The complex [Ir(ppy)₂(dp bpy)][PF₆] (Hppy = 2-phenylpyridine, dp bpy = 6,6'-diphenyl-2,2'-bipyridine) has been prepared and evaluated as an electroluminescent component for light-emitting electrochemical cells (LECs); the complex exhibits two intramolecular face-to-face π-stacking interactions and long-lived LECs have been constructed; the device characteristics are not significantly improved in comparison to analogous LECs with 6-phenyl-2,2'-bipyridine.

Light-emitting electrochemical cells (LECs) are a minimalistic derivative of organic light-emitting devices (OLEDs) and in their simplest form consist of a film of an ionic transition metal complex placed between two electrodes.¹,² LECs offer considerable technological advantages over OLEDs as they require a less reactive cathode material (Al instead of Ca or Mg) because the device is no longer dependent upon the work function of the electrode and hence do not require stringent protection from environmental oxygen or water. The disadvantage of LECs is the short operating lifetime, in the order of hours to days, compared to OLEDs.³–⁵ We have recently reported the use of intra- and intermolecular face-to-face π-stacking for the stabilisation of the ground and excited state of electroluminescent iridium complexes and shown that this leads to exceptionally long-living LEC devices.⁶–⁷ The long lifetimes of these devices establish LECs as a viable alternative to OLED technology. In [Ir(ppy)(ppy)]⁺ (Hppy = 2-phenylpyridine, pppy = 6-phenyl-2,2'-bipyridine) the pendant phenyl group of the ppy ligand forms a face-to-face π-stack with the metallated ring of a ppy ligand (3.2–3.5 Å). This interaction minimises the expansion of the metal–ligand bonds in the excited state and precludes the attack by water and other nucleophiles resulting in the long observed lifetimes. We concluded that analogous complexes with 6,6'-diphenyl-2,2'-bipyridine would have an even greater stabilisation of the excited state as the two pendant phenyl groups would stack with different ppy ligands giving a very "tight" complex.

The ligand 6,6'-diphenyl-2,2'-bipyridine, dp bpy, was obtained from the reaction of four equivalents of phenyllithium with 2,2'-bipyridine in THF followed by oxidation of the intermediate tetrahydro-species with MnO₂ according to the general procedure of Sauvage et al.⁸ The reaction of dp bpy with the chloro-bridged dimer [ppy]Ir(μ-Cl)₂Ir(ppy)₂]⁹ under standard conditions (reflux in 1 : 1 CH₂Cl₂–MeOH in the presence of Ag[PF₆]̅¹⁰ resulted in a smooth conversion to [Ir(ppy)₂(dp bpy)][PF₆] from which the orange salt [Ir(ppy)₂(dp bpy)][PF₆] was obtained in 89% yield. The complex was fully characterised by conventional methods and exhibited two peaks with similar intensity in the ESMS spectrum at m/z 809.2 [M – PF₆]⁺ and 501.1 [M – dp bpy – PF₆]⁺, respectively. The orange colour is associated with a weak and broad MLCT absorption at 474 nm (CH₃CN, ε = 920 M⁻¹ cm⁻¹) and the complex is luminescent exhibiting an emission in MeCN solution with a maximum at 595 nm with a lifetime τ = 0.6 μs and a quantum yield (PLQE) of 3%.

We have determined the structure of [Ir(ppy)₂(dp bpy)][PF₆] and the [Ir(ppy)₂(dp bpy)]⁺ cation present in the lattice is shown in Fig. 1a. The Ir–N(ppy) distances (2.0504(17), 2.0341(17) Å) and Ir–C(ppy) distances (2.0120(18), 2.0212(2) Å) closely resemble those previously reported for [Ir(ppy)₂(ppy)][PF₆]⁺ (Ir–N, 2.036(3)–2.068(3), Ir–C, 2.004(3)–2.025(3) Å). The Ir–N(dp bpy) distances (2.2017(18), 2.215(3) Å) are, however, significantly longer than those in [Ir(ppy)₂(dp bpy)][PF₆] (2.148(3)–2.215(3) Å), which is a consequence of the π-stacking of the pendant phenyl groups. The pendant phenyl rings exhibit the expected π-stacking with the ppy ligands; the phenyl ring containing C39 lies 3.1–3.3 Å from the ppy containing N1 and that containing C33 3.0–3.35 Å from the ppy containing N2. The dihedral angle between the two pyridine rings of the bpy is 20.7°, which compares favourably with angles of 18.4° and 20.7° for the two independent cations in the unit cell of [Ir(ppy)₂(dp bpy)][PF₆]. We stress here that the intramolecular π-stacking is a direct and inevitable consequence of the ligand structure and will be present in the solid state, solution and thin film phases. To summarise, as observed from the crystal structure, the use of the dp bpy ligand for optimising the

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**Two are not always better than one: ligand optimisation for long-living light-emitting electrochemical cells†**

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The light output is extremely slow resulting in turn-on times of the pbpy and dpbpy complexes. In both cases, the build-up of graphs is the similarity of the behaviour of the devices using methylimidazolium (BMIP) is incorporated to shorten the aluminium layer as cathode. The ionic liquid (IL) 1-butyl-3-methylimidazolium hexafluorophosphate and finally an anode of aluminium (Al) coated with a 0.1 μm spin-coated layer of polystyrene–polysulfonic acid (PEDOT:PSS) followed by a 80 nm spin-coated layer of 4 : 1 [Ir(ppy)2(dpbb)][PF6]–1-butyl-3-methylimidazolium hexafluorophosphonate and finally an aluminium layer as cathode. The liquid iodide (IL) 1-butyl-3-methylimidazolium (BMIP) is incorporated to shorten the turn-on time of the device. 1,2,3,4 Characteristics of devices at other ratios of IL to complex are presented in the ESI. 3

The temporal behaviour of electroluminescent devices containing [Ir(ppy)2(dpbb)][PF6] and [Ir(ppy)2(dpbb)][PF6] is depicted in Fig. 2. It is obvious from these two graphs is the similarity of the behaviour of the devices using the ppy and dpbb complexes. In both cases, the build-up of the light output is extremely slow resulting in turn-on times (t_on) of several days. It is possible to reduce this extremely long t_on either by adding more IL or by pre-biasing the devices with short intervals of higher voltages. 5

For a better comparison between the two devices we did not employ these methods for the data herein reported. After reaching the maximum luminance, both devices show a slow decrease in luminance versus time and reach a value of 50% of the maximum luminance after approximately 1300 h. This time is referred to as the lifetime (t_L). Thus, the turn-on time and the lifetime are similar for devices employing the ppy and dpbb complexes. In addition, the current efficacy (2.7 and 3.1 cd A⁻¹), the external quantum efficiency (EQE: 1% and 1.1%) and the power efficiency (2.8 and 3.3 Lm W⁻¹) displayed by the [Ir(ppy)2(dpbb)]⁺ and the [Ir(ppy)2(dpbb)]⁺ devices are also similar. We can categorically state that these device lifetime data are the best ever reported except those published with devices containing [Ir(ppy)2(dpbb)][PF6] operated with a pre-biasing method. 6

The main difference between the two devices lies in the absolute values of the observed luminances. The devices incorporating [Ir(ppy)2(dpbb)][PF6] have significantly higher luminance values than those using [Ir(ppy)2(dpbb)][PF6], 110 versus 70 cd m⁻², respectively. The lifetime of organic luminescent devices depends, in general, strongly on the luminance values at which they are operated. To take this into account in LEC devices, Kalyuzhny et al. proposed the use of the total photon flux (E_\text{t,0}) emitted up to the time the luminance reaches 1/5th of the maximum value (t_1/5) for a cell area of 3 mm². 1,2 By extrapolating the lifetime curves, E_\text{t,1/5} values of 13.6 and 6.9 J are obtained for the devices using [Ir(ppy)2(dpbb)][PF6] and [Ir(ppy)2(dpbb)][PF6], respectively, which are significantly larger than the best value reported in the literature (0.27 J). 2 Hence, taking the different luminance levels and the total photon flux data into account, one must conclude that the device using [Ir(ppy)2(dpbb)][PF6] is less stable than the device using [Ir(ppy)2(dpbb)][PF6], thereby showing that two phenyl groups on the bpy are not better than one phenyl group.

In an attempt to understand why one phenyl group is better than two, density functional theory (DFT) calculations were performed at the B3LYP/6–31G**/LANL2DZ level on [Ir(ppy)2(dpbb)]⁺ and [Ir(ppy)2(dpbb)]⁺ cations. The geometries of the singlet ground state (S₀), the emitting triplet state (T₁) and the metal-centered triplet state (3MC) were fully optimized for both complexes. Calculations accurately reproduce the face-to-face π–π stacking observed in the solid state for [Ir(ppy)2(dpbb)]⁺ between the pendant phenyl rings of the bpy ligand with the adjacent ppy ligands. The intramolecular π–π interaction is preserved for the lowest triplet excited state T₁. This makes the complex more robust in the excited emitting state and reduces the possibility of triplet-triplet annihilation. However, the planarity of the ppy ligands is not well satisfied for the [Ir(ppy)2(dpbb)]⁺ cation. This reflects the difference in the π–π interaction of the ppy ligands.
for ligand-exchange reactions leading to the degradation of the complex.

State $T_1$ originates from the HOMO $\rightarrow$ LUMO excitation and is calculated to lie 2.22 and 2.32 eV above the $S_0$ state (adiabatic energy difference) for $[\text{Ir}(\text{ppy})_2(\text{bppy})]^+$ and $[\text{Ir}(\text{ppy})_2(\text{dpbpy})]^+$, respectively. For both complexes, the emitting $T_1$ state is described as a mixture of metal-to-ligand and ligand-to-ligand charge transfer ($\text{MLCT}$ and $\text{LLCT}$, respectively) due to the participation of both the Ir atom and the phenyl rings of the ppy ligands in the HOMO while the LUMO is fully located on the bpy ligand (see Fig. S5 in the ESI†). The structure of the $[\text{Ir}(\text{ppy})_2(\text{dpbpy})]^+$ complex in its $^3\text{MC}$ state is similar to that obtained for the $[\text{Ir}(\text{ppy})_2(\text{bpy})]^+$ complex (Table S1†). However, the relative energy position of the $^3\text{MC}$ state with respect to the emitting triplet state is significantly different. In the case of the $[\text{Ir}(\text{ppy})_2(\text{bpy})]^+$ complex, the $^3\text{MC}$ state is calculated after geometry relaxation to lie at approximately 0.60 eV above the emitting $T_1$ state. In contrast, the $^3\text{MC}$ state in the $[\text{Ir}(\text{ppy})_2(\text{dpbpy})]^+$ complex is located at only 0.26 eV above the $T_1$ state (Fig. 3). Hence, the probability of populating the $^3\text{MC}$ states increases from $[\text{Ir}(\text{ppy})_2(\text{bpy})]^+$ to $[\text{Ir}(\text{ppy})_2(\text{dpbpy})]^+$, thereby increasing the probability for non-radiative decay to the ground state and degradation reactions.

In conclusion, a new cationic iridium(III) complex has been prepared that exhibits a double intramolecular $\pi$-stacking between the two phenyl groups of the dpbpy ligand and different cyclometallated ppy ligands, resulting in an intramolecular cage formation. When used as the active component in a light-emitting electrochemical cell, the device exhibits extraordinarily long lifetimes of around 1300 h. Although very long, somewhat surprisingly the lifetime is not increased with respect to a device using a similar complex that can only form one phenyl-ppy intramolecular $\pi$-stack. One reason for this is attributed to the distortion of the planarity of the bpy ligand domain when the two attached phenyl groups $\pi$-stack with the phenyl groups of the ppy ligands. Quantum chemical calculations show that the energy difference between the emitting triplet and the metal-centred triplet state has decreased which renders the complex more susceptible to emission losses and degradation reactions. Thus, in this case two is not better than one.

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Notes and references