to Ir–N as opposed to Ir–C bonds. The strong *trans* influence of the metallated carbon atoms of the diarylpyrimidine ligand is similarly evident from the longer Ir–Cl bond of 2.4406(15) Å in **A3** compared to a value of 2.334(5) Å in [Ir(tpy)(bpy)Cl]²⁺, where the chloride is *trans* to a pyridine rather than to a metallated aryl ring.

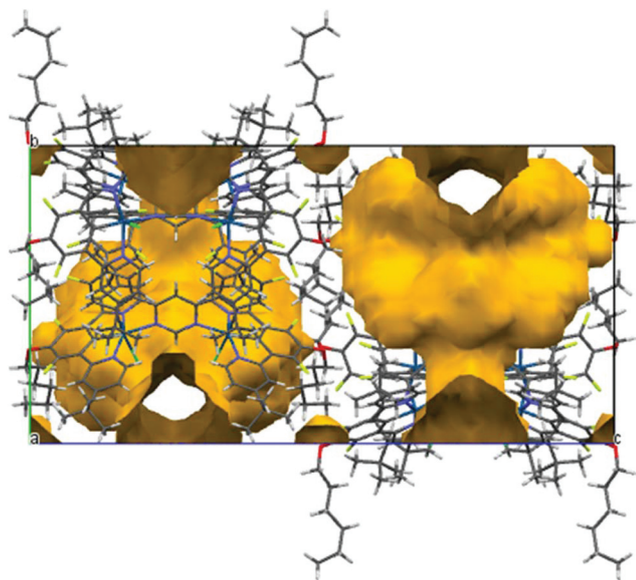


Fig. 3 The large voids (shown in yellow) in the crystal structure of the dinuclear complex **A3**.

Density functional theory calculations

Density functional theory (DFT) calculations have been carried out on each of the new complexes, as described in the Experimental section. In order to expedite the calculations, the *t*-butyl and OC₆H₁₃ substituents of the tridentate ligands and the thienyl C₆H₁₃ substituents of **A5** were truncated to Me, OMe and Me respectively. The *t*-butyl substituents of **L**¹ were retained in full. After an initial wavefunction stability check, molecules were subjected to full geometry optimisation. Due to the large size of these dinuclear iridium complexes, frequency calculations proved to be intractable and were therefore omitted.

All of the optimised geometries of these complexes showed a twist about the central pyrimidine ligand, such that the dihedral angles between the two phenyl rings varied from 7.9° for **A2** to 22.8° for **A1**'. The equivalent dihedral angle between the smaller thiophene rings in **A5** was much smaller, at 1.0°. In all cases, this distortion results in rotational symmetry about the central pyrimidine ring. In contrast, the X-ray crystal structure of **A3** shows mirror symmetry about the pyrimidine (Fig. 2), such that the central ligand remains planar rather than twisted. This results in different distances between the equivalent pyridine N atoms in the two outer ligands (6.293 and 7.433 Å) in the experimental structure, whereas these distances are equal (at 7.104 Å) in the calculated structure. A geometry optimisation starting from the X-ray crystal structure produced the same calculated structure; hence these slightly different conformations appear to be similar in energy. This conclusion was confirmed by re-optimising the geometry of **A3** with imposed mirror symmetry, by the use of z-matrix techniques; this approach gave an optimised structure that was closer to the experimental structure, but just 0.3 kJ mol⁻¹ higher in energy than the rotationally symmetric conformer. Hence, deformation about the central ligand is clearly a very low-energy process that is within both the margin of error of the calculations and the likely effects of molecular packing within the crystal structure.

In all other respects, the calculated structures are as expected (see Table 1 for a comparison of calculated and experimental bond lengths and angles). In each case, the metal is in a distorted octahedral geometry. The Ir–N(pyridine) bond lengths (range 2.079–2.096 Å) are shorter than the Ir–N

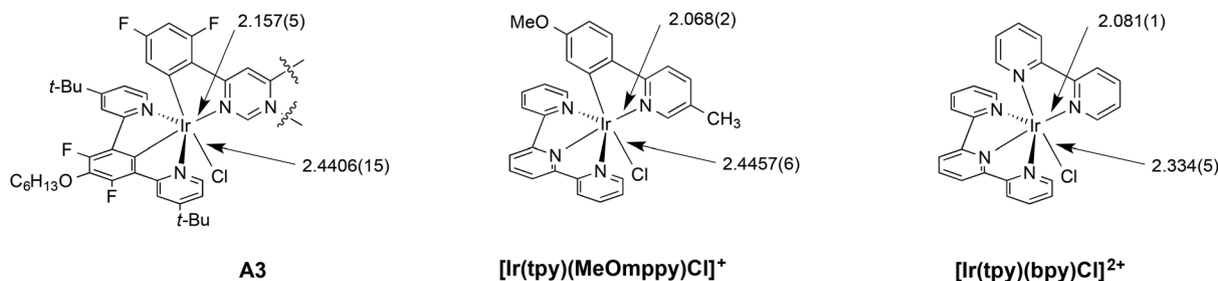


Fig. 4 Ir–N and Ir–Cl bond lengths (in Å) in **A3** and related complexes highlighting the *trans* influence of the metallated carbon atoms in elongating *trans*-disposed bonds.



(pyrimidine) bond lengths (2.208–2.263 Å), as are the Ir–C bond lengths for the tridentate compared to the bidentate cyclometallating ligands (1.917–1.936 Å and 2.017–2.085 Å respectively). These differences mirror those found for related mononuclear complexes and are consistent with the expected *trans* influences of the various donor atoms involved.^{12e,17} The Ir–Cl bond lengths cover a narrow range of 2.478–2.489 Å, whilst the Ir–CN bond length in **A1'** is 2.062 Å.

UV-visible absorption spectroscopy

In discussing the optical properties of the complexes, it is helpful to divide them into three series, with **A1** common to all. In the series **A1**–**A5**, the tridentate ligand (L^A) is common to each member, and so changes in properties within this series can be ascribed to the influence of the bis- N^A C ligand. Meanwhile, in the series **A1**–**C1**, the evolution of properties can be related to changing the tridentate ligand. Complex **A1** and its cyano derivative **A1'** form the third, small series.

The UV-visible absorption spectra of the complexes in dichloromethane solution at room temperature are shown in Fig. 5, and absorption maxima and extinction coefficients are tabulated in Table 2. All eight complexes show a series of very intense bands in the UV region (ϵ in the range 30 – $80 \times 10^3 \text{ M}^{-1} \text{ cm}^{-1}$) together with intense bands in the visible region up to around 500 or 530 nm (ϵ in the range 5 – $30 \times 10^3 \text{ M}^{-1} \text{ cm}^{-1}$). There is a general trend of decreasing molar absorptivities with increasing wavelength. The spectra are quite typical of cyclometallated iridium(III) complexes,^{1,2,5} although with higher extinction coefficients than those of related mononuclear complexes of the $\text{Ir}(N^A C^A N)(N^A C)Cl$ class, such as $\text{Ir}(\text{F}_2\text{dpyb})(\text{ppy})\text{Cl}$ (the structure of which is shown in the inset to Fig. 1 and for which UV-visible absorption data are listed in the footnote to Table 2;^{12e} $\text{F}_2\text{dpybH} = 1,3\text{-di}(2\text{-pyridyl})\text{-4,6-difluorobenzene}$).

Within the **A1**–**C1** series, it can be seen that the absorption spectra have very similar profiles, but the bands at $\lambda > 370 \text{ nm}$ are increasingly blue-shifted on going from **A1** to **B1** to **C1** (Fig. 5a and Table 2). The change from a difluorophenyl to difluoropyridyl ring (**B1** to **C1**) shifts the bands by around 500 cm^{-1} to higher energy. On the other hand, the presence of the alkoxy substituent in **A1** compared to **B1** lowers the energy of the bands by a smaller amount of around 200 cm^{-1} . Meanwhile, changing the monodentate ligand from chloride in **A1** to cyanide in **A1'** is seen to result in a larger blue-shift of the lowest-energy absorption bands of around 1000 cm^{-1} (Fig. 5b and Table 2). The overall spectral profile is, however, otherwise similar, and the higher-energy region $< 350 \text{ nm}$ is little affected. For the series **A1**–**A5**, in which the bis- N^A C ligand is modified, the spectra of **A1**–**A4** are all similar to one another in profile, with the lowest-energy absorption bands increasing in energy in the order $\text{A2} < \text{A1} < \text{A3} < \text{A4}$ (Fig. 5c and Table 2). The bis-thienylpyrimidine complex **A5**, on the other hand, is rather different from the others at wavelengths $> 450 \text{ nm}$, in that the two bands which appear in this region have much higher extinction coefficients than those of the other complexes. In addition, this complex displays a weak band at low

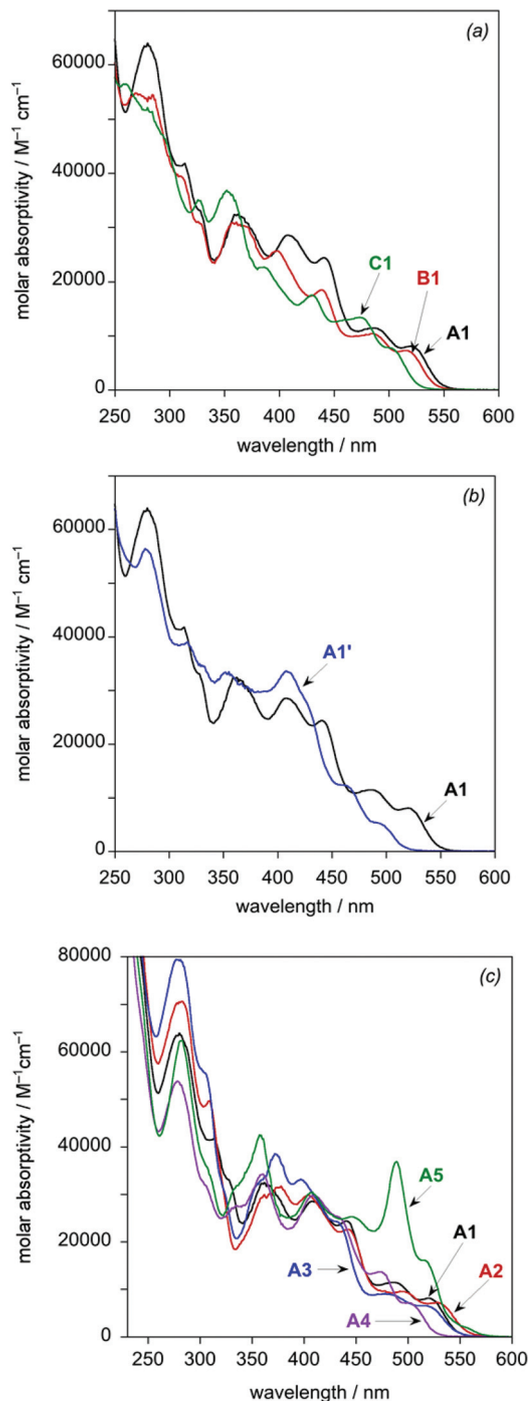


Fig. 5 UV-visible absorption spectra of the dinuclear Ir(III) complexes in CH_2Cl_2 at $298 \pm 3 \text{ K}$: (a) the **A1**–**C1** series; (b) **A1** and **A1'**; (c) the **A1**–**A5** series.

energy ($\lambda_{\text{max}} = 554 \text{ nm}$, $\epsilon = 2100 \text{ M}^{-1} \text{ cm}^{-1}$), which has no apparent counterpart in the other complexes.

Luminescence properties

All eight of the new complexes are intensely luminescent in deoxygenated solutions at room temperature. Key emission data are tabulated in Table 3, whilst the emission spectra are



Table 2 UV-visible absorption data for the complexes in CH₂Cl₂ at 298 ± 3 K

Complex ^a	λ_{\max}/nm ($\epsilon/\text{M}^{-1} \text{cm}^{-1}$)
A1	279 (63 400), 312 (41 500), 360 (32 300), 408 (28 600), 441 (24 400), 484 (11 400), 520 (8080)
B1	270 (54 700), 310sh (39 500), 326sh (31 100), 363 (30 400), 398 (25 600), 438 (18 500), 484 (10 300), 514 (7300)
C1	260 (56 500), 281 (51 500), 327 (34 900), 354 (36 500), 385 (22 500), 430 (17 500), 474 (13 400), 500 (7800)
A1'	279 (56 300), 315 (38 800), 353 (33 000), 409 (33 500), 460 (12 300), 491 (5390)
A2	281 (70 400), 309 (49 600), 376 (31 500), 404 (29 800), 442 (22 600), 494 (9510), 526 (7120)
A3	279 (79 300), 303sh (55 800), 358sh (32 700), 372 (38 600), 397 (33 200), 430 (24 300), 477 (8970), 514 (6560)
A4	277 (53 600), 335 (27 600), 360 (34 300), 409 (29 700), 433sh (25 200), 473 (13 700), 498 (7070)
A5	281 (62 300), 305sh (36 200), 337sh (31 800), 358 (42 200), 407 (30 300), 446 (25 300), 489 (36 800), 514 (16 100), 554 (2100)

^a For comparison, data for the related mononuclear complex Ir(F₂dpyb)(ppy)Cl (Fig. 1 inset) are as follows: λ_{\max}/nm ($\epsilon/\text{M}^{-1} \text{cm}^{-1}$) = 260 (44 800), 280 (40 300), 358 (9100), 381 (11 300), 402 (14 000), 438 (3640), 473 (1540) [ref. 12e].

Table 3 Emission data for the complexes in degassed CH₂Cl₂ at 298 K and in EPA^b at 77 K

Complex ^a	λ_{\max}/nm	Φ_{lum}^c	τ/ns^d	$k_r^e/10^6 \text{ s}^{-1}$	$\sum k_{\text{nr}}^e/10^6 \text{ s}^{-1}$	$k_Q^f/10^9 \text{ M}^{-1} \text{ s}^{-1}$	Emission at 77 K ^g	
							λ_{\max}/nm	τ/ns
A1	552, 582sh	0.94	440 [210]	2.14	0.14	1.13	513, 552, 594	6650
B1	544, 576sh	1.0	550 [160]	1.82	— ^h	2.01	512, 549, 593sh	7500
C1	526, 557sh	1.0	590 [230]	1.69	— ^h	1.21	501, 538, 578	7800
A1'	517, 547sh	0.88	590 [240]	1.49	0.20	1.12	488, 523, 560	5100
A2	568, 599sh	0.88	570 [230]	1.54	0.21	1.18	519, 559, 602	4200
A3	570	0.91	590 [220]	1.54	0.15	1.30	525, 563, 610sh	1300
A4	528, 557sh	1.0	360 [160]	2.78	— ^h	1.58	493, 530, 569	2400
A5	572, 611sh	0.94	4000 [380]	0.24	0.015	1.08	560, 582, 608, 633, 663, 695, 728	23 000

^a For comparison, data for the related mononuclear complex Ir(F₂dpyb)(ppy)Cl (Fig. 1 inset) are as follows: at 298 K, $\lambda_{\max} = 487, 516 \text{ nm}$; $\Phi_{\text{lum}} = 0.20$; $\tau = 390 \text{ ns}$; $k_r = 0.5 \times 10^6 \text{ s}^{-1}$; $\sum k_{\text{nr}} = 2.1 \times 10^6 \text{ s}^{-1}$. At 77 K, $\lambda_{\max} = 471, 508, 537, 548, 593 \text{ nm}$; $\tau = 2.5 \mu\text{s}$ [ref. 12e]. ^b EPA = diethyl ether/isopentane/ethanol (2 : 2 : 1 v/v). ^c Measured using Ir(ppy)₃ in 2-MeTHF as the standard, for which $\Phi = 0.97$. ^d Values in air-equilibrated solution in parentheses. ^e k_r and $\sum k_{\text{nr}}$ are the radiative and non-radiative decay rate constants, estimated from the quantum yield and lifetime assuming that the emissive state is formed with unitary efficiency. ^f Biomolecular rate constant for quenching by O₂, estimated from the luminescence lifetimes in degassed and air-equilibrated solutions, and taking [O₂] = 2.2 mM in CH₂Cl₂ at $p = 1 \text{ atm}$ air and $T = 298 \text{ K}$. ^g In EPA: see footnote a. ^h Since the measured quantum yield is unity within the error of the measurement, it is not possible to make a meaningful estimate of k_{nr} .

illustrated in Fig. 6 (subdivided into the same groups as those of Fig. 5). Most of the spectra show a well-defined maximum, accompanied by a shoulder to the low-energy side; such profiles are typical of rigid chromophores with a small Huang–Rhys factor, in which the emission is concentrated in the highest-energy 0,0 vibrational component band.²¹ The vibrational structure becomes much more pronounced at 77 K (Fig. 7), under which conditions all of the spectra show at least three components of a vibrational progression of $\sim 1400 \text{ cm}^{-1}$, typical of aromatic C=C vibrational modes. The thienyl complex additionally shows bands due to a second, lower-frequency progression. The following trends emerge from the spectra.

(i) For the **A1**–**C1** series, the emission energy increases in the order **A1** < **B1** < **C1**, mirroring the trend observed in absorption. Moreover, based on the 0,0 maxima, the changes in emission energy on going from **A1** to **B1** ($\sim 200 \text{ cm}^{-1}$) and from **B1** to **C1** ($\sim 500 \text{ cm}^{-1}$) are very similar to those observed for the lowest-energy absorption band.

(ii) The replacement of Cl by CN in going from **A1** to **A1'** blue-shifts the emission by $\sim 1000 \text{ cm}^{-1}$, similar in magnitude to the blue-shift observed in absorption.

(iii) Within the series **A1**–**A5**, the trend in emission energy at room temperature is **A5** \sim **A3** \sim **A2** < **A1** < **A4**. At this tempera-

ture, the lower-energy vibrational shoulder is only just detectable for **A3** and essentially not at all for **A2**. At 77 K, the 0,0 components are clearly resolved, however, and the order of emission energy is **A5** < **A3** < **A2** < **A1** < **A4**.

The excitation spectra closely match the absorption spectra in each case, indicating that the same emissive state is formed with high efficiency, irrespective of the excitation wavelength. Such behaviour is quite typical of cyclometallated Ir(III) and Pt(II) complexes, where the processes of internal conversion and intersystem crossing to the lowest-lying triplet state are all several orders of magnitude faster than the rate of fluorescence from the singlet manifold.²² The luminescence quantum yields in solution at room temperature are very high for all of the complexes, approaching unity, within the uncertainty on the measurement. The quantum yields are higher than those of previously explored members of the mononuclear Ir(N[^]C[^]N)(N[^]C)Cl series (e.g., see footnote to Table 3).^{12e} The values are comparable to those of *fac*-Ir(ppy)₃, despite the emission being significantly further towards the red than this archetypal complex (apart from **A1'**).²³ With the exception of the thienyl complex **A5**, the luminescence lifetimes are in the range 360–590 ns at room temperature, which are significantly shorter than that of *fac*-Ir(ppy)₃, for which a



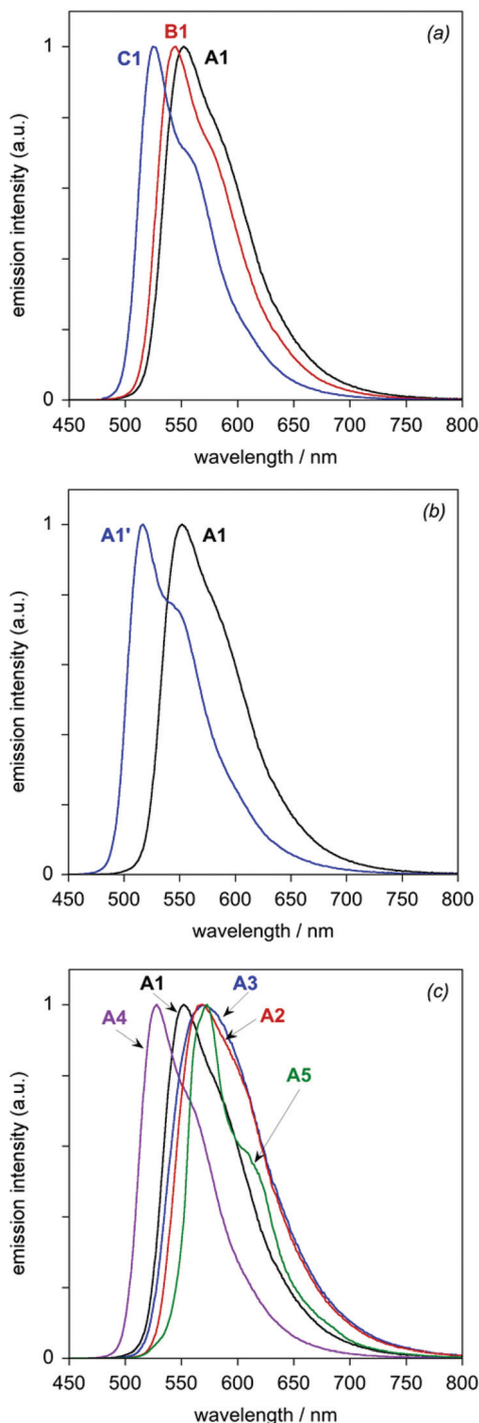


Fig. 6 Luminescence spectra of the dinuclear Ir(III) complexes in CH_2Cl_2 at 298 ± 3 K: (a) the A1–C1 series; (b) A1 and A1'; (c) the A1–A5 series.

value of $1.6 \mu\text{s}$ has been reported under comparable conditions.²³ Given that the quantum yields are comparable to that of *fac*-Ir(ppy)₃, this observation is suggestive of a significantly higher rate of radiative decay in the dinuclear complexes. Assuming that the emissive state is formed with unitary efficiency upon photo-excitation, the radiative and

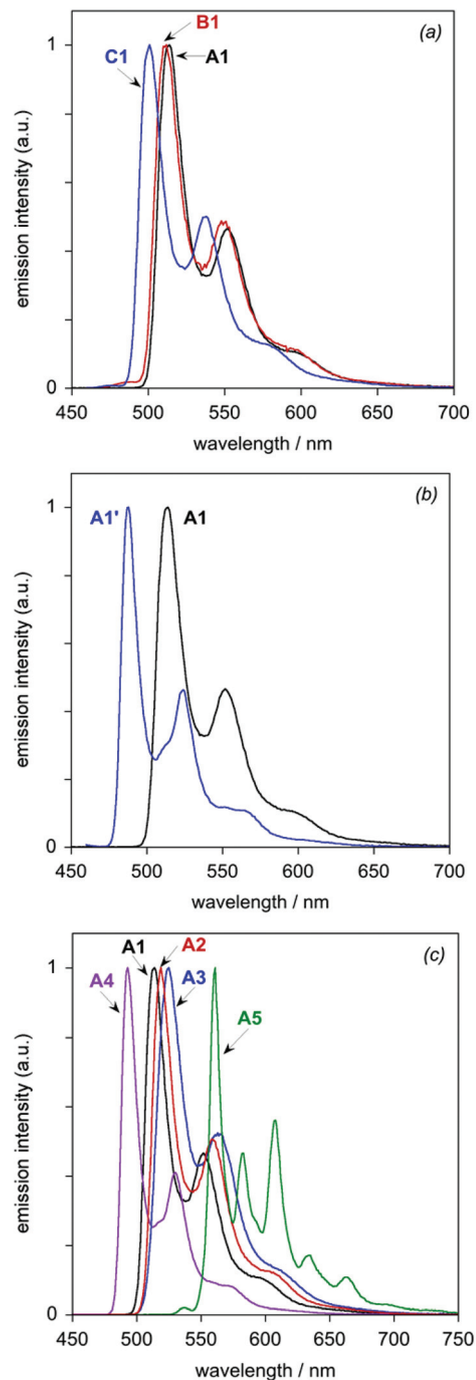


Fig. 7 Luminescence spectra of the Ir(III) complexes in diethyl ether/isopentane/ethanol (2 : 2 : 1) at 77 K: (a) the A1–C1 series; (b) A1 and A1'; (c) the A1–A5 series.

non-radiative rate constants, k_r and $\sum k_{nr}$ respectively, can be estimated through the relationships.

$$k_r = \Phi/\tau \text{ and } \sum k_{nr} = (1 - \Phi)/\tau$$

The values of the rate constants estimated in this way are included in Table 3. It is evident that, apart from A5, the k_r



values are increased by a factor of 2–5 compared to that of Ir(ppy)₃, for which k_r has been estimated to be $6 \times 10^5 \text{ s}^{-1}$.²³

The luminescence of all the complexes is quenched by oxygen, with a bimolecular rate constant of the order of $10^9 \text{ M}^{-1} \text{ s}^{-1}$, quite typical of cyclometallated Ir(III) and Pt(II) complexes. However, since k_r values are unusually high and the lifetimes correspondingly short (again with the exception of A5), the effect of oxygen at atmospheric pressure of air is unusually small, reducing the lifetimes and emission intensities by a factor of only about 2. Complex A5 is anomalous in that it has a much longer lifetime than the other complexes, both at room temperature and at 77 K ($\tau = 4$ and 23 μs respectively). Its emission in solution is correspondingly more sensitive to O₂, being quenched by a factor of ~ 10 under air-equilibrated conditions.

Interpretation of the photophysical data with reference to TD-DFT results

Time-dependent DFT (TD-DFT) calculations were carried out at the optimised geometries obtained as described in section 3. The main excitation energies, their oscillator strengths and the respective predominant molecular orbital contributions are compiled in Tables S1–S8 of the ESI.† Visible spectra were calculated in vacuum, and in dichloromethane solution using the polarised continuum model for the solvent²⁴ (Fig. S1 and S2†). In all cases, the general effect of solvent is to blue-shift the frequency of the highest energy singlet band by 15 to 38 nm and to increase its intensity. The experimental spectra show more structure in the absorption bands than those simulated using the TD-DFT data. Such structure is likely to arise as a result of absorption to vibrationally excited levels of the electronically excited states, which is not modelled by the theory. For example, the lowest-energy bands of A1 are separated by around 1400 cm^{-1} , which is typical of aromatic C=C vibrations. This mirrors the vibrational progression observed in the emission spectra (albeit arising in that case from transitions from the triplet state to the vibrationally excited levels of the ground state).

The calculations provide very useful insight into the trends in absorption and emission energies discussed earlier. For all of the complexes, the trend in the energy of the lowest-energy absorption band determined experimentally correlates well with the HOMO–LUMO gap, as does the trend in the experimental emission energy, as illustrated in Fig. 8c. Plots of the calculated S₁ and T₁ energies in vacuum and in solvent are shown in the ESI,† and likewise show a very good correlation with the experimentally determined data (Fig. S3†). Thus, it is appropriate to seek to understand the influence of the substituents in terms of their effects on the HOMO and LUMO energies. Molecular orbital plots of the HOMO and LUMO of all of the complexes (as well as HOMO–1 and LUMO+1) are shown in the ESI (Fig. S4–S11†). Typically for mononuclear cyclometallated Ir(III) and Pt(II) complexes with aryl-heterocycle derivatives (e.g. ppy), the LUMO is based largely on the heterocyclic ring (pyridine in the case of ppy), whilst the HOMO spans the metallated ring and the metal atom, reflecting the

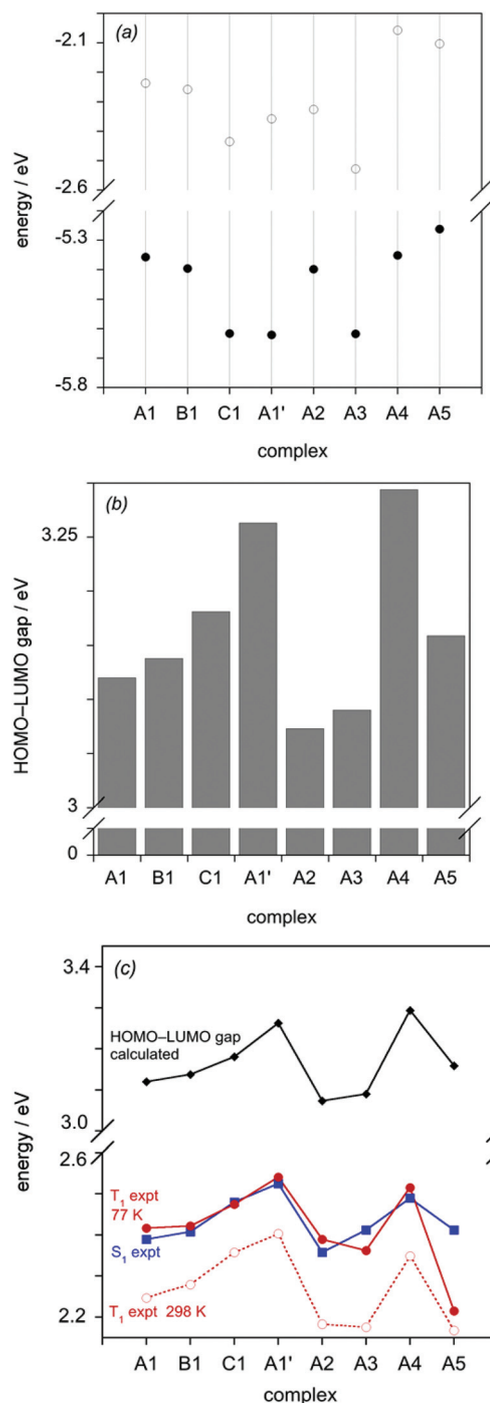


Fig. 8 (a) Calculated HOMO and LUMO energies for the complexes *in vacuo*, in eV (filled and empty circles respectively). (b) The HOMO–LUMO gap (eV) calculated from the respective energies in (a). (c) Plot of the experimental S₁ energy (blue squares), estimated based on λ_{max} of the lowest-energy absorption band (excluding the weak band at 554 nm for A5, which is concluded to be a triplet transition, S₀ → T₁ – see text); the T₁ energy estimated from experimental λ_{max} (0,0) values in the emission spectra at 298 K (open red circles) and at 77 K (filled red circles); and the calculated HOMO–LUMO gap (black diamonds). The lines between points are provided solely as a guide to aid the eye.



high degree of covalency of the M–C bond.^{2e,25} In the case of the series of dinuclear complexes reported here, one might intuitively expect the more electron-deficient pyrimidine heterocycle of the bis-N^{^C} ligand to offer lower-energy π^* orbitals than the pyridine rings of the tridentate ligand, and indeed the calculations confirm that the LUMO is predominantly localised on the pyrimidine ring in all eight complexes (Fig. S4–S11†). On the other hand, the HOMO spans the metal and the metallated ring of either the N^{^C}N or the N^{^C} ligand, according to the identity of the two ligands.

Fig. 8a shows the calculated HOMO and LUMO energies of the complexes and Fig. 8b illustrates the magnitude of the resulting gap. The blue-shift in absorption and emission that is observed on going from **A1** through **B1** to **C1** can be seen to arise from a larger stabilisation of the HOMO than the LUMO. Examining the HOMO orbital plots, it can be seen that the HOMO of **A1** involves the aryl ring of the N^{^C}N ligand with only a minor contribution from that of the N^{^C} unit (Fig. S4†). Removal of the electron-donating alkoxy group on going to **B1** switches round this distribution of electron density in the HOMO: it now involves almost primarily the N^{^C} aryl ring with only a minor contribution from the N^{^C}N (Fig. S5†). In **C1**, in which the central aryl ring of the N^{^C}N ligand is replaced by a more electron-deficient pyridyl ring, the contribution of the tridentate ligand to the HOMO becomes negligible (Fig. S6†).

The replacement of the chloride ligand by cyanide in (**A1** to **A1'**) can similarly be seen to have a much larger stabilising effect on the HOMO than the LUMO (Fig. 8b), accounting for the large blue-shift observed experimentally both in absorption and emission. This is consistent with the observation that the metal typically makes a much more significant contribution to the HOMO than the LUMO.

The small red-shift in absorption and emission on going from **A1** to **A2** can be seen from Fig. 8a to arise primarily from a stabilisation of the LUMO, larger than the effect on the HOMO, upon removal of the *t*-butyl substituents in the N^{^C} ligand. Meanwhile, the data in Fig. 8b would suggest that **A3** should be very similar to **A2**, which is indeed the case for the emission, see Fig. 6c (in absorption, a small blue shift is observed experimentally, Fig. 5c). At first sight, this is a somewhat surprising result, given that *fac*-Ir(dfppy)₃ is well-known to be strongly blue-shifted relative to *fac*-Ir(ppy)₃ for example, attributed to the effect of the fluorine atoms in stabilising the HOMO.²⁶ However, from Fig. 8a, it is apparent that the electron-withdrawing fluorine atoms in **A3** stabilise the LUMO and HOMO to similar extents, leading to the lack of significant effect on the optical properties. In contrast, the significant blue-shift observed in the absorption and emission of **A4** compared to **A1** is seen from Fig. 8a to arise almost exclusively from the destabilisation of the N^{^C}-based LUMO. In this regard, it is pertinent to note that the electron-donating methoxy substituents of **A4** are located *para* to the electron-deficient pyrimidine ring. Interestingly, inspection of the frontier orbitals of **A4** (Fig. S10†) shows that the methoxy substituent is sufficiently electron-rich to shift the HOMO to the N^{^C} as opposed to the N^{^C}N ligand.

Finally, we turn to complex **A5**. We noted its anomalously long luminescence lifetime of 4 μ s in section 5. At 77 K, the lifetime of this complex increases to 23 μ s, which is much longer than that of typical ³MLCT states in Ir(III) complexes, whilst its spectrum at 77 K displays much more vibrational structure. Such changes are consistent with an excited state of more thienyl-pyrimidine localised ³ π - π^* parentage, mirroring conclusions made on Ir(thpy)₂(acac) versus Ir(ppy)₂(acac) for example {and similarly for Pt(thpy)(acac) versus Pt(ppy)(acac)}.²⁷ It seems likely that at 298 K, states of predominant ³MLCT and ³ π - π^* character are close in energy, but the former is displaced to higher energy at 77 K, leading to the distinctly ³ π - π^* -like spectrum and lifetime observed at low temperature.²⁸ It is notable that the simplistic correlation with the HOMO–LUMO gap works well for the S₁ state of **A5** (from the absorption spectrum) but less well for the emission data, particularly at 77 K (Fig. 8c). In contrast, the more rigorous calculation of the T₁ energy captures the trend well, with **A5** predicted to have much the lowest-energy emission amongst all the complexes, as observed at 77 K (Fig. S3†). Indeed, we surmise that the additional weak absorption band observed at 554 nm for **A5** may be the direct excitation to the ³ π - π^* state.²⁹

What can be concluded about the influence of a second metal ion? It is clear from comparison with the most closely related mononuclear complexes available (see data in Table 3 and footnote) that the radiative rate constant in the dinuclear complexes is enhanced whilst the non-radiative decay is suppressed. This observation is all the more remarkable given that the excited state energies of these new complexes are significantly lower than in the model mononuclear complexes: typically, for a given class of complex, non-radiative decay processes increase with decreasing excited state energy, according to the energy-gap law, whilst a decrease in *k_r* may be anticipated, according to the Einstein coefficient which depends on ν^3 . The consistency with which *k_r* is enhanced in our recently studied multinuclear complexes featuring such bis-cyclometalating ligands supports the notion that spin–orbit coupling pathways may be enhanced by the presence of a second heavy metal centre.^{6d,h,15} That non-radiative decay is suppressed may, on the other hand, arise from the greater conformational rigidity of the dinuclear systems.

Concluding remarks

In summary, we have shown that 4,6-diarylpyrimidines are attractive bis-cyclometalating ligands that can be used to bind simultaneously to two iridium ions *via* N^{^C} coordination to each metal centre. The use of a tridentate, N^{^C}N-coordinating, terminal ligand ensures that the resulting dinuclear complexes are achiral. This contrasts with the mixtures of stereoisomers that are formed when tris-bidentate metal units are employed. Moreover, the strong *trans* influence of cyclometalated aryl rings leads to one unique product, in which the pyrimidine ring is *trans* to the central ring of the N^{^C}N ligand at both metal centres. The resulting products are very



intensely luminescent, with quantum yields close to unity and the emission energy being tuneable through substituents in either the bidentate or tridentate ligand or metathesis of the monodentate ligand. Whilst offering strong light absorption and intense emission in the visible region, dinuclear cyclo-metallated complexes clearly offer huge potential of structural diversity. The versatility of the design strategy, and ease of synthesis of suitable tri- and bidentate ligands, can be expected to lead to further new compounds, for example, designed as water-splitting catalysts or for specific interactions with biomolecules, in both of which fields multinuclear complexes are increasingly recognised to offer unique opportunities.^{30,31}

Experimental

Synthesis and characterisation

Full details are provided in the ESI.† ¹H and ¹³C NMR spectra were recorded on JEOL Delta-270 or JEOL ECS-400 spectrometers operating at the frequencies indicated in the ESI.† Chemical shifts (δ) are in ppm, referenced to tetramethylsilane Si(CH₃)₄, and coupling constants are in Hertz. Mass spectra were recorded at the EPSRC National Mass Spectrometry Service Centre on a Thermo Scientific LTQ Orbitrap XL Mass Spectrometer using low resolution electrospray (ESI) or high resolution nano-electrospray (NSI) ionisation techniques. Pro-ligands L¹H₂-L⁵H₂ were prepared from 4,6-dichloropyrimidine, using the procedure we reported previously.¹⁵

Photophysical measurements

Absorption spectra were measured on a Biotek Instruments XS spectrometer, using quartz cuvettes of 1 cm path length. Steady-state luminescence spectra were measured using a Jobin Yvon FluoroMax-2 spectrofluorimeter, fitted with a red-sensitive Hamamatsu R928 photomultiplier tube; the spectra shown are corrected for the wavelength dependence of the detector, and the quoted emission maxima refer to the values after correction. Samples for emission measurements were contained within quartz cuvettes of 1 cm path length modified with appropriate glassware to allow connection to a high-vacuum line. Degassing was achieved *via* a minimum of three freeze-pump-thaw cycles whilst connected to the vacuum manifold; final vapour pressure at 77 K was $<5 \times 10^{-2}$ mbar, as monitored using a Pirani gauge. Luminescence quantum yields were determined using *fac*-Ir(ppy)₃ in degassed 2-methyltetrahydrofuran solution as the reference ($\Phi_{\text{lum}} = 0.97$ (ref. 23)); estimated uncertainty in relative values of Φ_{lum} is $\pm 10\%$ or better. The luminescence lifetimes of the complexes were measured by time-correlated single-photon counting, following excitation at 405 nm with an EPL-405 pulsed-diode laser. The emitted light was detected at 90° using a Peltier-cooled R928 PMT after passage through a monochromator. The lifetime of A5 at 77 K was measured using the same detector operating in multi-channel scaling mode, following excitation with a microsecond pulsed xenon flash-lamp. The estimated uncertainty in the quoted lifetimes is $\pm 10\%$ or

better. Bimolecular rate constants for quenching by molecular oxygen, k_{Q} , were determined from the lifetimes in degassed and air-equilibrated solution, taking the concentration of oxygen in CH₂Cl₂ at 0.21 atm O₂ to be 2.2 mmol dm⁻³.

Density functional theory calculations

All DFT calculations were carried out using Gaussian 09W.³² The B3LYP hybrid functional was used,³³ together with the LanL2DZ basis set for iridium,³⁴ and 6-31+G(d) for all other atom types. Geometries were fully optimised *in vacuo*. UV-visible absorption spectra were obtained by single point TD-DFT calculations at the optimised geometries, using the same functional and basis sets, either *in vacuo* or with a PCM solvent correction for dichloromethane.²⁴ The molecular orbitals and spectra were visualised using GaussView 5.0 with Gaussian line shapes and an arbitrary half-width at half-height of 1000 cm⁻¹ for the latter.

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