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Introduction

Luminescent Cu(I) complexes have gained increasing interest for their application in organic light emitting devices (OLEDs),¹ as they are environmentally benign and economically viable compared with noble metal complexes of Ru(II), Pt(II), Os(II) and Ir(III).²⁻⁶ In addition, they possess interesting photophysical and photochemical properties associated with the closed-shell d¹⁰ electronic configuration, such as thermally activated delayed fluorescence (TADF), which is attributed to the moderate spin– orbit coupling ($\xi_{Cu} = 857 \text{ cm}^{-1}$) and the raised non-radiative deactivating ligand-field state.⁷ Since the highly luminescent mononuclear Cu(I) complex, [Cu(dmp)(POP)]⁺ (where dmp = 2,9-dimethyl-1,10-phenanthroline, POP = bis[2-(diphenylphosphino)phenyl]ether) was reported by McMillin's group,⁸ a large

Effect of the dangling aromatic ring on neutral luminescent bis(phosphine) Cu(ı)/Ag(ı) complexes with the asymmetric pyridyl-tetrazolate ligands*

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A series of neutral luminescent bis(phosphine) Cu(i) complexes of pyridyl-tetrazolate ligands (L^1-L^3) with the general formula $[Cu^1(L^n)(P^AP)]$ (1–6) were synthesized, which have been well characterized by IR, UV/vis, CV, ¹H NMR and ³¹P NMR. For comparison, an Ag(i) complex $[Ag^1(L^2)(PPh_3)_2]$ (7) was also synthesized. The crystal structures of 2 and 7 have been further determined by X-ray crystallography. All these Cu(i) compounds show bright luminescence in the solid state with photoluminescence quantum yields (PLQYs) in the range of 25.8% to 85.0%. More interestingly, the Cu(i) complexes bearing an additional dangling aromatic ring on the diimine ligands exhibit enhanced luminescent performance in various solutions and their PLQYs are significantly higher than those of related Cu(i) complexes without steric protection. Compared with 1, the Cu(i) complexes with an additional dangling tetrazole moiety show a significant solvatochromic effect, which is uncommon for luminescent Cu(i) complexes. Moreover, $[Cu^1(L^2)(PPh_3)_2]$ (2) was further designed as an OLED material, which showed a high external quantum efficiency of 7.7%.

number of luminescent Cu(1) complexes have been reported so far.⁹ Among these mononuclear Cu(I) complexes, the distorted coordinated tetrahedrally heteroleptic complexes $[Cu^{I}(N^{N})(P^{P})]^{+/0}$ (N^N = various diimine ligands and P^P = diphosphine ligands) have been intensively studied, mainly because their emission properties are readily tuned by the systematic modification of electronic and steric properties of the diimine and diphosphine ligands.¹⁰ Accordingly, some luminescent heteroleptic Cu(1) complexes were designed as LED materials with a high external quantum efficiency (EQE > 15%).11 Compared with the traditionally well-established diimine ligands, such as the 2,2'-bipyridine (bpy), 1,10-phenanthroline (phen) and their derivatives, Cu complexes of tetrazole (tz)containing ligands, e.g. 5-(2-pyridyl)tetrazolate (L¹), were relatively less studied. However, these ligands possess some obvious advantages: (1) tz has a stronger coordination ability to Cu(1) centers than the pyridyl ring, due to its soft base property; (2) the ligand can be readily protonated/deprotonated due to the mesomeric structure of tz rings; (3) neutral Cu(1) complexes are readily formed, which enhance the post-processability for their application as an OLED material; (4) non-radiative decay



Fig. 1 The structures of HL¹-HL³.

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Fig. 2 Structures of neutral Cu(i)/Ag(i) complexes 1--7 containing various tz ligands.

through C-H vibrational quenching can be reduced by replacing the pyridine (py) ring with a tz unit; (5) the asymmetric coordination environment will improve their stability in the excited states.12 A typical example is the well-studied neutral heteroleptic Cu(I) complex $[Cu^{I}(L^{1})(P^{P})]$ (L¹ = 5-(2-pyridyl)tetrazolate).13 As revealed by DFT study, the HOMO is mainly confined to Cu(I) and the LUMO is located primarily on the pyridine ring and to a less extent on the tz ring. The MLCT absorption band mainly involves the metal d orbital and π^* orbital of the pyridine ring. Thus, the M-N(py) bond is weakened on the excited state, while the M-N(tz) bond remains almost unchanged. Recent study on the TADF properties of $[Cu^{I}(L^{1})(POP)]$ in the solid state have found an interesting direct intersystem crossing process.14 These unique properties have aroused our interest for the in-depth study of luminescent Cu(1) complexes containing this class of ligands.

Although $[Cu^{I}(L^{1})(PPh_{3})_{2}]$ and its related complexes exhibit excellent luminescent properties in the solid state, their fluid luminescent properties are usually very poor, mainly due to the geometrical distortion from tetrahedron to square plane on the excited state. One of the traditional strategies to improve the luminescent properties of Cu(I) complexes is to employ the sterically demanding a,a'-disubstituted diimine ligands, which effectively suppress the deformation of excited states. However, both the a,a'-positions in these diimine ligands should be occupied by bulky functional groups to suppress the excited geometric distortion, while the existence of a bulky group on only one of the a,a'-positions is not enough.¹⁵ Nevertheless, the charge-neutrality of resultant Cu(1) complexes is not easy to maintain, if the bulky alkyl groups are added on the a,a'-positions of these pyridyl tetrazolate ligands. In order to maintain the charge neutrality of Cu(1) complexes and improve their emission properties simultaneously, an additional aromatic ring was introduced on the 5-position of 5-(2-pyridyl)tetrazolate (L¹) (Fig. 1). Accordingly, a series of Cu(1) complexes, $[Cu^{I}(L^{n})(P^{P})]$ (2-6, Fig. 2) containing tz ligands $(L^{1}-L^{3})$ have been prepared. It is noted that these new Cu(I) complexes exhibit excellent luminescence not only in the solid state, but also in solution, including polar protic solvents. These Cu(I) complexes have no steric protection on the a,a'-positions, and exhibit enhanced solution-state luminescence. These compounds represent a class of neutral mononuclear Cu(1) complexes with the interesting TADF behaviour. To provide more insight on their luminescence, analogous Ag complex

 $[Ag(L^2)(PPh_3)_2]$ (7) was also synthesized as a comparison. Moreover, 2 has been further used to fabricate light-emitting diodes (OLEDs).

Results and discussion

The ligands HL¹, HL² and HL³ were synthesized by similar procedures via [3+2] cycloaddition reactions of organonitriles and N_3^{-17} Treatment of $[Cu^{I}(MeCN)_4](ClO_4)$ with phosphine ligands and various asymmetric diimine ligands afforded the neutral Cu(1) complexes 1-6 with the general formula $[Cu^{I}(L^{n})(P^{P})]$ (n = 1-3), which were isolated as yellow microcrystalline solids with moderate yields. In the similar condition, treatment of HL² with AgNO₃ afforded the colourless mononuclear neutral complex $[Ag(L^2)(PPh_3)_2]$ (7). All these complexes have been characterized by IR, UV/vis, ¹H NMR and ³¹P NMR spectra, satisfactory elemental analysis. The IR spectra of these compounds show strong stretches at \sim 1618 and 1435 cm⁻¹, which are assigned to ν (C=N) and ν (N=N) stretching bands from tz moieties and comparable with most related tz compounds.94,16 Moreover, a v(N-H) stretch at \sim 3125 cm⁻¹ is found in 2–5. The absence of stretching band of the anion (ClO_4^{-}) proves that these compounds are neutral in the solid state. The ¹H NMR spectra show that there are three well distinguished sharp proton signals in the normal range, which are consistent with their diamagnetic d¹⁰ electronic configurations. These complexes have also been characterized by ³¹P{¹H} NMR spectroscopy and broadening of their ³¹P signals were observed, which is similar to those reported in related Cu(1) and Ag(1) complexes (Fig. S3-S7[†]).^{9d,10b,21}

Crystal structures of 2 and 7

The crystal structures of 2 and 7 have been determined by X-ray crystallography, and selected bond parameters are summarized in Table 1. As shown in Fig. 3, the two compounds are iso-structural, and the metal centres are coordinated by a bidentate diimine ligand and two PPh₃ ligands in a distorted tetrahedron. In 2, the Cu1–N2 bond length (2.049(3) Å) is shorter than that of Cu1–N1 (2.160(3) Å), indicating that the coordinated tz moiety is deprotonated and forms the stronger back-bonding interaction with Cu atom. In the structurally similar complex [Cu(N^N)(POP)]⁺ (N^N = 2-(2-tert-butyl-2H-tetrazol-5-yl)pyridine),¹⁶ the Cu–N_(py) and Cu–N_(tz) bond lengths are essentially identical, since both heterocycles are neutral. Compared with pyridine ring, the deprotonated tz moiety has stronger coordination



Fig. 3 The ORTEP drawing of structures for 2 (a) and 7 (b).

Table 1 Selected bond parameters (Å, $^{\circ}$) for 2 and 7

	7	
2.049(3)	Ag1-N6	2.308(6)
2.160(3)	Ag1-N7	2.513(6)
2.235(2)	Ag1-P1	2.419(2)
2.265(2)	Ag1-P2	2.471(2)
79.16(12)	N6-Ag1-N7	69.84(19)
122.48(4)	P1-Ag1-P2	128.27(7)
	2.160(3) 2.235(2) 2.265(2) 79.16(12)	2.160(3) Ag1-N7 2.235(2) Ag1-P1 2.265(2) Ag1-P2 79.16(12) N6-Ag1-N7

ability and higher electron density, resulting in an asymmetric coordination environment. Compared with $[Cu^{I}(N^{N})(PPh_{3})_{2}]^{\dagger}$ with symmetric diimine ligands, the Cu-N_(tz) bond length is also shorter. The shortening of Cu-N bond distance leads also to a slightly smaller P-Cu-P bite angle (122.48(4)°). Similar result is also found in 7, where the Ag-N6 and Ag-N7 bond distances are 2.308(6) and 2.513(6) Å, respectively and the P-Ag-P bond angle is $128.27(7)^{\circ}$. Although 2 and 7 are isostructural, the N-M-N bite angle of 2 (79.16(12)°) is significantly larger than the corresponding angle $(69.84(19)^\circ)$ in 7. The pyridyl ring is coplanar with the deprotonated tz ring with small dihedral angles of 1.8 and 2.8°, for 2 and 7 respectively. It is noted that the dangling tz ring and the coordinating pyridyl tetrazole ligands have dihedral angles of 9.0 and 7.9° respectively. Extensive weak interactions, such as π - π stacking interactions of aromatic rings and H-bondings via NH…N of adjacent Cu(I) complexes, have been found in their solid structures. (Fig. S9 and S10[†])

Electronic absorption and emission properties

The UV/vis spectra of these compounds in CH_2Cl_2 solution and the ligands $HL^{1}-HL^{3}$ in MeOH solution have been collected at room temperature. As shown in Fig. S1,† these ligands exhibit the strong ligand-centred π - π * absorption bands <330 nm. The overlaid absorption spectra of 1–7 are shown in Fig. 4. Apart from the intense ligand-centred (LC) π - π * transitions of phosphine and tz ligands in CH_2Cl_2 solutions with molar extinction coefficients on the order of $10^4 \text{ M}^{-1} \text{ cm}^{-1}$ in the range of 220–330 nm, 1–6 also show moderate absorption bands with molar extinction coefficients on the order of $10^3 \text{ M}^{-1} \text{ cm}^{-1}$ from 330 to 400 nm tailing down off 450 nm. Compared with related cationic complexes [Cu^I(N^N)(PPh_3)2]⁺, these absorption bands are obviously blue-shifted, in agreement with the raising



Fig. 4 The UV/vis spectra of 1-7 in CH₂Cl₂ solution.

of π^* -orbitals of diimine ligands. Since neither phosphine ligands nor tz ligands show absorption bands in these ranges, these absorption bands are mainly assigned to metal-to-ligand charge transfer (MLCT) of $\{d(Cu) \rightarrow \pi^*(tz)\}$, possibly mixed with somewhat contributions of inter-ligand charge transfer. As suggested by TD-DFT studies, the lowest-lying transitions of 1-6 are assigned as the ¹MLCT/¹LLCT charge-transfer (Cu/PPh₃-N^N) transitions, mixed with some intra-ligand (¹IL) transitions of the anionic chelated ligands. These assignments are further supported by the dependence of the absorption energy on the π accepting ability of phosphine ligands, with the order of $P(PhMe)_3 < POP < PPh_3 < P(C_6H_4Cl)_3$ for compounds 2-5 containing the same diimine ligand L^2 . The increase of the π accepting ability of phosphine ligands leads to the blue-shifted of the lowest-energy absorption bands. The assignment of the lowest energy absorption bands of these Cu(I) complexes are also supported by the dependence of the absorption energy on the electronic nature of diimine ligands, which is in line with the order of expected π^* orbital energy level $[L^2 < L^1 < L^3]$ for the complexes 1, 2 and 6 bearing the same PPh_3 ligand. For Ag(1)complex 7, the corresponding MLCT absorption band is not observed due to the lower-lying d orbital of the Ag⁺ ion.

The emission properties of these complexes have been investigated in CH₂Cl₂ solution at room temperature (Table 2, Fig. 5b). All these Cu(i) complexes exhibited yellow to orange emissions (539–595 nm) with photoluminescence quantum yields (PLQYs) up to 4.7% upon excitation with $\lambda_{ex} = 380$ nm. Although the PLQYs of these compounds in solution are lower than that of [Cu^I(dmp)(PPh₃)₂]⁺ (dmp = 2,9-dimethylphenan-throline),¹⁸ they are significantly higher than most of the Cu(i) complexes [Cu(N^N)(PPh₃)₂]⁺ without the sterically-protected

Table 2Photophysical data for 1–7

		Emission ^{<i>a</i>} λ /nm (τ_0 /	
	Medium	μs)	$\phi_{ m em} \left(imes 10^3 ight)$
1	CH_2Cl_2 (298)	571 (0.48)	2
	Solid $(298)^b$	512 (20.6)	850
	Glass (77)	534 (230.47)	
2	CH_2Cl_2 (298)	600 (0.52)	31
	Solid (298)	567 (1.65)	258
	Glass (77)	578 (112.10)	
3	CH_2Cl_2 (298)	595 (0.65)	12
	Solid (298)	544 (8.23)	393
	Glass (77)	569 (75.43)	
4	CH_2Cl_2 (298)	590 (0.72)	47
	Solid (298)	548 (2.31)	403
	Glass (77)	543 (15.42)	
5	CH_2Cl_2 (298)	589 (0.07)	0.3
	Solid (298)	546 (8.72)	628
	Glass (77)	568 (102.50)	
6	CH_2Cl_2 (298)	539 (2.02)	5.2
	Solid (298)	524 (6.69)	486
	Glass (77)	528 (263.39)	
7	CH_2Cl_2 (298)	438 (<10 ns)	1
	Solid (298)	528 (15.78)	66
	Glass (77)	517 (208.28)	

 a Excitation at 380 nm in $\rm CH_2Cl_2$ solution and 355 nm for glassy state at 77 K. b Ref. 13.



Fig. 5 (a) Emission spectra Cu(I) complexes 1-6 in the solid state at room temperature (b) in CH₂Cl₂ at room temperature and (c) in low-temperature 77 K glassy media EtOH–MeOH (4 : 1, v/v) ($\lambda_{ex} = 355$ nm).

diimine ligands.^{18,19} All these emission spectra are broad and structureless (with ~110 nm of the full width at half-maximum), consistent with the MLCT assignment with the large structural distortion on the excited state. The emission maxima of 2–5 with the same L² ligand are in the range of 589 to 600 nm and red-shifted as compared with **1**. In contrast to the observed trend in the lowest energy absorption, the emission energies of 2–5 varied with the substituents on phosphine ligands, which is in the trend of **2** (600 nm, PPh₃) > **3** (595 nm, P(PhMe)₃) > **4** (590 nm, P(PhCl)₃) > **5** (589 nm, POP), as reflected from the overlaid emission spectra of **1–6** (Fig. 5b). Similar observation is also recently reported in our luminescent Cu(i) complexes containing various phosphine ligands.^{10b}

The luminescence of 2-4 in CH₂Cl₂ is significantly enhanced as compared with 1. It is noted that although 1 exhibits the excellent PLQY of \sim 85% in the solid state, its PLQY is only 0.2% in degassed CH2Cl2 solution. On the introduction of an additional aromatic ring into the pyridyl ring, the PLQYs in CH₂Cl₂ solution increase from 1.2% to 4.7%. It is noted that most Cu(1) complexes are nearly non-emissive or poorly emissive without the bulky substituents on the $a_{a}a'$ -positions of dimine ligands.¹⁸ The emission properties of these luminescent Cu(I) complexes reported herein are attributed to the dangling aromatic rings on the pyridyl ring, which enhance the π -conjugation system. Moreover, 5 with a POP ligand is nearly non-emissive with PLQY \sim 0.03% and its lifetime ($\tau = 0.07 \ \mu s$) is also much shorter than the other Cu(1) complexes in the range of 0.48 to 2.02 µs. Considering the spectroscopic studies on the related Cu(1)complexes,19 as well as the microsecond emission lifetime and tunable emission, these emissions are ascribed to ³MLCT $\{d(Cu) \rightarrow \pi^*[N^N]\}$ probably admixed with somewhat ³LLCT $\{\pi(\text{phosphine}) \rightarrow \pi^*[N^N] \text{ excited states. All the emission}\}$ maxima of Cu(1) complexes in solutions are relatively red-shifted as compared with those in solid state, which result from to deformation of excited states due to lack of steric protection. On the contrary, the Ag(I) complex 7 shows a blue emission at $\lambda_{em} =$

438 nm with lifetime less than 10 ns in CH₂Cl₂ solution (Fig. S11a[†]). Thus, it is tentatively assigned to metal-perturbed ligand-centred π - π * transitions, which are similarly observed in structurally related mononuclear Ag(1) complexes.²⁰

In the solid state, these complexes show intense broad green (512 to 567 nm) unstructured photoluminescence, typically observed in luminescent Cu(1) complexes (Fig. 5a and Table 2). The lifetimes are in the range of 1.65 to 20.6 µs. Moreover, the PLQYs in solid state for 2-6 ranging between 25.8% and 62.8%, which are lower than the reported PLQY of 1 (85%), possibly due to the non-radiative decay through C-H/N-H vibrational quenching of the dangling aromatic rings. The enhanced solid emissions of these Cu(I) complexes are attributed to the decrease in the excited state distortion of Cu(I) complexes and non-radiative decay in the solid state. Since the rigidochromic effect impedes the structural distortion in the excited state, the solid-state emissions for most of these Cu(I) complexes are blueshifted as compared with their emissions in solution. In contrast to the solution emissions, the solid-state emissions show a different trend on the electronic properties of phosphine ligands and diimine ligands. The deviation can be explained by the presence of various factors such as inter- and innermolecular π -stacking in the solid state, as previously reported in the solid-state emissions of related Cu(1) complexes.^{9d,10b} In contrast, the emission maxima of complexes 2-5 with the same L² ligand range from 544 to 567 nm and are red-shifted compared with that of **1**. Upon excitation at $\lambda_{ex} = 380$ nm, **7** exhibits a green emission peaking at 528 nm. Based on the long decay times, large Stokes shifts, and spectroscopic studies of the related Ag(1) complexes,20 the emission of 7 in the solid state is assigned as intra-ligand ³LC phosphorescence. (Fig. S11b[†])

In 77 K EtOH–MeOH (4:1, v/v) glassy medium, all these complexes display intense luminescence (Fig. 5c and Table 2). Compared with their emissions in solution, the emissions in the glassy state are generally blue-shifted and longer-lived. This is attributed to the rigidochromic effect, which reduces the



Fig. 6 Frontier molecular orbitals for the triplet state of complexes 2 and 7 calculated at the B3LYP/6-31+G(d)/LANL2DZ level of theory.

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structural distortion on the excited state and is also commonly observed in MLCT emitters. However, compared with the emission spectra in the solid state at room temperature, the emission spectra at 77 K are red-shifted, which are normally observed for $[Cu^{I}(N^{N})(P^{P})]^{+}$ complexes due to the presence of TADF behaviour.8 The longer excited state lifetimes in the glassy medium at 77 K further support the assignment of triplet excited state. These emissions are also tentatively assigned to ³MLCT [d(Cu) $\rightarrow \pi^*(N^N)$] excited state. The emission band of 7 in low-temperature glassy medium is highly structured with a vibrational progression spacing of $\sim 1200 \text{ cm}^{-1}$. The significantly different low-temperature emission properties of 7 is suggestive of a different emission origin, which is ascribed to the ³LC excited state of the L² ligand. (Fig. S11c[†]) The emission maximum of 7 is very close to those of the Cu(I) complexes of L^2 in glassy state. Thus, there may be contributions from ³LC excited state to the photoluminescence of Cu(I) complexes at low temperature.

Electronic structure calculations

Density functional theory (DFT) calculations of the complexes were performed by using the Gaussian 09 program to determine the nature of frontier orbitals. The lowest energy triplet excited states of 2 and 7 were computed and the two singly occupied molecular orbitals (SOMOs) were identified. Contour plots of the two SOMO and the spin densities of the complexes were shown in Fig. 6. Hydrogen atoms in the structures were omitted for clarity. For the Cu(1) complex 2, the lower energy SOMO of the complex consist of $d_{\pi}(Cu)$ with contributions by lone pairs of P (from PPh_3) and N (from L^2), while the higher one consist of the π^* orbital of L² ligand, which suggests the triplet excited state of the complex is MLCT $[Cu \rightarrow L^2]$ in nature, with some mixing of LLCT [PPh₃ \rightarrow L²]. For the Ag(1) complex 7, the two SOMOs consist of π and π^* orbital of the L² ligand, which suggests the triplet excited state of the complex is ligandcentred (IL) in nature.

Solvatochromic effect

Many Cu(1) complexes exhibit low stability in various solutions, especially in their excited states, which limit their applications as probes/sensors for O2, pH, and various ions. Herein, the asymmetric ligands were used instead of bipyridine and phenanthroline diimine ligands. These asymmetric diimine ligands may provide obvious advantages over the symmetric one. In the ground state, the electron-rich tz moiety acts as a soft-base, which could form stronger bonds with Cu(I) ions. In the excited state, the MLCT transition could be attributed to the Cu(I) centre and pyridyl group of the ligands, while the tz moiety is not involved significantly due to its high lying π^* orbital and effectively stabilize the excited state of these Cu(I) complexes. Moreover, the L² in complexes 2-5 have an additional protonated tz moiety, whose pK_a value (~4.5) is comparable with the carboxylic acid. Thus, their emissions may be sensitive towards the pH variation.

As expected, the MLCT absorption bands of these Cu(I) complexes showed a dependence on the nature of the solvents.



Fig. 7 The absorption spectra (a) and emission spectra (b) of ${\bf 3}$ in various solvents.

Taking 3 as an example (Fig. 7), the lowest-energy MLCT absorption bands in the range 350-400 nm are slightly dependent on the solvents used, while the LC absorption bands remain almost unchanged in different solvents. In contrast, the emission spectra of 3 show a stronger solvatochromic effect in various solvents. The emission maxima (λ_{em}) follow the order of DMF (553 nm) < MeOH (560 nm) < CH₃CN (564 nm) < CHCl₃ $(584 \text{ nm}) < CH_2Cl_2$ (594 nm). Such a variation trend is mainly attributed to the existence of an acidic N-H bond in the dangling tz moiety, which is readily affected by solvent polarity and H-bonding interaction. In MeOH, the acidic N-H group forms strong H-bonding with solvents, which raises the π^* orbital of ligand. Thus a higher-energy emission could be expected, which is supported by the emission observed in basic DMF with higher electron-density. This hypothesis is also supported by the negligible solvatochromic effect for 1 which does not bear a dangling tz unit (Fig. S2[†]).

Electroluminescence performance

As a result of the excellent solid-state luminescent properties of these neutral Cu(i) complexes, OLED containing 2 was fabricated to evaluate their EL properties. A structure of ITO/HAT-



Fig. 8 (a) The OLED device structure of 2-based device; (b) the EL spectra of 20 wt% doping device under different voltages of 2; (c) current density–voltage–brightness (I-V-B) curves of 2-based device; (d) the dependence of the current efficiency of 2 on the current density.

CN/TAPC/TCTA/mCP/mCP: PO-T2T: 20 wt% of 2/PO-T2T/LiF/Al was designed. Among them, HAT-CN, TAPC, TCTA, mCP, PO-T2T, and LiF are used as hole injection layer, hole transport layer, electron blocking layer, exciton blocking layer, electron transport layer, and electron injection layer, respectively. The corresponding energy level diagrams and molecular structures of the charge-transport materials are shown in Fig. 9. ITO conductive glass was used as the anode. The ITO conductive glass was treated with UV ozone and then placed in an organic electroluminescence integrated deposition system. Each organic material was formed on the ITO conductive glass by solution spin coating method on a 2500 r/s spin coater in turn. The film thickness was monitored by a quartz oscillating plate, and the light-emitting area was 3 mm \times 3 mm. The HOMO and LUMO levels of these Cu(1) complexes were determined using the cyclic voltammetry (CV) and the optical bandgaps are estimated (Fig. S12 and Table S1[†]). As shown in Fig. 8b, the device containing 2 emits bright green emission peaking at 526 nm, and the emission spectrum of the electroluminescent devices is comparable with the solid-state emission spectrum, indicating that EL emission comes directly from 2. However, the emission maxima in these spectra exhibit a slight blue shift at different voltages, possibly due to the difficulty in managing excitons in a single emission layer. Since the primary mCP and charge transfer material used are non-emissive, an effective combination of holes and electrons occurred in the dopant. The Commission Internationale de L'Eclairage (CIE) coordinates of this device is (0.365, 0.551) (Fig. 9d). The current densityvoltage-brightness curves are displayed in Fig. 8c, revealing a turn-on voltage at 2.4 V and a luminance of 1000 cd m⁻² for lighting at 3.5 V. For most Cu(1) devices, a 2.4 V turn-on voltage is very low. The low resistance at doping concentration of 20 wt% suggests that 2 is an efficient charge carrier, as high doping concentration usually leads to the concentration quenching effect. The maximum current efficiency was 22.96 cd A^{-1} at a current density of 0.01 mA cm⁻² (Fig. 8d). At the



Fig. 9 (a) EQE–voltage characteristics of **2**-based device; (b) the molecular structures of the charge transfer compound in the device; (c) the energy band diagram for **2**; (d) CIE diagram of **2**-based device.

maximum current efficiencies, the maximum external quantum efficiency (EQE) was 7.7% (Fig. 9a), comparable to the EQEs of most Cu(i)-doped OLEDs. The good performance of this device could be attributed to the TADF-type launch.

Conclusion

Functionalized pyridyl-tetrazole ligands have been studied for the design of neutral heteroleptic Cu(I)-based compounds with the general formula $[Cu(N^N)(P^P)]$ (1–6). All these Cu(I) complexes exhibit strong luminescence in the solid state. Upon the introduction of a dangling aromatic moisty on the pyridyl ring, the luminescent performance in solutions is significantly enhanced as compared to parent pyridyl-tetrazole complex 1. OLED containing 2 exhibits high EQE and current efficiency. These features suggest that heteroleptic Cu(I) complexes with pyridyl-tetrazole ligands could be used as an economic luminescent material.

Experimental section

Materials and methods

Triphenylphosphine (PPh₃), tri(*p*-tolyl)phosphine [P(PhMe)₃], tri(4-chlorophenyl)phosphine [P(PhCl)₃], bis{2-(diphenylphosphanyl)phenyl} ether (POP), were purchased from Acros Organic Chemical Company. 2-(1H-tetrazol-5-yl)pyridine (HL¹), 2,4-di(1H-tetrazol-5-yl)pyridine (HL³) were synthesized by the similar procedures *via* [3 + 2] cycloaddition reactions of organonitriles and N₃⁻.¹⁷ [Cu(MeCN)₄](ClO₄)²¹ and [Cu^I(L¹)(PPh₃)₂] (1)¹³ were synthesized according to literature procedures. The commercially available reagents and solvents of analytical grade were used without further purification.

Synthesis and characterization

 $[Cu^{I}(L^{2})(PPh_{3})_{2}]$ (2). $[Cu(MeCN)_{4}]ClO_{4}$ (50 mg, 0.153 mmol), PPh₃ (80.3 mg, 0.31 mmol) were added to CH₃CN (15 mL) and the mixture was stirred at room temperature for 1 h under nitrogen atmosphere. Then, HL² (32.9 mg, 0.15 mmol) was added and was further stirred for 1 h. After removal of solvent under reduced pressure, the residue was purified by column chromatography on silica gel eluted with acetone/ dichloromethane (v:v, 1:4). The bright yellow layer was collected. The light yellow microcrystalline solid was obtained and was further purified by slow evaporation of a mixed CH₂Cl₂/ MeOH solvent of 2. Yield: 105 mg, 64.0%. Elemental analysis for C₄₃H₃₄CuN₉P₂: calcd C 64.37, H 4.27, N 15.71%; found C 64.33, H 4.15, N 15.74%, selected IR (KBr, cm⁻¹): v(N–H) 3139, v(N=N) 1435, ν (C=N) 1619. UV/vis (CH₂Cl₂): λ_{max}/nm (ϵ/mol^{-1} dm³ cm⁻¹):228 (71 110), 256sh (48 870), 366sh (7000). ¹H NMR $(400 \text{ MHz}, \text{CDCl}_3)$: 9.41 (s, 1H, L²-H), 9.12 (d, $J = 5.4 \text{ Hz}, 1H, \text{ L}^2$ -H), 8.18 (dd, J = 5.4, 1.7 Hz, 1H, L²-H), 7.38–7.30 (m, 6H, PPh₃-H), 7.23–7.08 (m, 24H, PPh₃-H); ³¹P{¹H} NMR (162 MHz, CDCl₃): δ: 0.68 (s, PPh₃).

 $\{Cu^{I}(L^{2})[P(PhMe)_{3})_{2}\}\$ (3). The synthetic procedure of 3 is similar to that of 2 except that $P(PhMe)_{3}$ (93.07 mg, 0.306 mmol)

was used instead of PPh₃. Yield: 104 mg, 52.8%. Elemental analysis for C₄₉H₄₆CuN₉P₂: calcd C 66.39, H 5.23, N 14.22%; found C 66.54, H 5.28, N 14.43%. Selected IR (KBr, cm⁻¹): ν (N–H) 3128, ν (N=N) 1436, ν (C=N) 1621. UV/vis (CH₂Cl₂): λ _{max}/nm (ε /mol⁻¹ dm³ cm⁻¹): 230 (102 300), 263sh (42 980), 371sh (2900). ¹H NMR (400 MHz, CDCl₃): δ 7.97 (d, *J* = 8.3, 1H, L²-H), 7.76 (d, *J* = 5.4 Hz, 1H, L²-H), 7.56 (dd, *J* = 11.8, 8.1 Hz, 10H), 7.32 (s, 1H), 7.26 (d, *J* = 2.6 Hz, 6H), 7.12 (d, *J* = 16.5 Hz, 8H), 1.62 (s, 18H, -CH₃). ³¹P{¹H} NMR (162 MHz, CDCl₃): δ 29.41 (s, P(PhMe)₃).

{Cu^I(L²)[P(PhCl)₃)₂]} (4). The synthetic procedure of 4 is similar to that of 2 except that P(PhCl)₃ (112.0 mg, 0.307 mmol) was used instead of PPh₃. Yield: 124 mg, 55.3%. Elemental analysis for C₄₃H₂₈Cl₆CuN₉P₂: calcd C 51.19, H 2.80, N 12.49%; found C 52.23, H 2.68, N 12.54%. Selected IR (KBr, cm⁻¹): ν (N-H) 3130, ν (N=N) 1435, ν (C=N) 1620. UV/vis (CH₂Cl₂): λ _{max}/nm (ε /mol⁻¹ dm³ cm⁻¹):232 (88 390), 264sh (43 000), 357sh (5600). ¹H NMR (400 MHz, CDCl₃): 10.66 (s, 1H, L²-H), 8.25 (d, *J* = 8.0 Hz, 1H, L²-H), 7.84 (d, *J* = 8.0 Hz, 1H, L²-H), 7.72–7.29 (m, 4H, PPh₃-H), 7.13–7.01 (m, 10H, PPh₃-H), 6.93–6.80 (m, 10H, PPh₃-H). ³¹P{¹H} NMR (162 MHz, CDCl₃): δ –2.01 (s, P(PhCl)₃).

[**Cu^I**(**L**²)(**POP**)] (5). The synthetic procedure of 5 is similar to that of 2 except that POP (82.33 mg, 0.153 mmol) was used instead of PPh₃. Yield: 87 mg, 50.9%. Elemental analysis for C₄₃H₃₂CuN₉OP₂: calcd C 63.27, H 3.95, N 15.44%; found C 63.56, H 3.76, N 15.54%. Selected IR (KBr, cm⁻¹): ν(N−H) 3132, ν(N=N) 1435, ν(C=N) 1622. UV/vis (CH₂Cl₂): λ_{max}/nm (ε/mol⁻¹ dm³ cm⁻¹): 228 (76 630), 254sh (49 280), 290sh (27 892), 380 (6503). ¹H NMR (400 MHz, CDCl₃): 9.83 (s, 1H, L²-H), 8.10 (d, *J* = 5.4 Hz, 1H, L²-H), 8.05 (d, *J* = 5.4 Hz, 1H, L²-H), 7.40–7.66 (m, 4H, POP-H), 7.35–7.25 (m, 4H, POP-H), 7.22–7.11 (m, 8H, POP-H), 7.06–6.98 (m, 4H, POP-H), 6.98–6.90 (m, 4H, POP-H), 6.80–6.66 (m, 4H, POP-H). ³¹P{¹H} NMR (162 MHz, CDCl₃): δ −12.98 (s, POP).

[**Cu^I**(**L**³)(**PPh**₃)₂] (6). The synthetic procedure of 6 is similar to that of 1 except that HL³ (34.13 mg, 0.153 mmol) was used instead of HL¹. Yield: 65 mg, 52.4%. Elemental analysis for C₄₈H₃₈CuN₅P₂: calcd C 71.14, H 4.73, N 8.64%; found C 71.62, H 4.62, N 8.48%. Selected IR (KBr, cm⁻¹): ν (N=N) 1434, ν (C=N) 1613. UV/vis (CH₂Cl₂): λ _{max}/nm (ε /mol⁻¹ dm³ cm⁻¹): 230 (98 420), 260sh (62 450), 353sh (3000). ¹H NMR (400 MHz, CDCl₃): 8.52 (s, 1H, L³-H) 7.91 (d, *J* = 5.5 Hz, 1H, L³-H), 7.71 (dd, *J* = 8.0, 1.5 Hz, 2H, L³-H), 7.54–7.45 (m, 4H, L³-H), 7.32–7.26 (m, 6H, PPh₃), 7.15–7.13 (m, 24H, PPh₃-H); ³¹P{¹H} NMR (162 MHz, CDCl₃): δ 10.25 (s, PPh₃).

[Ag^I(L²)(PPh₃)₂] (7). AgBF₄ (50 mg, 0.25 mmol), PPh₃ (134.0 mg, 0.5 mmol) were added to CH₃CN (15 mL) and was stirred at room temperature for 1 h under nitrogen atmosphere. After that, HL² (54 mg, 0.25 mmol) was added, and the mixture was further stirred for 1 h. After removal of the solvent under reduced pressure, the residue was further purified by slow diffusion of diethyl ether into a CH₂Cl₂ solution of 7 to give colorless crystals. Yield: 96 mg, 59.4%. Elemental analysis for C₄₃H₃₄AgN₉P₂: calcd C 61.00, H 4.05, N 14.89%; found C 61.43, H 4.32, N 14.98%. Selected IR (KBr, cm⁻¹): ν (N–H) 3135, ν (N=N) 1436, ν (C=N) 1618. UV/vis (CH₂Cl₂): λ_{max} /nm (ε /mol⁻¹ dm³ cm⁻¹): 230 (52 327), 252sh (38 046). ¹H NMR (400 MHz,

 $\begin{array}{l} {\rm CDCl}_3{\rm):} \ 9.63 \ ({\rm s}, 1{\rm H}, {\rm L}^2{\rm -H}), \ 8.11 \ ({\rm d}, J=5.1 \ {\rm Hz}, 1{\rm H}, {\rm L}^2{\rm -H}), \ 8.03 \ ({\rm d}, J=5.4 \ {\rm Hz}, 1{\rm H}, {\rm L}^2{\rm -H}), \ 7.42{\rm -4.34} \ ({\rm m}, \ 6{\rm H}, \ {\rm PPh}_3{\rm -H}), \ 7.34{\rm -7.29} \ ({\rm m}, \ 8{\rm H}, \ {\rm PPh}_3{\rm -H}), \ 7.28{\rm -7.22} \ ({\rm m}, \ 16{\rm H}, \ {\rm PPh}_3{\rm -H}); \ {}^{31}{\rm P}\{{}^{1}{\rm H}\} \ {\rm NMR} \ (162 \ {\rm MHz}, \ {\rm CDCl}_3){\rm :} \ \delta \ 9.22 \ ({\rm s}, \ {\rm PPh}_3). \end{array}$

Conflicts of interest

There are no conflicts to declare.

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