Phosphane tuning in heteroleptic [Cu(N^N)(P^P)]⁺ complexes for light-emitting electrochemical cells†

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The synthesis and characterization of five [Cu(P^P)(N^N)][PF₆] complexes in which P^P = 2,7-bis(tert-butyl)-4,5-bis(diphenylphosphino)-9,9-dimethylxanthene (8Bu₂xantphos) or the chiral 4,5-bis(mesityl-phenylphosphino)-9,9-dimethylxanthene (xantphosMes₂) and N^N = 2,2’-bipyridine (bpy), 6-methyl-2,2’-bipyridine (6-Mebpy) or 6,6’-dimethyl-2,2’-bipyridine (6,6’-Me₂bpy) are reported. Single crystal structures of four of the compounds confirm that the copper(i) centre is in a distorted tetrahedral environment. In [Cu(xantphosMes₂)(6-Mebpy)][PF₆], the 6-Mebpy unit is disordered over two equally populated orientations and this disorder parallels a combination of two dynamic processes which we propose for [Cu(xantphosMes₂)(N^N)][PF₆] cations in solution. Density functional theory (DFT) calculations reveal that the energy difference between the two conformers observed in the solid-state structure of [Cu(xantphosMes₂)(6-Mebpy)][PF₆] differs in energy by only 0.28 kcal mol⁻¹. Upon excitation into the MLCT region (λexc = 365 nm), the [Cu(P^P)(N^N)][PF₆] compounds are yellow to orange emitters. Increasing the number of Me groups in the bpy unit shifts the emission to higher energies, and moves the Cu⁺/Cu²⁺ oxidation to higher potentials. Photoluminescence quantum yields (PLQYs) of the compounds are low in solution, but in the solid state PLQYs of up to 59% (for [Cu(8Bu₂xantphos)(6,6’-Me₂bpy)][PF₆]) are observed. Increased excited-state lifetimes at low temperature are consistent with the complexes exhibiting thermally activated delayed fluorescence (TADF). This is supported by the small energy difference calculated between the lowest-energy singlet and triplet excited states (0.17–0.25 eV). The compounds were tested in simple bilayer light-emitting electrochemical cells (LECs). The optoelectronic performances of complexes containing xantphosMes₂ were generally lower with respect to those with 8Bu₂xantphos, which led to bright and efficient devices. The best performing LECs were obtained for the complex [Cu(8Bu₂xantphos)(6,6’-Me₂bpy)][PF₆] due to the increased steric hindrance at the N^N ligand, resulting in higher PLQY. (LECs) and interest in these devices has grown tremendously in the last few years.¹–⁴ OLEDs are now well established and are widely employed in display applications. LECs feature many of the advantages of OLEDs including direct electron-to-photon conversion and the possibility of fabrication employing flexible surfaces and thin-film processing. Additionally, the simple device architecture of LECs and the use of air-stable electrode materials might reduce the manufacturing cost of electroluminescent devices and widen their field of applications.⁵–⁷ LECs incorporating ionic transition-metal complexes (iTMCs) based on iridium (and to a lesser extent ruthenium) have been the focus of intense investigations and show good performances in terms of colour tunability, brightness and device lifetime.⁸–¹⁴ However, the limited availability of iridium and ruthenium in the Earth’s crust motivated the search for alternative emissive materials.
LEC devices employing copper(i) iTMCs show promising characteristics. McMillin and co-workers first reported the photoluminescent properties of copper(i) complexes containing 2,2′-bipyridine (bpy) or 1,10-phenanthroline (phen) and PPh₃ or chelating bis(phosphane) ligands, and the applications of which boosts device performance. Indirect harvesting of fluorescence from the triplet state, from the triplet state at room temperature. This enables the indirect harvesting of fluorescence from the triplet state, which boosts device performance.

Here, we report on the effects of modifying the xantphos ligand and investigate the influence on both the structural and electronic properties. It has previously been reported that in complexes containing Cu(P^P)(N^N)], and Cu(xantphos)(N^N)]⁺ species is significantly reduced when the phenyl groups in xantphos (Scheme 1) are replaced by tert-butyl substituents, attributed to vibrational quenching effects. Thus, for good photophysical properties it appears crucial to retain ary1 substituents on the phosphorus atoms in ligands related to xantphos or POP.

Experimental

Details of general instruments are given in the ESL. tert-Bu₂xantphos has previously been used in combination with palladium(u) in vinylarene hydroamination catalysts. The introduction of peripheral tert-butyl groups was expected to additionally result in a larger spatial separation of complex ions in the active layer in a LEC and therefore have an influence on the electroluminescent properties. The other \( \text{P}^\ast \) ligands investigated in this work were 4,5-bis(dimethylphosphino)-9,9-dimethylxanthene (xantphos), and 4,5-bis(di-tert-butylphosphino)xanthene. In the latter case, it has been observed that the photoluminescence (PL) of \([\text{Cu}(\text{P}^\ast \text{P})(\text{N}^\ast \text{N})]⁺\) species is significantly reduced when the phenyl groups in xantphos (Scheme 1) are replaced by tert-butyl substituents, attributed to vibrational quenching effects. Thus, for good photophysical properties it appears crucial to retain ary1 substituents on the phosphorus atoms in ligands related to xantphos or POP.

For this reason, we have studied the influence on both the structural and electronic properties. It has previously been reported that in complexes containing Cu(P^P)(N^N)]⁻ and Cu(xantphos)(N^N)]⁻ complexes, the HOMO is mainly centred on copper with small contributions from phosphorus, while the LUMO is localized on the N^N ligand. Structural modifications made to the P^P ligand will have little if any effect on the energy level of the HOMO and any structural tuning should lead primarily to changes in steric effects. We have introduced tert-butyl groups into the 2,7-positions of xantphos to give \( \text{Bu}_2\text{xantphos} \) (Scheme 1). The \( \text{Bu}_2\text{xantphos} \) has previously been used in combination with palladium(u) in vinylarene hydroamination catalysis. The introduction of peripheral tert-butyl groups was expected to additionally result in a larger spatial separation of complex ions in the active layer in a LEC and therefore have an influence on the electroluminescent properties. The other \( \text{P}^\ast \) ligands investigated in this work were 4,5-bis(dimethylphosphino)-9,9-dimethylxanthene (xantphosMes₂) and 4,5-bis(dimethylphosphino)-9,9-dimethylxanthene (xantphosMes₂), in which the Ph₂ groups in xantphos were replaced by either PMes₂ or PPhMes units (Scheme 1), thereby tuning the Tolman cone angle of the phosphane. The N^N ligands chosen for the investigation were bpy, 6-Mebpy and 6,6′-Me₂bpy (Scheme 1). The photophysical properties of \([\text{Cu}(\text{xantphos})(\text{N}^\ast \text{N})][\text{PF}_6] \) (N^N = bpy, 6-Mebpy and 6,6′-Me₂bpy) have been previously reported, giving a benchmark series for the present investigation.
and were used as received. Details of the syntheses and characterizations of 6-Mebpy, xantphosMes4.Cl, chloro(mesityl) phenylphosphane, xantphosMes2, [Cu(Bu2xantphos)[bpy]][PF6], [Cu(Bu2xantphos)(6-Mebpy)][PF6], [Cu(Bu2xantphos)(6,6'-Me-bpy)][PF6], [Cu(xantphosMes2)[bpy]][PF6] and [Cu(xantphosMes2)(6-Mebpy)][PF6] are given in the ESL.†

Crystallography

Data were collected on a Bruker Kappa Apex2 diffractometer with data reduction, solution and refinement using the programs APEX43 and CRYSTALS.42 Structural analysis was carried out using Mercury v. 3.7,31,32 In [Cu(Bu2xantphos)(6-Mebpy)][PF6], one CH2Cl2 molecule was refined and SQUEEZE35 was used to treat part of the solvent region; formulae and numbers were modified in the cif to keep this result into account. In [Cu(xantphosMes2)(6-Mebpy)][PF6], the 6-Mebpy is orientationally disordered over two orientations and the relevant aromatic rings were refined as rigid bodies.

**xantphosMes2**

C14H14O4P2, M = 662.79, colourless plate, monoclinic, space group P21/c, a = 10.6380(9), b = 15.0841(13), c = 23.5452(2) Å, β = 102.385(3)°, U = 6387.4(5) Å3, Z = 4, Dc = 1.194 Mg m–3, μ(Cu-Ko) = 1.317 mm–1, T = 123 K. Total 40993 reflections, 6810 unique, Rint = 0.035. Refinement of 6264 reflections (433 parameters) with I > 2σ(I) converged at final R1 = 0.0336 (R1 all data = 0.0365), wR2 = 0.0776 (wR2 all data = 0.0794), GOF = 0.9775. CCDC 1860879.

**[Cu(Bu2xantphos)(bpy)][PF6]·0.5Et2O**

C35H63Cl3Cu6F6N2O4s·P21, M = 1092.60, yellow block, triclinic, space group P1, a = 12.0991(12), b = 13.3253(13), c = 18.6750(18) Å, α = 91.353(3), β = 90.939(3), γ = 115.475(2)°, U = 2716.2(5) Å3, Z = 2, Dc = 1.336 Mg m–3, μ(Cu-Ko) = 1.932 mm–1, T = 123 K. Total 35686 reflections, 9826 unique, Rint = 0.029. Refinement of 9540 reflections (664 parameters) with I > 2σ(I) converged at final R1 = 0.0416 (R1 all data = 0.0423), wR2 = 0.1012 (wR2 all data = 0.1016), GOF = 0.9707. CCDC 1844060.

**[Cu(Bu2xantphos)(6-Mebpy)][PF6]·1.5CH2Cl2·0.5H2O**

C39H64Cl6Cu6F6N2O4s·P21, M = 1205.97, yellow block, triclinic, space group P1, a = 10.7080(9), b = 13.4475(12), c = 22.1672(7) Å, α = 73.142(6), β = 79.483(6), γ = 86.606(6)°, U = 3003.4(5) Å3, Z = 2, Dc = 1.33 Mg m–3, μ(Cu-Ko) = 2.997 mm–1, T = 123 K. Total 35561 reflections, 10903 unique, Rint = 0.082. Refinement of 7619 reflections (700 parameters) with I > 2σ(I) converged at final R1 = 0.1178 (R1 all data = 0.1490), wR2 = 0.1151 (wR2 all data = 0.1470), GOF = 1.0317. CCDC 1844063.

**[Cu(xantphosMes2)[bpy]][PF6]**

C35H52Cu6F6N2O3s·P21, M = 1027.49, yellow block, monoclinic, space group C2/c, a = 38.341(2), b = 11.8342(7), c = 26.5208(14) Å, β = 126.4524(19)°, U = 9679.2(10) Å3, Z = 8, Dc = 1.410 Mg m–3, μ(Cu-Ko) = 2.126 mm–1, T = 123 K. Total 29726 reflections, 8726 unique, Rint = 0.027. Refinement of 7795 reflections (613 parameters) with I > 2σ(I) converged at final R1 = 0.0320 (R1 all data = 0.0362), wR2 = 0.0851 (wR2 all data = 0.0870), GOF = 1.0356. CCDC 1844062.

**[Cu(xantphosMes2)(6-Mebpy)][PF6]**

C39H64Cu6F6N2O3s·P21, M = 1041.51, yellow block, monoclinic, space group C2/c, a = 38.1932(19), b = 12.0648(6), c = 26.5270(13) Å, β = 127.0427(19)°, U = 9756.6(9) Å3, Z = 8, Dc = 1.418 Mg m–3, μ(Cu-Ko) = 2.117 mm–1, T = 123 K. Total 31926 reflections, 8749 unique, Rint = 0.033. Refinement of 7701 reflections (685 parameters) with I > 2σ(I) converged at final R1 = 0.0467 (R1 all data = 0.0531), wR2 = 0.0586 (wR2 all data = 0.0623), GOF = 0.9991. CCDC 1844061.

Computational details

A set of density functional theory (DFT) calculations were performed for the [Cu(P(=P)[N=N])2]+ cations (P(=P) = Bu2xantphos and xantphosMes2; N=N = bpy, 6-Mebpy, and 6,6'-Me-bpy), using the A.03 revision of Gaussian 16.46 The Becke’s three-parameter B3LYP exchange–correlation functional17,48 was used in all the calculations. The “double-ζ” quality def2svp basis set was employed for C, H, P, N and O atoms, whereas the “triple-ζ” quality def2tvp basis set was used for the Cu atom.49,50 Intramolecular non-covalent interactions are expected to play a relevant role in determining the molecular geometry of the studied complexes owing to the presence of the bulky xantphos-derived ligands. To get a better description of those interactions, the D3 Grimme’s dispersion term with Becke–Johnson damping was added to the B3LYP functional (B3LYP-D3).51,52 The geometries of all the complexes in both their singlet ground electronic state (S0) and their lowest-energy triplet excited state (T1) were optimized without imposing any symmetry restriction. For T1 the spin unrestricted UB3LYP approximation was used with a spin multiplicity of three. The lowest-lying excited states of each complex, both singlets and triplets, were computed at the minimum-energy geometry optimized for S0 using the time-dependent DFT (TD-DFT) approach.53–55 All the calculations were performed in the presence of the solvent (CH2Cl2). Solvent effects were considered within the self-consistent reaction field (SCRF) theory using the polarized continuum model (PCM) approach.56–58

Device preparation and characterization

LEC s were prepared on top of patterned indium tin oxide (ITO, 15 Ω sq1) coated glass substrates previously cleaned by chemical and UV-ozon methods. Prior to the deposition of the emitting layer, 80 nm thick films of poly-(3,4-ethylenedioxythiophene):poly(styrenesulfonate) (PEDOT:PSS) (CLEVIOSTM P VP AI 4083, Heraeus) were coated in order to flatten the ITO electrode and to increase its work function. The emitting layer (100 nm thick) was prepared by spin-coating a dichloromethane solution of the emitting compound with the addition of the ionic liquid 1-ethyl-3-methylimidazolium hexafluorophosphate [Emim][PF6] (>98.5%, Sigma-Aldrich), in a 1:4 molar ratio. The devices were then transferred to an inert atmosphere glovebox (<0.1 ppm O2 and H2O), where the aluminium cathode (100 nm) was thermally deposited in high
vacuum using an Edwards Auto500 chamber integrated in the glovebox. The thickness of all films was determined with an Ambios XP-1 profilometer. The active area of the devices was 6.5 mm². LECs were not encapsulated and were characterized inside the glovebox at room temperature. The device lifetime was measured by applying a pulsed current and monitoring the voltage and luminance versus time by a True Colour Sensor MAZeT (MTCSiCT Sensor) with a Botest OL1 OLED Lifetime-Test System. The electroluminescence (EL) spectra were measured using an Avantes AvaSpec-2048 Fiber Optic Spectrometer during device lifetime measurement.

Results and discussion

Preparation and characterization of P^P ligands

The P^P ligand ^Bu_xantphos was synthesized using the literature procedure. The synthetic routes to xantphosMes_4 and xantphosMes_2 were based on the strategy of Hamann and Hartwig to prepare bidentate phosphines with varying steric properties. The syntheses of xantphosMes_4 and xantphosMes_2 are summarized in Scheme 2. Both compounds were isolated as white solids, but facile oxidation to the phosphane oxides made it difficult to obtain analytically pure samples. The electrospray (ESI) mass spectra of xantphosMes_4 and xantphosMes_2 showed base peaks at m/z 747.3 and 663.5, respectively, arising from the [M + H]⁺ ions. The solid-state IR spectra of ^Bu_xantphos and xantphosMes_2 are shown in Fig. S1 and S2. Single crystals of xantphosMes_2 were grown from an Et_2O solution of the compound by slow evaporation. Fig. S3† shows an ORTEP-style plot of the molecule and important bond parameters are given in the figure caption. Few chiral phosphane-derived ligands have been reported in the literature, and the solid-state structure of xantphosMes_2 represents the (rac)-form of xantphosMes_2. It crystallizes in the monoclinic P2_1/a space group with both the (R,R)- and (S,S)-enantiomers in the unit cell; the (S,S)-enantiomer is shown in Fig. 1. The xanthene unit deviates very slightly from planarity, in contrast to the ‘bowl’ shape that is commonly adopted (see discussion below).

The ^31P NMR spectra of xantphosMes_4 and xantphosMes_2 exhibit resonances at δ −36.2 and −25.8 ppm, respectively, consistent with one phosphorus environment in each compound. ^1H and ^13C NMR spectra (see Experimental section in the ES†) were assigned by 2D methods and were in accord with functionalization in the 4,5-positions of the xanthene unit (Scheme 2). The ^1H NMR spectra are shown in Fig. S4 and S5†. The ^1H NMR spectrum of xantphosMes_2 (Fig. S5†) also shows the presence of a subspecies in solution, present in <10% based on integration. The chemical shifts of the low intensity signals and the presence of diagnostic NOESY peaks suggest the major and minor species are structurally related, and we assign them to the (rac)- and (meso)-forms, respectively. Based on the preference seen in the solid-state, we propose that the dominant species is the (rac)-form. Thus, the bisphos- phane is preorganized to give particular diastereoisomers upon complexation with copper(i) and this indeed is the case as discussed later.

Preparation and characterization of copper(i) complexes

Attempts to prepare [Cu(xantphosMes_4)(bpy)][PF_6], [Cu(xantphosMes_4)(6-Mebp)][PF_6] and [Cu(xantphosMes_4)(6,6'-Mebpy)][PF_6] by reaction of [Cu(MeCN)_4][PF_6] and xantphosMes_4 followed by the corresponding 2,2'-bipyridine ligand failed to yield the desired heteroleptic complexes. The ^31P spectrum of each crude reaction mixture was dominated by a peak envelope at δ ~36.2 ppm, and also exhibited several other unassigned signals. Evidence for the formation of [Cu(xantphosMes_4)]⁺ came from ESI mass spectrometry with a peak envelope at m/z 809.5. Mononuclear [Cu(P^P)]⁺ complexes containing sterically demanding substituents attached to the phosphorus atoms are known, for example [Cu(Bu-xantphos)x^10.P][PF_6]¹⁷ ([Bu-xantphos = 9,9-dimethyl-4,5-bis(di- tert-butlyphosphino)xanthene] and 3-coordinate copper(i) complexes with P^N^P pincer ligands. Since xantphosMes_4 proved to be too sterically demanding for the formation of [Cu(xantphosMes_4)(N^N)]⁺ species, we turned our attention to the use of xantphosMes_2.

Heteroleptic [Cu(P^P)(N^N)]⁺ complexes with P^P = ^Bu_xantphos or xantphosMes_2 were prepared using the established procedure²⁴ by the addition of a mixture of the xanthos and bpy ligands to a solution of [Cu(MeCN)_4][PF_6] in

Scheme 2 Syntheses of xantphosMes_4 and xantphosMes_2. Conditions: (i) t-BuLi, dry heptane, TMEDA, reflux, 20 min; (ii) Mes_2PCI, THF, 0 °C, 1 h; (iii) MesPhPCL, THF, 0 °C, 1 h.

Fig. 1 Crystallographically determined structure of the (S,S)-enantio- mer of xantphosMes_2. See also Fig. S3†.
CH₂Cl₂. This procedure avoids competitive formation of homoleptic [Cu(P^P)₃]⁺ complexes.⁶³ Homoleptic [Cu(N^N)₂]⁺ complexes where N^N is a bpy derivative, and heteroleptic [Cu(xantphos[N^N])₂]⁺ species are typically red and yellow respectively. In all cases, the reaction mixture remained orange after being stirred for 1–2 hours, suggesting incomplete conversion to the heteroleptic complex. To force full conversion, an additional 0.2 equivalents of the biphosphine ligand were added, resulting in the solution turning yellow. After being stirred for a further 2.5 hours, the solvent was removed and the excess ligand was removed by washing with Et₂O. This procedure yielded [Cu(Bu₂xantphos)[bpy]][PF₆], [Cu(Bu₂xantphos)(6-Me bpy)][PF₆], [Cu(xantphosMes₂)(bpy)][PF₆], [Cu(xantphosMes₂)(6,6′-Me₂bpy)][PF₆], [Cu(xantphosMes₂)(6-Mebpy)][PF₆], and [Cu(xantphosMes₂)(6-Mebpy)][PF₆] as yellow powders. In the reaction between [Cu(MeCN)₄][PF₆], xantphosMes₂ and 6,6′-Me₂bpy, the yellow colour corresponding to the heteroleptic complex was not observed and only a red solid, identified by NMR spectroscopy as [Cu(6,6′-Me₂bpy)]+[PF₆], could be isolated. We suggest that the steric demands of the substrates in 6,6′-Me₂bpy combined with the two mesityl groups in xantphosMes₂ mitigate against the formation of [Cu(xantphosMes₂)(6,6′-Me₂bpy)][PF₆].

The five heteroleptic compounds were characterized by ¹H, ¹³C and ³¹P NMR spectroscopies, elemental analysis, ESI mass spectrometry and IR spectroscopy (see Fig. S6–S10†), as well as representative single crystal structures. The ESI mass spectrum of each complex containing ‘Bu₂xantphos exhibited a base peak corresponding to the [M–PF₆]⁺ ion. The ESI mass spectrum of [Cu(xantphosMes₂)[bpy]][PF₆] showed a base peak envelope at m/z 725.4 assigned to [Cu(xantphosMes₂)]⁺ and a lower intensity peak envelope at m/z 881.5 arising from [M–PF₆]⁻. For [Cu(xantphosMes₂)(6-Mebpy)][PF₆], the ESI mass spectrum exhibited only a peak envelope for the [Cu(xantphosMes₂)]⁺ ion (m/z 725.4). Elemental analysis and the NMR spectra confirmed the formation of a heteroleptic complex. The solution NMR spectroscopic properties are discussed after the solid-state structures.

**Single crystal structures of copper(I) complexes**

Single crystals of [Cu(Bu₂xantphos)[bpy]][PF₆]-0.5Et₂O, [Cu(Bu₂xantphos)(6-Me bpy)][PF₆]-1.5CH₂Cl₂-0.5H₂O, [Cu(xantphosMes₂)[bpy]][PF₆] and [Cu(xantphosMes₂)(6-Mebpy)][PF₆] were obtained by layering Et₂O over a CH₂Cl₂ solution of each complex. The ‘Bu₂xantphos-containing compounds crystallized in the triclinic space group P₁, whereas [Cu(xantphosMes₂)(bpy)][PF₆] and [Cu(xantphosMes₂)(6-Mebpy)][PF₆] crystallized in the monoclinic space group C2/c. Fig. S11–S14† show ORTEP-style plots of the [Cu(Bu₂xantphos)(bpy)]⁺, [Cu(Bu₂xantphos)(6-Mebpy)]⁺, [Cu(xantphosMes₂)(bpy)]⁻ and [Cu(xantphosMes₂)(6-Mebpy)]⁻ cations. Selected structural parameters are given in the captions to Fig. S11–S14† and important parameters are compared in Table 1. In each cation, the copper(i) centre is in a distorted tetrahedral geometry and the xanthene backbone adopts a ‘bowl’ (boat) conformation, which is observed for the majority of free and coordinated xantphos ligands. A search of the Cambridge Structural Database (CSD) showed no relevant structurally analogous complexes.

On going from [Cu(P^P)(bpy)]⁺ to [Cu(P^P)(6-Mebpy)]⁺, the introduction of the 6-methyl group in the bpy unit lowers the symmetry of the cation. The methyl substituent can, in principle, lie over the xanthene ‘bowl’ or be remote from it.²⁶ Fig. 2a shows that in [Cu(Bu₂xantphos)(6-Mebpy)][PF₆], the methyl group lies over the xanthene unit. In contrast, in [Cu(xantphosMes₂)(6-Mebpy)][PF₆], the 6-Mebpy ligand is orienta-
tionally disordered over two sites, each with 50\% occupancy. Fig. 2b depicts the conformer in which the Me group is remote from the xanthene unit; the second conformer is structurally related to [Cu(tBu₂xantphos)(bpy)][PF₆] \( \text{[Fig. 2a].} \) The N–C–C–N torsion angles in Table 1 demonstrate that the bpy unit is significantly more twisted in the cations containing the xantphosMes₂ ligand than those with tBu₂xantphos. This appears to be associated with the fact that in both [Cu(xantphosMes₂)(bpy)][PF₆] \( \text{[Fig. 3] and [Cu(xantphosMes₂)(6-Mebpy)][PF₆],} \) one methyl group of one mesityl substituent is directed towards the middle of the bpy domain (Fig. 3b). This spatial proximity is characterized by \( C_{\text{Me(Mes)}} \text{⋯centroidpyridine} = 3.98 \) and \( 4.37 \) Å (\( H_{\text{Me(Mes)}} \text{⋯centroidpyridine} = 3.16 \) and \( 3.56 \) Å) in the [Cu(xantphosMes₂)(bpy)]\( ^{+} \) cation. Corresponding separations in [Cu(xantphosMes₂)(6-Mebpy)]\( ^{+} \) are \( 4.00 \) and \( 4.38 \) Å (\( 3.24 \) and \( 3.57 \) Å) and \( 3.85 \) and \( 4.39 \) Å (\( 3.27 \) and \( 3.48 \) Å) for the two partial occupancy 6-Mebpy sites.

**Solution NMR spectroscopy**

The aromatic regions of the solution \(^1\)H NMR spectra of [Cu(tBu₂xantphos)(bpy)][PF₆], [Cu(xantphosMes₂)(bpy)][PF₆] and [Cu(tBu₂xantphos)(6,6′Me₂bpy)][PF₆] are shown in Fig. 4 (see also Fig. S15\( \dagger \), S16 and S17\( \dagger \) respectively. All signals are sharp and well-resolved at room temperature and were assigned using COSY, NOESY, HMBC and HMOC methods. The tBu substituents give rise to a singlet at \( \delta = 1.11 \) ppm in [Cu(tBu₂xantphos)(bpy)][PF₆] and \( \delta = 1.16 \) ppm in [Cu(tBu₂xantphos)(6,6′Me₂bpy)][PF₆]. Fig. 4 and S16\( \dagger \) reveal that the two pyridine rings of bpy in [Cu(tBu₂xantphos)(bpy)][PF₆] and [Cu(xantphosMes₂)(bpy)][PF₆] are magnetically equivalent. Similarly, the 6,6′-Me₂bpy ligand in [Cu(tBu₂xantphos)(6,6′-Me₂bpy)][PF₆] is symmetric on the NMR timescale at room temperature (Fig. S17\( \dagger \)). We have previously detailed a solution dynamic behaviour for [Cu(xantphos)(N^N)]\( ^{+} \) complexes involving inversion of the xanthene unit (‘bowl’),\(^{26,32}\) and this is depicted in the first dynamic process illustrated in Fig. 5. The scheme demonstrates that inversion of the xanthene bowl exchanges the environments of

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**Fig. 3** Structure of the [Cu(xantphosMe₂)(bpy)]\( ^{+} \) cation (a) emphasizing the relative positions of the phenyl and mesityl substituents with respect to the bpy unit, and (b) with the bpy (blue) and one mesityl group (red) shown in space-filling representation to emphasize their spatial proximity.

**Fig. 4** Aromatic region of the \(^1\)H NMR spectrum (500 MHz, acetone-\(d_6\)) of [Cu(tBu₂xantphos)(bpy)][PF₆]. See Fig. S15\( \dagger \) for the full spectrum. See Scheme 3 for atom labelling.

**Fig. 5** Upper: Proposed dynamic processes in [Cu(xantphos)(bpy)]\( ^{+} \) type compounds. \( \text{N}_A \) and \( \text{N}_B \) represent pyridine rings A and B. For tBu₂xantphos, \( \text{R}_1 = \text{R}_2 = \text{Ph}. \) For xantphosMes₂, \( \text{R}_1 = \text{Mes}, \text{R}_2 = \text{Ph}. \) Lower: Crystallographically determined structure of the [Cu(xantphosMes₂)(6-Mebpy)]\( ^{+} \) cation in which the 6-Mebpy is orientationally disordered: (a) orientation 1 of 6-Mebpy, and (b) orientation 2 of 6-Mebpy shown in two views of the cation (see text).
the xanthphos methyl groups (green and magenta in Fig. 5) between axial and equatorial sites, but does not render pyridine rings A and B (represented by NA and NB) equivalent. By invoking a second process involving movement of the \( \{\text{Cu(bpy)}\} \) unit (i.e., a change in conformation of the chelate ring, Fig. 5), NA and NB experience both sides of the xanthene bowl and are equivalent on the NMR timescale.

On going from \([\text{Cu(Bu}_2\text{xantphos}](\text{bpy})][\text{PF}_6]\) to \([\text{Cu(Bu}_2\text{xantphos}(6-\text{Mebpy})][\text{PF}_6]\), the symmetry of the cation is lowered and phenyl rings D (see Scheme 3) split into two sets, those proximate to the methyl group of 6-Mebpy and those on the side of the unsubstituted pyridine ring (Fig. 2a). Fig. 6 shows the aromatic region of the solution \(^1\text{H}\) NMR spectrum of \([\text{Cu(Bu}_2\text{xantphos}(6-\text{Mebpy})][\text{PF}_6]\), in which the sets of D rings are labelled D and D’. In the NOESY spectrum at 298 K, exchange (EXSY) peaks are observed between pairs of signals for protons D2/D2’ and D3/D3’; the D4/D4’ EXSY peaks appear too close to the diagonal in the NOESY spectrum to be clearly resolved. NOESY cross peaks (no EXSY) are observed between MeCq1 and MeCq1’ (Fig. S18†). These observations are consistent with inversion of the chelate ring (‘copper flip’ in Fig. 5) and no inversion of the xanthene bowl.

The single crystal structures of \([\text{Cu(xantphosMes}_2](\text{bpy})][\text{PF}_6]\) and \([\text{Cu(xantphosMes}_2](6-\text{Mebpy})][\text{PF}_6]\) reveal that the two PPhMes groups of the xanthphosMes\(_2\) ligand are mutually oriented as shown in Fig. 3a and 5. This desymmetrizes the xanthene unit (labelled rings C and C’). In addition, the equatorial and axial positions of the Ph and Mes substituents with respect to the chelate ring leads to chemical shift differences in the \(^1\text{H}\) NMR spectrum for pairs of phenyl rings (D and D’) and mesityl groups (E and E’). Fig. S19† shows the \(^1\text{H}\) NMR spectrum of \([\text{Cu(xantphosMes}_2](6-\text{Mebpy})][\text{PF}_6]\), and Fig. 7 and S20† show exchange peaks observed in the NOESY spectrum. Exchange peaks between the signals for phenyl proton D2/D2’ and D4/D4’ is consistent with the ‘copper flip’ shown in Fig. 5. This leads to equivalence of the outer rings of the xanthene unit as confirmed by the EXSY peak between the signals for protons C3/C3’. The EXSY peak between signals for the xanthene methyls MeCq1 and MeCq1’ (Fig. S20†) confirms the inversion of the xanthene bowl (Fig. 5). This contrasts with \([\text{Cu(Bu}_2\text{xantphos}(6-\text{Mebpy})][\text{PF}_6]\) where no exchange (only NOESY) peaks are observed (see above and Fig. S18†).

Earlier, we noted an orientational disorder of the 6-Mebpy ligand in the solid-state structure of \([\text{Cu(xantphosMes}_2]\)
(6-Mebpy)][PF₆]}. The disorder was modelled with a 50% occupancy of each orientation and Fig. 5a and b show the [Cu(xantphosMes₂)(6-Mebpy)]⁺ with the two orientations of 6-Mebpy. The structure in Fig. 5a corresponds to the top diagram in the scheme in Fig. 5, while Fig. 5b corresponds to the bottom diagram in the scheme. The disorder therefore parallels a combination of the two dynamic processes which we propose the cation undergoes in solution.

Variable temperature (VT) NMR spectra were recorded for an acetone-d₆ solution of [Cu(xantphosMes₂)(6-Mebpy)][PF₆]. The ³¹P NMR spectrum (Fig. S21) shows only one signal over the range 298–180 K. Fig. 8 shows the effect of temperature on the alkyl region of the ¹H NMR spectrum of [Cu(xantphosMes₂)(6-Mebpy)][PF₆]. The collapse of the signals for mesityl-methyl protons MeE₂ and MeE₂' and the appearance of four signals for these methyls below 218 K are consistent with freezing out the rotation of the mesityl groups. A similar temperature dependence is observed for the mesityl E₃ protons in the aromatic region of the spectrum (Fig. S22). Significant shifting of the xanthene methyl protons MeCq₁ and MeCq₁' (Fig. 8) and 6-Mebpy protons A₆ and MeB₆ (Fig. S22) can be attributed to changes in their magnetic environments as the mesityl groups adopt a static configuration. Both ³¹P and ¹H VT NMR spectra are consistent with the presence of only one conformer in solution.

Electrochemistry

The electrochemical behaviour of the [Cu(N^N)(P^P)][PF₆] complexes was studied using cyclic voltammetry and the results are summarized in Table 2 and Fig. 9. All compounds with the exception of [Cu(xantphosMes₂)(bpy)][PF₆] exhibit a quasi-reversible process in the range +0.76 to +0.85 V, which is assigned to a copper-centred oxidation. For [Cu(xantphosMes₂)](bpy)][PF₆] the oxidation at +0.80 V is irreversible. Although the corresponding reduction peak could not be resolved, the position of this oxidation peak (E_loop) is similar to the one observed for [Cu(Bu₂xantphos)(bpy)][PF₆] (+0.80 V, Fig. 9).

Table 2 Cyclic voltammetric data for [Cu(N^N)(P^P)][PF₆] complexes referenced to internal Fc/Fc⁺ = 0 V; degassed HPLC grade CH₂Cl₂ solution with [Bu₄N][PF₆] as supporting electrolyte and a scan rate of 0.1 V s⁻¹. Processes are quasi reversible unless otherwise stated (ir = irreversible). Data for [Cu(xantphos)(N^N)][PF₆].

<table>
<thead>
<tr>
<th>Complex cation</th>
<th>E_loop/N (E_pc – E_pn/mV)</th>
<th>E_loop/N</th>
<th>E_pool/N</th>
</tr>
</thead>
<tbody>
<tr>
<td>[Cu(Bu₂xantphos)(bpy)]⁺</td>
<td>+0.76 (90)</td>
<td>–2.20</td>
<td></td>
</tr>
<tr>
<td>[Cu(Bu₂xantphos)(6-Mebpy)]⁺</td>
<td>+0.83 (90)</td>
<td>–2.22</td>
<td></td>
</tr>
<tr>
<td>[Cu(Bu₂xantphos)(6,6-Mebpy)]⁺</td>
<td>+0.85 (100)</td>
<td>–2.28</td>
<td></td>
</tr>
<tr>
<td>[Cu(xantphosMes₂)(bpy)]⁺</td>
<td>+0.84 (70)</td>
<td>+0.80†</td>
<td></td>
</tr>
<tr>
<td>[Cu(xantphos)(bpy)]⁺</td>
<td>+0.76 (110)</td>
<td>–2.28</td>
<td></td>
</tr>
<tr>
<td>[Cu(xantphos)(6-Mebpy)]⁺</td>
<td>+0.85 (100)</td>
<td>–2.28</td>
<td></td>
</tr>
<tr>
<td>[Cu(xantphos)(6,6-Mebpy)]⁺</td>
<td>+0.90 (150)</td>
<td>–2.28</td>
<td></td>
</tr>
</tbody>
</table>

† Values taken from ref. 24.

When scanning beyond +1.0 V, additional oxidation waves appear and the copper oxidation is no longer reversible. Instead a reduction signal appears at around +0.25 V. This indicates decomposition of the complexes at potentials higher than +1.0 V. Complexes containing Bu₂xantphos also show an irreversible reduction wave at around –2.2 V (Table 2) arising from a ligand-based process. For the complexes containing xantphosMes₂, no reduction processes were observed within the solvent accessible window.

Along the series [Cu(Bu₂xantphos)(N^N)][PF₆] with N^N = bpy to 6-Mebpy to 6,6-Mebpy, the copper oxidation shifts to higher potentials (as observed for the analogous [Cu(xantphos)(N^N)][PF₆] series, Table 2). This trend was also observed for a related series of compounds and has been rationalized using DFT calculations.
Theoretical calculations: geometry and molecular orbitals

The geometries of all the \([\text{Cu}^\prime\text{P}^\prime\text{P}][\text{N}^\prime\text{N}]\) cations in their electronic ground state \((S_0)\) were optimized at the DFT B3LYP-D3/ (def2svp + def2tvp) level in the presence of the solvent \((\text{CH}_2\text{Cl}_2)\) and without imposing any symmetry restriction (see the Experimental section for full computational details). The values obtained for the most representative geometrical parameters defining the \(\text{Cu}(i)\) coordination sphere are summarized in Table S1.† Calculations successfully reproduce the distorted tetrahedral structures observed experimentally around the \(\text{Cu}\) centre for all the studied complexes. Compared with the values from X-ray diffraction, the \(\text{Cu}–\text{N}\) and \(\text{Cu}–\text{P}\) bond distances and the \(\text{P}–\text{Cu}–\text{P}\) and \(\text{N}–\text{Cu}–\text{N}\) chelating angles are calculated with errors below 0.05 Å and 2°, respectively. The angle formed by the \(\text{P}–\text{Cu}–\text{P}\) and \(\text{N}–\text{Cu}–\text{N}\) planes, which can be used as an indication of the deviation from the orthogonal disposition of the \(\text{P}^\prime\text{P}\) and \(\text{N}^\prime\text{N}\) ligands, has values lying between 82 and 89°, in good agreement with those observed experimentally and with those computed in previous works for similar complexes.24,26,66 The \(\text{N}–\text{C}–\text{C}–\text{N}\) torsion angles remain in a small range between −10 and 14° indicating that the bpy ligand is essentially planar in all the cations. In contrast to the experimental results (Table 1), the cations containing the xantphosMes\(_2\) ligand do not feature significantly more twisted bpy ligands (Table S1†), suggesting that the packing forces play an important role in determining the structure in the crystal. Theoretical calculations correctly reproduce the longer \(\text{Cu}–\text{P}\) bond distances observed for these complexes, and the spatial proximity of the equatorial mesityl group to the bpy unit shown in Fig. 3. The differences in the torsion angles between theoretical and X-ray geometries are probably due to the fact that the former are obtained for an isolated molecule optimized in solution and do not take into account the packing forces and intermolecular interactions acting in the solid state.

Two geometry minima were found for \([\text{Cu}(\text{xantphosMes}_2)(6\text{-Me}bpy)]^+\), which show a different relative orientation of the 6-Mebpy ligand and correspond to the two conformations observed in the single-crystal structure determination (Fig. 5). They possess close energies, the conformation with the 6-methyl group lying over the xanthene bowl being more stable with the \(\text{Me}\) group away from the xanthene unit by only 0.28 kcal mol\(^{-1}\). This is in good agreement with the occupancy of 50% experimentally found for each conformation as discussed above.

The geometry of the first triplet excited state \((T_1)\) was also optimized at the UB3LYP level for all the \([\text{Cu}^\prime\text{P}^\prime\text{P}][\text{N}^\prime\text{N}])^+\) cations, and the most significant geometry parameters are also included in Table S1.† The molecular geometries in the \(T_1\) state significantly differ from those in the ground state \(S_0\). As discussed below, the \(T_1\) state implies a charge transfer from a molecular orbital that mainly involves a \(d\) orbital of the \(\text{Cu}\) atom to a molecular orbital spreading over the bpy ligand. Consequently, the metal atom is partially oxidized and tends to adopt the square-planar coordination sphere expected for four-coordinate \(d^9\) \(\text{Cu}(i)\) complexes, instead of the tetrahedral conformation typical of \(d^{10}\) \(\text{Cu}(i)\) coordination complexes. This effect can be studied by following the changes in the angle formed by the \(\text{N}–\text{Cu}–\text{N}\) and \(\text{P}–\text{Cu}–\text{P}\) planes, which decreases in going from \(S_0\) to \(T_1\) as the molecule becomes more planar (Table S1†). The distortions from the tetrahedral structure in going from \(S_0\) to \(T_1\) is indeed limited by number of methyl groups in the 6,6′-positions of the bpy ligand, because substituents in these positions impede the movement of the ligands towards more planar dispositions.24 In this way, the \([\text{Cu}^\prime(\text{Bu}_2\text{xantphos})(\text{bpy})]^+\) complex, with no substituent in the 6,6′-positions, shows the largest reduction \((25.6°)\) passing from 82.8° in \(S_0\) to 57.2° in \(T_1\). The \([\text{Cu}^\prime(\text{Bu}_2\text{xantphos})(6\text{-Me}bpy)]^+\) complex, including one \(\text{Me}\) group in the 6-position, undergoes a smaller reduction of 23.3° \((\text{from 87.9 to 64.6°})\), and the \([\text{Cu}^\prime(\text{Bu}_2\text{xantphos})(6,6\text{-Me}_2\text{bpy})]^+\) complex, featuring \(\text{Me}\) substituents in both the 6- and 6′-positions, shows a reduction of only 15.4° \((\text{from 86.1 to 70.7°})\). Thus, the presence of \(\text{Me}\) groups in the 6- and 6′-positions strongly affects the degree of geometrical relaxation of the \(T_1\) excited state, and limits its stabilization. The energy position of the \(T_1\) state relative to \(S_0\) and thereby the emission properties of the complexes, therefore depend not only on the electron-donating or electron-withdrawing character of the substituent groups present in the ligands but also on the positions where the substituents are introduced and on the structural effects they induce.

Fig. 10 shows the evolution of the energy calculated for the highest-occupied (HOMO) and lowest-unoccupied molecular
orbital (LUMO) of the five complexes. The atomic orbital composition of the molecular orbitals remains almost unchanged along the series and only the contour plots computed for [Cu (‘Bu₂xantphos)(bpy)]⁺ are displayed as a representative example. As reported previously for similar copper(I) complexes,²⁴,²⁶,⁶⁶ the HOMO appears mainly centred on the metal with a small contribution from the phosphorus atoms, and the LUMO spreads over the bpy ligand. The addition of ‘Bu groups to the xantphos moiety has a negligible effect on the energy of the HOMO, and the [Cu(‘Bu₂xantphos)(N=Ν)]⁺ (N=Ν = bpy, 6-Me bpy and 6,6'-Me₂bpy) complexes have a slightly higher HOMO energy, around −5.97 eV (Fig. 10), than that calculated for the reference complex [Cu(xantphos)(bpy)]⁺ (−6.00 eV) at the same theoretical level.⁶⁶ Replacement of PPh₂ moieties by PhMes groups in the xantphos ligand has a more significant effect moving up the HOMO of [Cu(xantphosMes₂) (bpy)]⁺ and [Cu(xantphosMes₂)(6-Me bpy)]⁺ to −5.81 and −5.89 eV, respectively. There are small changes in this energy within each series, as the structural changes introduced in the complexes are made in regions where the HOMO is not centered. The HOMO energies, even considering the small changes described, are quite close, in good agreement with the close \( E_{1/2} \) values reported for the complexes in the Electrochemistry section. The LUMO undergoes a small destabilization when a Me group (a weak electron donor) is added to the bpy ligand, a destabilization that becomes more pronounced when a second group is added. The HOMO–LUMO energy gap increases in the [Cu(‘Bu₂xantphos)(N=Ν)]⁺ series with the number of Me substituents of the bpy ligand in agreement with the experimental CV data (Table 2). It is therefore expected that excited states described by HOMO → LUMO transitions appear at bluer wavelengths as more Me groups are attached to the N=Ν ligand.

**Photophysical properties**

The solution absorption spectra of the [Cu(P=P)(N=Ν)] [PF₆] complexes in CH₂Cl₂ are shown in Fig. 11. The intense bands below 330 nm are assigned to ligand-based n → π* and π → π* transitions and vary little within a series of a certain phosphine ligand. The broad absorption at around 380 nm (Table 3) arises from a metal-to-ligand charge transfer (MLCT) excitation. In the series [Cu(‘Bu₂xantphos)(N=Ν)] [PF₆], the MLCT absorption slightly shifts to higher energies on going from bpy to 6-Me bpy and to 6,6'-Me₂bpy (see Fig. S23†) as has previously been observed for the analogous [Cu(xantphos) (N=Ν)] [PF₆] series.²⁴ This behaviour is also observed in passing from [Cu(xantphosMes₂)(bpy)] [PF₆] to [Cu(xantphosMes₂) (6-Me bpy)] [PF₆], and is in good agreement with the electrochemical properties discussed above.

To get a better understanding of the nature of the electronic excited states involved in the absorption spectra, the time-dependent DFT (TD-DFT) approach was used to study the lowest lying singlet (S₀) and triplet (T₁) excited states. Table 4 collects the energies and oscillator strengths (f) computed for the S₀ and T₁ states of all the complexes, together with those obtained for [Cu(xantphos)(bpy)]⁺ included as a reference. In all cases, both the S₁ and the T₁ state result from the HOMO → LUMO monoexcitation, with a contribution exceeding 95%. This excitation implies an electron transfer from the Cu(P=P) moiety of the complex to the bpy ligand, supporting the MLCT character of S₁ and T₁. The broad absorption band observed experimentally at around 380 nm therefore originates in the S₀ → S₁ transition, and the calculated values, although somewhat displaced to the red, reproduce the experimentally observed shift of the absorption maxima to bluer wavelengths as the number of Me substituents of the bpy ligand increases. This is also in good accord with the increase of the HOMO–LUMO gap along each series of complexes predicted above (Fig. 10).

The powder and solution emission spectra are displayed in Fig. 12 and S24† respectively, and data are given in Table 3. Upon excitation into the MLCT region (\( \lambda_{exc} = 365 \) nm), all the compounds show an emission in the orange to yellow region. On going from [Cu(‘Bu₂xantphos)(bpy)] [PF₆] to [Cu(‘Bu₂xantphos)(6-Me bpy)] [PF₆] to [Cu(‘Bu₂xantphos)(6,6'-Me₂bpy)] [PF₆], or from [Cu(xantphosMes₂)(bpy)] [PF₆] to [Cu(xantphosMes₂)(6-Me bpy)] [PF₆], the introduction of additional methyl groups in the bpy ligand shifts the emission to higher energies (Table 3). Additionally, the photoluminescent quantum yield (PLQY) increases for solution and especially for powder emission along the series (Table 3). [Cu(‘Bu₂xantphos) (bpy)] [PF₆] and [Cu(xantphosMes₂)(bpy)] [PF₆], just like [Cu(xantphos)(bpy)] [PF₆],²⁴ exhibit low PLQY values, especially in solution. This is most probably due to the accessibility of the copper centre, which leads to solvent quenching of the excited state. Consequently, PLQY values in the powder samples are higher than in solution. The emission in the solid state is in all cases blue-shifted compared to the solution emission. This trend has also been observed for related complexes containing POP or xantphos.²⁵,²⁶

The theoretical results reproduce the trends observed in the experimental emission spectra. The emitting T₁ state shifts to
Table 3 Photophysical properties of the \([\text{Cu}(\text{P}^\text{P})(\text{N}^\text{N})][\text{PF}_6]\) complexes compared to \([\text{Cu}(\text{xantphos})(\text{N}^\text{N})][\text{PF}_6]\) (\(\text{N}^\text{N} = \text{bpy}, 6\text{-Mebpy}, 6,6'\text{-Me}_2\text{bpy}\))

<table>
<thead>
<tr>
<th>Complex cation</th>
<th>(S_1) (E (eV\ nm^{-1})) (f)</th>
<th>(T_1) (E (eV))</th>
</tr>
</thead>
<tbody>
<tr>
<td>([\text{Cu}(\text{xantphos})(\text{bpy})]^+)</td>
<td>2.816/440 (0.08)</td>
<td>2.569</td>
</tr>
<tr>
<td>([\text{Cu}(\text{Bu}_2\text{xantphos})(\text{bpy})]^+)</td>
<td>2.803/442 (0.09)</td>
<td>2.554</td>
</tr>
<tr>
<td>([\text{Cu}(\text{Bu}_2\text{xantphos})(6\text{-Mebpy})]^+)</td>
<td>2.823/439 (0.10)</td>
<td>2.577</td>
</tr>
<tr>
<td>([\text{Cu}(\text{Bu}_2\text{xantphos})(6,6'\text{-Me}_2\text{bpy})]^+)</td>
<td>2.923/424 (0.07)</td>
<td>2.694</td>
</tr>
<tr>
<td>([\text{Cu}(\text{xantphosMes}_2)(\text{bpy})]^+)</td>
<td>2.664/465 (0.06)</td>
<td>2.470</td>
</tr>
<tr>
<td>([\text{Cu}(\text{xantphosMes}_2)(6\text{-Mebpy})]^+)</td>
<td>2.750/451 (0.06)</td>
<td>2.582</td>
</tr>
</tbody>
</table>

Table 4 Vertical excitation energies (\(E\)) calculated at the TD-DFT B3LYP/(def2svp + def2tzvp) level for the lowest singlet (\(S_1\)) and triplet (\(T_1\)) excited states of complexes \([\text{Cu}(\text{N}^\text{N})(\text{P}^\text{P})]^+\) in CH$_2$Cl$_2$ solution. Oscillator strengths (\(f\)) are given within parentheses for the \(S_0 \rightarrow S_1\) transition.

<table>
<thead>
<tr>
<th>Complex cation</th>
<th>(E (eV))</th>
<th>(f)</th>
</tr>
</thead>
<tbody>
<tr>
<td>([\text{Cu}(\text{xantphos})(\text{bpy})][\text{PF}_6])</td>
<td>2.816</td>
<td>440</td>
</tr>
<tr>
<td>([\text{Cu}(\text{Bu}_2\text{xantphos})(\text{bpy})][\text{PF}_6])</td>
<td>2.803</td>
<td>442</td>
</tr>
<tr>
<td>([\text{Cu}(\text{Bu}_2\text{xantphos})(6\text{-Mebpy})][\text{PF}_6])</td>
<td>2.823</td>
<td>439</td>
</tr>
<tr>
<td>([\text{Cu}(\text{Bu}_2\text{xantphos})(6,6'\text{-Me}_2\text{bpy})][\text{PF}_6])</td>
<td>2.923</td>
<td>424</td>
</tr>
<tr>
<td>([\text{Cu}(\text{xantphosMes}_2)(\text{bpy})][\text{PF}_6])</td>
<td>2.664</td>
<td>465</td>
</tr>
<tr>
<td>([\text{Cu}(\text{xantphosMes}_2)(6\text{-Mebpy})][\text{PF}_6])</td>
<td>2.750</td>
<td>451</td>
</tr>
</tbody>
</table>

As discussed above, the geometry relaxation of the emitting \(T_1\) state leads to the flattening of the tetrahedral coordination environment. This flattening is more hindered as the number of methyl substituents attached to positions 6 and 6' of the bpy ligand is increased, and the relaxation of the \(T_1\) triplet is impeded thus leading to higher emission energies. Inspection of Table 3 shows that increasing the steric hindrance of the bpy ligand is beneficial for the emissive properties. Less flattening of the tetrahedral coordination environment of the copper centre gives rise to higher emission energies and, as
expected, longer excited state lifetimes and higher PLQYs. The same is true for the series of \([\text{Cu(xantphos)}(\text{N}^\text{N})][\text{PF}_6]\) (Table 3).

However, from a synthetic point of view, the unsuccessful attempt to \([\text{Cu(xantphosMes}_2)(6,6'\text{-Me}_2\text{bpy})][\text{PF}_6]\) demonstrates a limitation in the combined steric properties of the xantphosMes$_2$ and 6,6'-Me$_2$-bpy ligands. The photophysical properties of \([\text{Cu(tBu$_2$xantphos)}(\text{bpy})][\text{PF}_6]\) and \([\text{Cu(xantphosMes}_2)(\text{bpy})][\text{PF}_6]\) are similar both in solution and in the solid state (Table 3). In the case of the \([\text{Cu}(\text{P}^\text{P})(6\text{-Meppy})][\text{PF}_6]\) complexes, a similar behaviour is observed in the solid state emission. Both complexes are blue shifted compared to their \([\text{Cu}(\text{P}^\text{P})(\text{bpy})][\text{PF}_6]\) analogues but the emission profile is still very similar and the difference in peak position is again low (\(\Delta\lambda = 5\) nm). It is notable that the shift difference when going from \([\text{Cu}(\text{P}^\text{P})(\text{bpy})][\text{PF}_6]\) to \([\text{Cu}(\text{P}^\text{P})(6\text{-Meppy})][\text{PF}_6]\) is larger in the complexes containing xantphosMes$_2$ (\(\Delta\lambda = 42\) nm) compared to tBu$_2$xantphos complexes (\(\Delta\lambda = 32\) nm). On the other hand, the solution emission profiles differ significantly for the two \([\text{Cu}(\text{P}^\text{P})(6\text{-Meppy})][\text{PF}_6]\) complexes (Fig. S24†). The emission maximum of \([\text{Cu}(\text{tBu$_2$xantphos})(6\text{-Meppy})][\text{PF}_6]\) is blue shifted compared to \([\text{Cu}(\text{tBu$_2$xantphos})(\text{bpy})][\text{PF}_6]\) (\(\Delta\lambda = 47\) nm) as expected, but in the case of \([\text{Cu}(\text{xantphosMes}_2)(6\text{-Meppy})][\text{PF}_6]\) the emission maximum is blue-shifted very little compared to \([\text{Cu}(\text{tBu$_2$xantphos})(\text{bpy})][\text{PF}_6]\) (\(\Delta\lambda = 10\) nm). Both \([\text{Cu}(\text{tBu$_2$xantphos})(6\text{-Meppy})][\text{PF}_6]\) and \([\text{Cu}(\text{xantphosMes}_2)(6\text{-Meppy})][\text{PF}_6]\) also show a second unstructured emission feature at 628 nm and 623 nm respectively, which is not observed in the solid-state emission profile.

To investigate whether the compounds showed thermally activated delayed fluorescence (TADF) at room temperature, low temperature emission spectra and excited state lifetimes were recorded in frozen Me-THF at 77 K (Table 3 and Fig. 13). All complexes show a red shift in emission of 5-40 nm compared to the solid state emission and a greatly enlarged excited state lifetime. This indicates the possibility that all complexes are TADF emitters at room temperature. The energy difference between the lowest energy singlet and triplet excited states has been calculated to lie between 0.17 and 0.25 eV (Table 4), and is small enough to allow the occurrence of TADF processes.\(^{22}\)

Low temperature data further support the idea that the position of the emission bands of the \([\text{Cu}(\text{tBu$_2$xantphos})(\text{N}^\text{N})][\text{PF}_6]\) complexes is strongly affected by the flattening of the tetrahedral environment in the T$_1$ state. In solution, this flattening is not impeded and the difference in peak position between \([\text{Cu}(\text{tBu$_2$xantphos})(\text{bpy})][\text{PF}_6]\) and \([\text{Cu}(\text{tBu$_2$xantphos})(6\text{-Meppy})][\text{PF}_6]\) is 86 nm (0.30 eV) (Table 3). In powder, the flattening is more restricted and the difference decreases to 62 nm (0.25 eV). Finally, at 77 K, where the relaxation is even more impeded for all the complexes, the emission maxima range between 597 and 555 nm, in a window of just 42 nm (0.15 eV).

**Device performances**

The series of compounds was tested in LECs using ITO/PEDOT:PSS as the anode, an emitting layer consisting of the complex in the presence of [Emim][PF$_6$] (4:1 molar ratio) and an aluminium cathode. Devices were tested monitoring the electroluminescence and voltage over time, and were driven with a pulsed current (100 A m$^{-2}$ average, 50% duty cycle, 1 kHz). The main device parameters obtained for the entire samples series are reported in Table 5. The time evolution of the voltage and luminance for the LECs are reported in Fig. 14, together with the electroluminescence spectra. All LECs based on copper(i) complexes with tBu$_2$xantphos as the P$^\text{P}$ ligand show a fast luminescence turn-on time ($t_{\text{on}}$, defined here as the time to reach the maximum luminance), ranging from 1 minute for the complex with 6,6'-Me$_2$-bpy to 4.5 minutes for the one with 6-Meppy (Fig. 14b). The maximum luminance registered for these compounds increases with increasing substitution to the N$^\text{N}$ ligand, going from 20 cd m$^{-2}$ for the complex with unsubstituted bpy, to 230 cd m$^{-2}$ and 370 cd m$^{-2}$ for the ones containing 6-Meppy and 6,6'-Me$_2$-bpy, respectively. This trend follows that of the PLQY registered for the same compounds (Table 3) and is consistent with an augmented steric hindrance of the N$^\text{N}$ ligand, resulting in a higher stabilization of the tetrahedral complex geometry. The highest efficiency of 3.7 cd A$^{-1}$ for the LECs using \([\text{Cu}(\text{tBu$_2$xantphos})(6,6'\text{-Me}_2\text{bpy})][\text{PF}_6]\) corresponds to an external quantum efficiency of 1.0%, which is substan-

![Fig. 13 Normalized emission spectra of \([\text{Cu}(\text{P}^\text{P})(\text{N}^\text{N})][\text{PF}_6]\) complexes in Me-THF at 77 K (\(\nu_{\text{exc}} = 410\) nm).](image-url)

<table>
<thead>
<tr>
<th>Complex</th>
<th>$t_{\text{on}}$ (min)</th>
<th>$L_{\text{max}}$ (cd m$^{-2}$)</th>
<th>$t_{1/2}$ (min)</th>
<th>Efficiency (cd A$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>([\text{Cu}(\text{tBu$_2$xantphos})(\text{bpy})][\text{PF}_6])</td>
<td>1.1</td>
<td>19</td>
<td>5.1</td>
<td>0.2</td>
</tr>
<tr>
<td>([\text{Cu}(\text{tBu$_2$xantphos})(6\text{-Meppy})][\text{PF}_6])</td>
<td>4.5</td>
<td>230</td>
<td>53.8</td>
<td>2.3</td>
</tr>
<tr>
<td>([\text{Cu}(\text{tBu$_2$xantphos})(6,6'\text{-Me}_2\text{bpy})][\text{PF}_6])</td>
<td>1.0</td>
<td>370</td>
<td>4.9</td>
<td>2.3</td>
</tr>
<tr>
<td>([\text{Cu}(\text{xantphosMes}_2)(6\text{-Meppy})][\text{PF}_6])</td>
<td>0.7</td>
<td>50</td>
<td>34.6</td>
<td>0.5</td>
</tr>
</tbody>
</table>
tially lower as compared to the PLQY of the same compound in thin-film (23%). This implies that even in such bright device, non-radiative losses dominate the recombination of the injected electrons and holes. Interestingly, the quantum efficiencies for photo- and electro-luminescence of the [Cu(′Bu₂xanthphos)(6,6′-Me₂bpy)][PF₆] are very close to the unsubstituted xanthphos analogue (PLQY = 22% and maximum EQE = 1.0%),²⁶ highlighting the dominant role of the N′N ligands on the optical and optoelectronic properties of these compounds.

In general, the device lifetime (t₁/₂, time to decay to one-half of the peak luminance) for complexes containing ′Bu₂xanthphos was found to be low, approximately 5 minutes in the cases of bpy and 6,6′-Me₂bpy and above 50 minutes for the LECs with [Cu(′Bu₂xanthphos)(6-Mebpy)][PF₆]. The low lifetime of the complexes with bpy and 6,6′-Me₂bpy might be due to a reduced stability of the materials toward charge transport, as seen from the corresponding LECs voltage profile which drastically increases after only few minutes of operation (Fig. 14a).

The optoelectronic performance of complexes containing the xanthphosMes₂ ligand were in general lower as compared to those involving ′Bu₂xanthphos. We could not observe any electroluminescence from [Cu(xanthphosMes₂)(bpy)][PF₆], perhaps due to its low PLQY both in solution and in the solid state. Moderate electroluminescence was measured for [Cu(xanthphosMes₂)(6-MeMebpy)][PF₆], with fast turn-on (<1 minute) and a maximum lumiance of 50 cd m⁻².

The spectral shape and position of the electroluminescence (EL, Fig. 14c) signals correlate with the PL maxima observed for the complexes in solution and in the solid state. For the ′Bu₂xanthphos-containing complexes, the EL maxima blue-shift from 584 nm to 575 and 557 nm when increasing the substitution at the bpy, i.e. going from bpy to 6-Mebpy and 6,6′-Me₂bpy, respectively. As highlighted in the inset of Fig. 14c, this shift corresponds to a colour variation from the orange to the green region of the CIE 1931 colour space. The EL spectrum of [Cu(xanthphosMes₂)(6-Mebpy)][PF₆] peaks at 582 nm, in agreement with the PL signal of the thin-film (Table 3).

**Conclusions**

We have prepared [Cu(′Bu₂xanthphos)(bpy)][PF₆], [Cu(′Bu₂xanthphos)(6-Mebpy)][PF₆], [Cu(′Bu₂xanthphos)(6,6′-Me₂bpy)][PF₆], [Cu(xanthphosMes₂)(bpy)][PF₆] and [Cu(xanthphosMes₂)(6-Mebpy)][PF₆], but steric effects militate against the isolation of [Cu(xanthphosMes₂)(6,6′-Me₂bpy)][PF₆]. To prepare the latter, the chiral xanthphosMes₂ was prepared and characterized, and the single crystal structure reveals the presence of the (rac)-form. In solution, one dominant diastereoisomer is observed, proposed as the (rac) rather than the (meso)-form. This makes the bisphosphane preorganized to give particular diastereoisomers when coordinated to copper(I) in [Cu(xanthphosMes₂)(N′N′)] complexes. Single crystal structures of four complexes were determined. In [Cu(xanthphosMes₂)(6-Mebpy)][PF₆], the 6-Mebpy unit is disordered over two orientations with 50% occupancies. The disorder corresponds to a combination of two dynamic processes, which the [Cu(xanthphosMes₂)(N′N′)] cations undergo in solution. DFT calculations reveal that the energy difference between the two conformers observed in the solid-state structure differ only by 0.28 kcal mol⁻¹.

The [Cu(P⁺P)(N′N′)][PF₆] compounds show a broad MLCT-absorption around 380 nm which shifts to higher energies on going from bpy to 6-Mebpy to 6,6′-Me₂bpy. Upon excitation into the MLCT band, the [Cu(P⁺P)(N′N′)][PF₆] complexes emit in the yellow to orange region; additional Me groups in the bpy ligand result in a blue-shift in the emission. The MLCT nature of the absorption and emission is supported by DFT calculations, which associate the lowest-energy S₁ and T₁ excited states to the HOMO → LUMO monoexcitation implying...
an electron transfer from the Cu(P^P) moiety to the bpy ligand. In solution, PLQY values are low, but in the solid state, PLQYs of 26 and 59% are observed for [Cu(xantphosMes$_2$)(6-Me bpy)][PF$_6$] and [Cu($^2$Bu$_2$xantphos)[6,6'-Me$_2$ bpy]][PF$_6$], respectively, compared to benchmark values of 34 and 37% for [Cu(xantphos)(6-Me bpy)][PF$_6$] and [Cu(xantphos)[6,6'-Me$_2$ bpy]][PF$_6$]. Increased excited state lifetimes at low temperature are consistent with the complexes being TADF emitters and this is supported by a calculated energy difference between S$_1$ and T$_1$ of 0.17–0.25 eV.

The compounds were tested in simple bilayer LECs. The optoelectronic performance of complexes containing the xantphosMes$_2$ ligand were generally lower than those with $^2$Bu$_2$xantphos, which led to bright and efficient devices. The current efficiency of the LECs follows the trend observed for the PLQY, increasing with increasing substitution at the bpy ligand. In particular, luminances as high as 370 cd m$^{-2}$ were obtained for the complex [Cu($^2$Bu$_2$xantphos)[6,6'- Me$_2$ bpy]][PF$_6$], which correspond to an efficiency of 3.7 cd A$^{-1}$. These encouraging results suggest that $^2$Bu$_2$xantphos is a promising ligand to develop novel and efficient copper emitters for LECs and OLEDs.

Conflicts of interest

There are no conflicts to declare.

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Notes and references
