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## Peripheral engineering of platinum(II) dicarbene pincer complexes for efficient blue hyperphosphorescent organic light-emitting diodes<sup>†</sup>

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Pt(II) dicarbene pincer complexes have been explored as deep-blue phosphors with narrowband emissions. Generally, they form excimers easily and their relatively long emission lifetimes also limit their applications as OLED emitters. Herein, we developed a simple method to modify 1,3-bisimidazol-3-ium pro-chelates and obtained three Pt(II) dicarbene pincer complexes, thus studying the issue of excimer formation vs. peripheral appendages. As a result, blue emissions can be maintained at doping concentrations of up to 20 wt% for IPrtBuPt. Moreover, the emission of IPrtBuPt was found to significantly overlap with the absorption of  $\nu$ -DABNA, an efficient blue boron/nitrogen based multi-resonant TADF molecule. A hyper-OLED using IPrtBuPt as a sensitizer and  $\nu$ -DABNA as a terminal emitter achieved EQEs of up to 33.59%, FWHM of 20 nm and CIE coordinates of (0.124, 0.148). These results demonstrated a successful example of utilizing both structural engineering and smart device architecture for getting efficient blue devices with suppressed efficient roll-off.

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### Introduction

While there has been significant progress in the utilization of both red and green phosphors for fabrication of organic light-emitting diodes (OLEDs),<sup>1–5</sup> the development of blue phosphors is significantly delayed. Therefore, blue fluorophores, which can only utilize one quarter of electrogenerated excitons, are still being used in modern commercial OLED products.<sup>6</sup> Thus, novel blue emitters and device architectures are urgently needed for boosting the performance of blue OLED devices.<sup>7,8</sup>

Multi-resonant thermally activated delayed fluorescent (MR-TADF) molecules have become the most intensively researched topics due to their merits including fast radiative decay,<sup>9–11</sup> narrow full width at half maximum (FWHM), proneness to

adapt horizontal alignment and high luminescent efficiency.<sup>12,13</sup> Thus, MR-TADF molecules possess great potential to improve external quantum efficiency (EQE) towards 40% with color purity comparable to quantum dot LEDs,<sup>14–17</sup> simultaneously. Another type of promising emitter is an N-heterocyclic carbene ( $C^NHC$ ) coordinated iridium(III) complex ( $Ir(C^NHC)_3$ ),<sup>18</sup> a representative emitter of which, *i.e.*, *mer*-Ir(CF<sub>3</sub>ppb)<sub>3</sub>, has delivered an EQE<sub>max</sub> of 21.2% with Commission Internationale de L'Eclairage (CIE) coordinates of (0.16, 0.05),<sup>19</sup> showing performances comparable to the state-of-the-art TADF emitters. This result is ascribed to the strong  $\sigma$ -donating  $C^NHC$  chelate, which greatly destabilizes the metal-centered (MC) dd excited state and enlarged energy offset between the emitting excited states and upper lying MC dd quenching states.<sup>7,8</sup>

Compared to TADF emitters and Ir(III) complexes, the development of deep-blue Pt(II) complexes is relatively lagging behind. In fact, the large spin-orbital coupling (SOC) constant of the Pt atom (4881 vs. 3450 cm<sup>−1</sup> for Ir)<sup>20</sup> confirmed its potential for achieving efficient phosphorescence. Incorporating an N-heterocyclic carbene entity, Li *et al.* demonstrated a tetradentate  $N^C^NHC$  coordinated Pt(II) complex with an EQE<sub>max</sub> of 24.8% and CIE coordinates of (0.148, 0.079).<sup>21,22</sup> However, its square-planar geometry and long radiative lifetime resulted in a low luminance and increased efficiency roll-off at high luminance (*e.g.*, 1000 cd m<sup>−2</sup>). Modifications to alleviate this problem are impeded by the sophisticated synthetic procedure of

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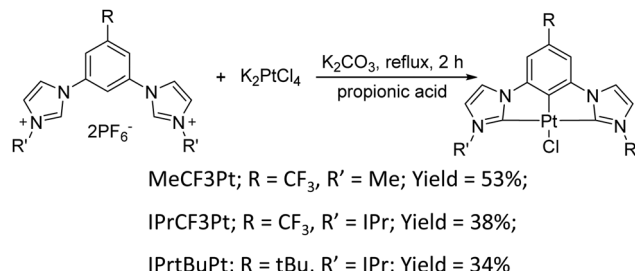
tetradentate chelates.<sup>23,24</sup> In contrast, the dicarbene pincer chelate can be modified much more easily. A pioneering research study shows a decent blue emission with a narrow FWHM down to  $\sim 20$  nm and a blue device with an  $\text{EQE}_{\text{max}}$  of 14.8% and CIE coordinates of (0.16, 0.13) using Pt(II) based  $\text{C}_{\text{NHC}}^{\wedge}\text{C}^{\wedge}\text{C}_{\text{NHC}}$  pincer complexes as dopants,<sup>25,26</sup> although they gave dimers (or excimers) at higher doping ratios. On the other hand, lowered doping ratios may lead to unbalanced transport of carriers, which undermine device performance at high brightness (e.g., 100 and 1000  $\text{cd m}^{-2}$ ). In fact, the emission of our Pt(II) pincer complexes exerts good spectral overlaps with absorption bands of the blue terminal emitter, *i.e.*,  $\nu$ -DABNA ( $N^7,N^7,N^{13},N^{13}$ ,5,9,11,15-octaphenyl-5,9,11,15-tetrahydro-5,9,11,15-tetraaza-19*b*,20*b*-diboradiphtho [3,2,1-*de*:1',2',3'-*jk*]pentacene-7,13-diamine).<sup>15</sup> This encouraged us to employ the Förster resonance energy transfer (FRET) process in realizing efficient blue OLEDs.<sup>27-29</sup>

Herein, we modify the peripheral groups of the  $\text{C}_{\text{NHC}}^{\wedge}\text{C}^{\wedge}\text{C}_{\text{NHC}}$  chelate to minimize intermolecular interactions and to develop three Pt(II) ( $\text{C}_{\text{NHC}}^{\wedge}\text{C}^{\wedge}\text{C}_{\text{NHC}}$ ) pincer complexes, namely MeCF3Pt, IPrCF3Pt and IPrtBuPt. These modifications have successfully addressed the issue on excimer formation and given pure blue emission with  $\text{EQE}_{\text{max}} = 13.96\%$  and CIE coordinates of (0.159, 0.241) from an IPrtBuPt-based OLED device. More importantly, the emission of IPrtBuPt shows a significant overlap with the absorption of MR-TADF blue emitter –  $\nu$ -DABNA. A hyperphosphorescent device with IPrtBuPt as a dopant sensitizer and  $\nu$ -DABNA as a terminal emitter is demonstrated to show a maximum EQE of 33.59% and color coordinates of (0.124, 0.148). It also revealed very mild efficiency roll-offs comparable to other hyper-OLED devices in the literature,<sup>7,8,15,23,24,30-38</sup> together with a high max. brightness of  $\sim 30\,000 \text{ cd m}^{-2}$ . Eventually, this study shows that a combination of simple structural modifications of Pt(II) dicarbene pincer complexes and smart device engineering can be promising in the generation of high performance blue OLEDs.

## Results and discussion

Synthesis of the dicarbene pincer pro-chelates follows the same protocol reported in our previous works.<sup>39,40</sup> The Pt(II) complexes are obtained by refluxing  $\text{K}_2\text{PtCl}_4$  and hexafluorophosphate salts of their corresponding 1,3-bisimida-zol-3-ium pro-chelates (Scheme 1 and the Experimental section) in the presence of potassium carbonate in propionic acid (yield: 34–54%, Scheme 1). It is noted that this method is simpler and more straightforward than the other literature precedents, which needs expensive reagents and afforded low product yields (<10%).<sup>25,26</sup> Finally, all products were characterized using  $^1\text{H}$  and  $^{19}\text{F}$  NMR spectroscopies, high-resolution mass spectrometry (HR-MS), and single-crystal X-ray crystallographic analysis.

UV-Vis absorption spectra of these Pt(II) complexes show two main absorption bands (Fig. 1), namely  $\pi-\pi^*$  absorption of the aromatic entities at  $\sim 320$  nm and the mixed ligand-centered  $\pi-\pi^*$  absorption and metal-to-ligand charge transfer (MLCT) at the region  $\geq 330$  nm. Compared to the  $\text{CF}_3$ -substituted complexes



Scheme 1 Synthetic route and chemical structures of MeCF3Pt, IPrCF3Pt and IPrtBuPt.

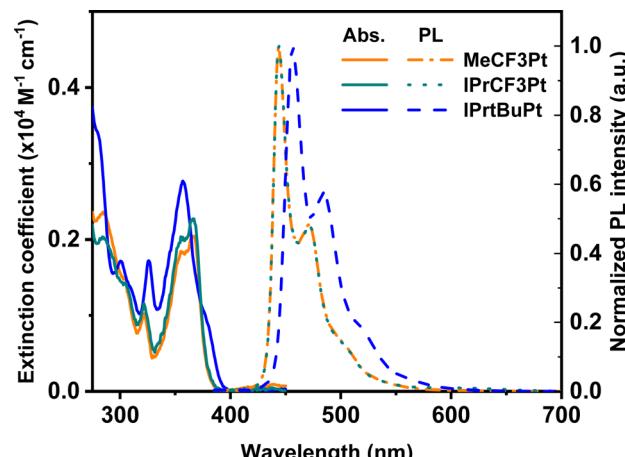


Fig. 1 Normalized UV-vis absorption (solid) and emission (dashed) spectra of the new Pt(II) complexes in DCM solution.

(MeCF3Pt and IPrCF3Pt), *t*Bu-substituted IPrtBuPt presents a slightly redshifted absorption peak (Table 1) and decreased MLCT intensity, implying a smaller energy gap and less MLCT involvement in transition. Thus, the substitution of the electron-rich group at the benzene ring will reduce the mix between the  $\pi^*$  orbital of the ligand and  $d\pi$  orbital of the metal center.<sup>41</sup> Consistently, a slightly redshift emission in IPrtBuPt (458 *vs.* 444 nm) with an enhanced 0–1 transition band (FWHM increases from 16 to 44 nm, ascribed to a greater contribution from ligand-centred  $\pi-\pi^*$  transition)<sup>42</sup> is observed in the photoluminescent (PL) spectrum in  $\text{CH}_2\text{Cl}_2$  solution ( $10^{-5} \text{ M}$ ). Concomitantly, the longest radiative lifetime (7.84  $\mu\text{s}$ ) is observed from IPrtBuPt. Isopropyl substituents in IPrCF3Pt and IPrtBuPt introduce an extra vibrational degree of freedom to the molecule and result

Table 1 Photophysical data of the Pt(II) complex in DCM ( $10^{-5} \text{ M}$ )

Name	$\lambda_{\text{Abs}}^a$ [nm]	$\lambda_{\text{em}}^a$ [nm]	PLQY <sup>ab</sup>	FWHM <sup>a</sup>	$\tau^a$ [ns]
MeCF3Pt	320, 356, 367	444, 471	73.2	16	3187
IPrCF3Pt	320, 356, 367	444, 471	46.4	16	2334
IPrtBuPt	326, 357, 376 (sh)	458, 485	46.7	44	3660

<sup>a</sup> Measured in degassed DCM. <sup>b</sup> Coumarin 102 (C102) in MeOH ( $10^{-5} \text{ M}$ , Q.Y. = 87% and  $\lambda_{\text{max}} = 480 \text{ nm}$ ) was employed as the standard.

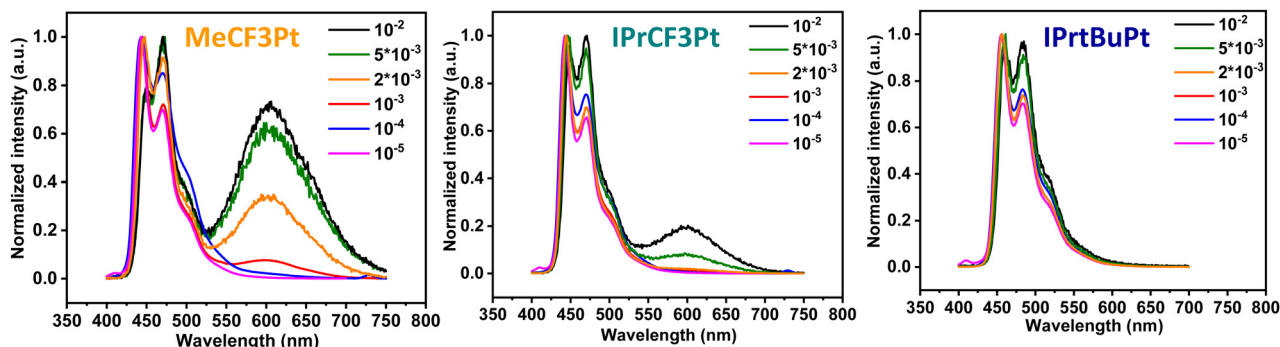


Fig. 2 Concentration-dependent PL spectra of the new Pt(II) complexes in DCM solution.

in lower photoluminescence quantum yields (PLQYs) compared to MeCF3Pt. Upon increasing the concentration from  $10^{-5}$  to  $10^{-2}$  M (Fig. 2), obvious excimer emission (peak at  $\sim 600$  nm) is observed for MeCF3Pt and IPrCF3Pt. Notably, no such lower energy shoulder was observed for IPtBuPt.

To better understand their photophysical properties, density functional theory (DFT) and time dependent-DFT (TD-DFT) calculations were carried out at the B3LYP/def2SVP level using the Gaussian16 program package.<sup>43</sup> As presented in Fig. 3, the highest occupied molecular orbitals (HOMOs) are localized on the central phenyl group of the pincer chelate and the Pt-Cl vector, while the lowest unoccupied molecular orbitals (LUMOs) are distributed over the whole molecules. Calculated HOMO and LUMO gaps followed an ascending order MeCF3Pt < IPrCF3Pt < IPtBuPt, confirming the increasing electron-donating ability of alkyl substituents between the chelates (*i.e.*, CF3/Me to CF3/IPr then *t*Bu/IPr). As the HOMO-LUMO hopping account for over 80% of the lowest energy process, the transitions between  $S_0$  and  $S_1/T_1$  are consequently assigned to an admixture of MLCT/ILCT/XLCT (X denotes a halide), and with MLCT contributions being 12.9%, 11.5% and 11.8% for MeCF3Pt, IPrCF3Pt and IPtBuPt (Table S1, ESI<sup>†</sup>), respectively. These percentages are considerably small

compared to those of Pt(II) complexes documented in literature reports (*ca.* 20%),<sup>44–46</sup> which explains the relatively long radiative lifetimes of our Pt(II) carbene pincer emitters.

Single crystal analysis reveals that MeCF3Pt, IPrCF3Pt and IPtBuPt possessed similar planar geometries but distinct packing behaviours. Selected bond lengths and angles are summarized in Table S2 (ESI<sup>†</sup>). The Pt-Cl, Pt-C<sub>Ph</sub> and Pt-C<sub>NHC</sub> lengths are 2.38–2.41, 1.93–1.94 and 2.02–2.04 Å, respectively. The Cl-Pt-C<sub>Ph</sub> and C<sub>NHC</sub>-Pt-C<sub>NHC</sub> bond angles were in the range of 177–179° and 156–158° respectively, presenting similar coordinating properties to other carbene Pt(II) pincer complexes in literature reports.<sup>25,26</sup> However, as shown in Fig. 4, intimately close Pt···Pt and  $\pi$ - $\pi$ /Pt- $\pi$  interactions are found in the unit cell of MeCF3Pt, confirming formation of excimers even at low doping concentrations. When the bulky *tert*-butyl group was added, only  $\pi$ - $\pi$  interactions were observed, as indicated in the crystal cells of IPrCF3Pt. The addition of the *tert*-butyl group in IPtBuPt also eliminated the respective face-to-face stacking interaction, explaining the photophysical properties observed earlier.

Before device fabrication, decomposition temperatures ( $T_d$ ) of all studied complexes were analysed *via* thermogravimetric measurements (Fig. S1, ESI<sup>†</sup>) and all of them present good thermal stability with  $T_d$  exceeding 300 °C, indicating they are suitable for vacuum-deposition processes. These Pt(II) complexes were employed as dopant emitters with an OLED configuration of ITO/TAPC(20 nm)/TCTA(10 nm)/mCP(10 nm)/EML (20 nm)/3TPYMB(40 nm)/LiF(1 nm)/Al, in which bipolar mCPCN is employed as the host material in the EML. As presented in Fig. S2 (ESI<sup>†</sup>) and Table 2, being consistent in their packing properties, MeCF3Pt exhibited a broadened lower energy emission even at a doping level of 1 wt%. This excimer emission (peak at 597 nm, similar to the PL observed in doped thin films, excluding the contribution of the electromer, Fig. S3, ESI<sup>†</sup>) has gained intensity in comparison to the monomer emission (peak at 447 nm) *vs.* doping concentration dramatically. The higher intensity of long wavelength peak in EL (Fig. S2c, ESI<sup>†</sup>) compared to PL (Fig. S3, ESI<sup>†</sup>) revealed using EL spectra to detect the excimer signal is more sensitive and efficient. Its EQE<sub>max</sub> was increased from 3.51 to 17.0%, which is ascribed to a favourable energy transfer from the host to the dopant and balanced carrier transport when doping concentration increased from 1 to 5 wt% and greater. Similarly, IPrCF3Pt exhibited excimer emissions starting from

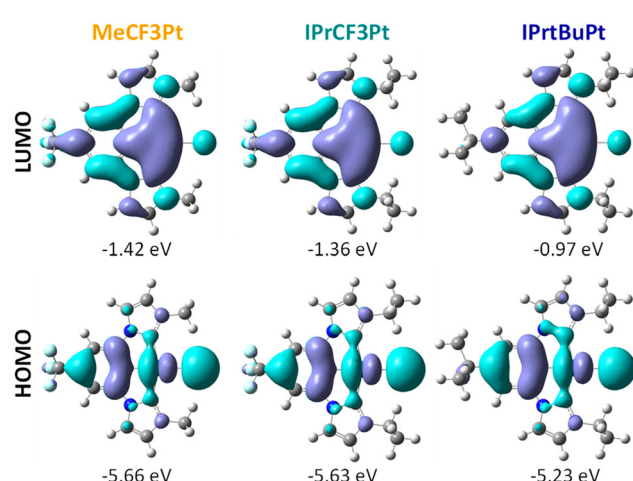


Fig. 3 Frontier molecular orbital distributions of MeCF3Pt, IPrCF3Pt and IPtBuPt.

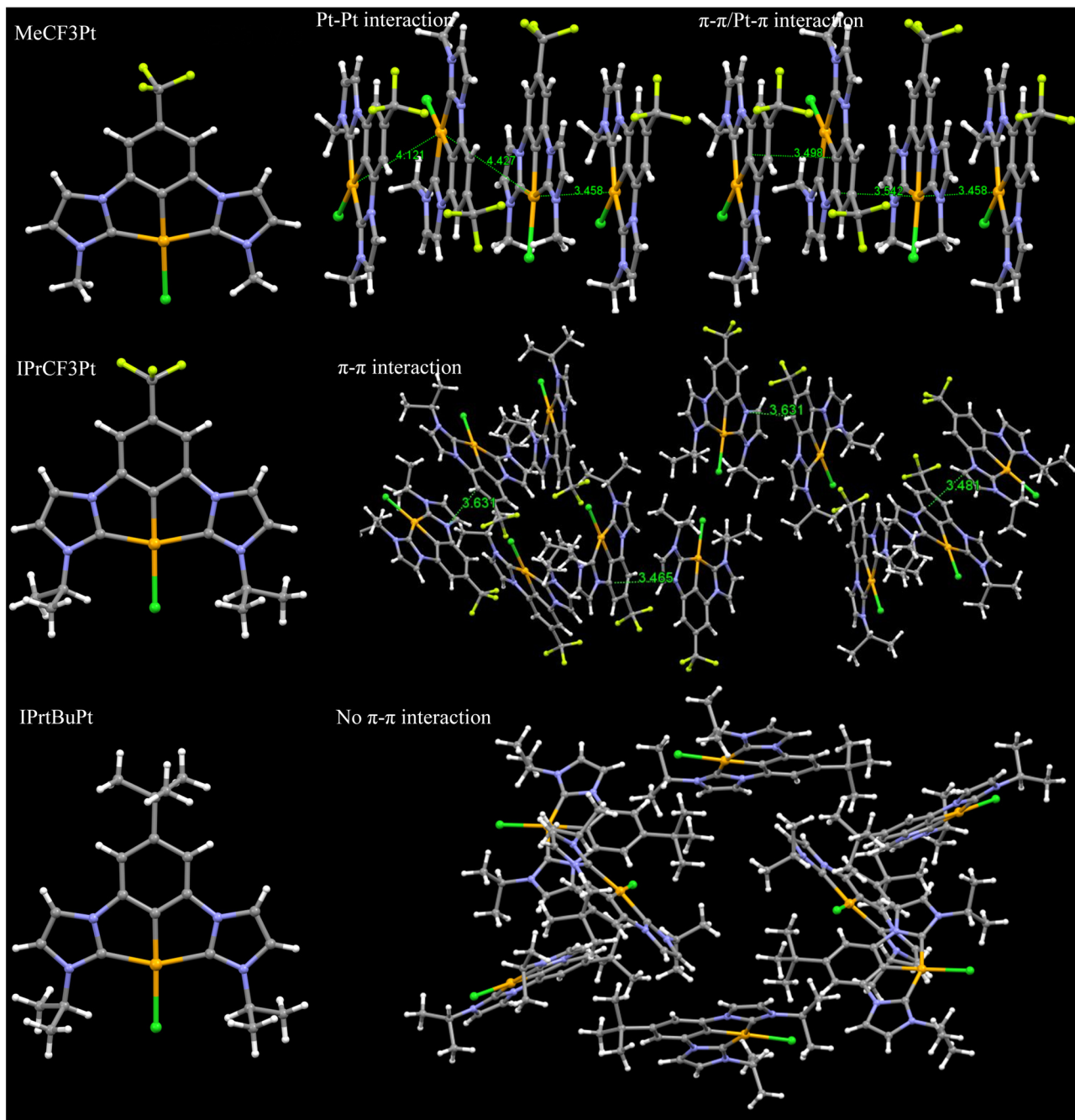


Fig. 4 Structure and intermolecular packing observed in single crystals of MeCF3Pt, IPrCF3Pt and IPrtBuPt.

8 wt% (Fig. S2f, ESI<sup>†</sup>). In sharp contrast, IPrtBuPt-based device displayed only higher energy, structured monomer emission even at 20 wt% (Fig. S2i, ESI<sup>†</sup>) and maintained a fairly stable CIE<sub>y</sub> and EQE<sub>max</sub> data of 0.221 and 12.38% @5 wt%, to 0.241 and 13.96% @20 wt%. However, the blue devices have low maximum brightness and show a dramatic efficiency roll-off with increased current density.

In view of its good resistance to excimer formation and good EL performance at high concentrations, IPrtBuPt was further employed as a dopant sensitizer for a narrow band MR-TADF terminal emitter:  $\nu$ -DABNA. Importantly, the emission spectrum

of IPrtBuPt shows a good overlap with the absorption spectrum of  $\nu$ -DABNA. By integrating the overlap as shown in the ESI,<sup>†</sup> a large FRET radius of 7.1 nm (Fig. S4, ESI<sup>†</sup>) is estimated,<sup>47,48</sup> implying the potentially effective energy transfer processes. A hyperphosphorescent device with a configuration of ITO/HATCN/HATCN:TPAPB (3:7 30 nm)/TPAPB (10 nm)/PCzAc (10 nm)/16 wt% IPrtBuPt & 1 wt%  $\nu$ -DABNA in mCBP:SiCzTrz 1:1 (30 nm)/DBFTrz (5 nm)/TmPyPB/LiF/Al was next fabricated. As shown in Fig. 5, a sharp EL profile peak at 471 nm and a narrow FWHM of 20 nm were achieved. EL spectra of the hyper device remained the same until high voltage ( $\geq 13$  V), where the

**Table 2** Summary of device performance of MeCF3Pt, IPrCF3Pt and IPrtBuPt with different doping ratios

Dopant	$V_{on}^a$ [V]	$CE_{max}^b$ [cd A $^{-1}$ ]	$EQE_{max}$ [%]	CIE $^c$ [x, y]
MeCF3Pt 1 wt%	3.5	5.36	3.51	0.326, 0.26
MeCF3Pt 3 wt%	3.5	20.36	9.77	0.454, 0.363
MeCF3Pt 5 wt%	3.5	37.52	17.0	0.529, 0.419
IPrCF3Pt 5 wt%	3.5	5.29	5.03	0.163, 1.18
IPrCF3Pt 8 wt%	3.5	10.26	9.12	0.187, 0.138
IPrCF3Pt 10 wt%	3	17.39	13.3	0.191, 0.145
IPrtBuPt 5 wt%	3.5	20.48	12.38	0.153, 0.221
IPrtBuPt 10 wt%	3.5	23.2	13.93	0.156, 0.223
IPrtBuPt 20 wt%	3.0	24.26	13.96	0.159, 0.241

<sup>a</sup> Turn-on voltage at 1 cd m $^{-2}$ . <sup>b</sup> Maximum current efficiency.

<sup>c</sup> Recorded at 100 cd m $^{-2}$ .

brightness was too high and pixels near the peaks were saturated (Fig. S5, ESI $\dagger$ ). Impressively, a much-relieved roll-off with  $EQE_{max}$  and  $EQE_{1000}$  (EQE at 1000 cd m $^{-2}$ ) of 33.59 and 26.61%, respectively, comparable to the best hyper-OLED ever reported in the literature (Fig. S6, ESI $\dagger$ ) were achieved. $^{7,8,15,24,30-34,36-38,49,50}$  However, the device lifetime of the hyper-device is only  $\sim 0.3$  h (Fig. S7, ESI $\dagger$ ), which may be attributed to the nucleophilic vulnerability of the Pt–Cl bond.

## Conclusions

In summary, Pt(II) dicarbene pincer complexes were modified using different peripheral bulky substituents. The annoying issue of excimer formation for planar Pt(II) complexes is addressed by these peripheral groups, and blue emissions can be maintained even at a high concentration of 20 wt% in the doped device of IPrtBuPt. Notably, IPrtBuPt has successfully served as a dopant sensitizer for a  $\nu$ -DABNA based hyper-OLED, giving max. EQE up to 33.59%, FWHM of 20 nm, and CIE coordinates of (0.124, 0.148). These results demonstrated a utilization of both the engineered emitters and smart device architecture to foster efficient blue OLED devices with mild efficiency roll-offs.

## Experimental details

### Synthesis

All solvents and reagents were purchased from commercial resources and used as received. All reactions were carried out in

the air unless otherwise stated. Hexafluorophosphate salts of 1, 3-bisimidazol-3-ium pro-chelates (denoted as  $[(mimf)H_3\cdot(PF_6)_2]$ ,  $[(pimf)H_3\cdot(PF_6)_2]$  and  $[(pimb)H_3\cdot(PF_6)_2]$ ), namely 1,1'-(5-(trifluoromethyl)-1,3-phenylene)bis(3-methyl-1H-imidazol-3-ium), 1,1'-(5-(trifluoromethyl)-1,3-phenylene)bis(3-isopropyl-1H-imidazol-3-ium) and 1,1'-(5-(*tert*-butyl)-1,3-phenylene)bis(3-isopropyl-1H-imidazol-3-ium), were obtained using the method depicted in our previous report. $^{40,41}$

### Synthesis of the Pt(II) chlorides

A mixture of  $[(mimf)H_3\cdot(PF_6)_2]$  (1.1 eq.),  $K_2PtCl_4$  (1 eq.) and  $K_2CO_3$  (10 eq.) was heated in refluxing propionic acid for 2 h. After cooled to RT, propionic acid was evaporated and the residue was dissolved in  $CH_2Cl_2$ . The solution was washed with deionized water, dried over anhydrous  $Na_2SO_4$ , filtered and concentrated to dryness. The crude product was purified by silica gel column chromatography, eluting with a (1 : 4) mixture of ethyl acetate and hexane to afford the targeted Pt(II) carbene pincer complexes.

**Spectral data of MeCF3Pt.**  $^1H$  NMR (400 MHz,  $CDCl_3$ , 296 K):  $\delta$  7.37 (d,  $J$  = 2.0 Hz, 2H), 7.06 (s,  $J_{PtH}$  = 16.8 Hz, 2H), 6.98 (d,  $J$  = 2.0 Hz, 2H), 4.26 (s, 6H);  $^{19}F$  NMR (376 MHz,  $CDCl_3$ , 296 K):  $\delta$  -61.17 (s, 3F). HRMS (Q-TOF)  $m/z$  for  $[M - Cl]^+$  calcd 500.0657, found 500.0639.

**Spectral data of IPrCF3Pt.**  $^1H$  NMR (400 MHz,  $CDCl_3$ , 296 K):  $\delta$  7.43 (d,  $J$  = 2.0 Hz, 2H), 7.14 (d,  $J$  = 2.0 Hz, 2H), 7.12 (s,  $J_{PtH}$  = 19.2 Hz, 2H), 6.18 (q,  $J$  = 6.8 Hz, 2H), 1.52 (d,  $J$  = 6.8 Hz, 12H);  $^{19}F$  NMR (376 MHz,  $CDCl_3$ , 296 K):  $\delta$  -61.22 (s, 3F). HRMS (Q-TOF)  $m/z$  for  $[M - Cl]^+$  calcd 556.1283, found 556.1272.

**Spectral data of IPrtBuPt.**  $^1H$  NMR (400 MHz,  $CDCl_3$ , 296 K):  $\delta$  7.39 (d,  $J$  = 2.0 Hz, 2H), 7.07 (d,  $J$  = 2.0 Hz, 2H), 6.90 (s,  $J_{PtH}$  = 16.0 Hz, 2H), 6.15 (q,  $J$  = 6.8 Hz, 2H), 1.50 (d,  $J$  = 6.8 Hz, 12H), 1.36 (s, 9H). HRMS (Q-TOF)  $m/z$  for  $[M - Cl]^+$  calcd 544.2040, found 544.2020.

### Author contributions

Y. C. and C. S. L. initiated and provided financial support for the project. Y. C. designed the complexes. J. Y., Z. L. Z. and L.W. F. synthesized and characterized the new molecules. Y. J., C. C. and Z. L. Z. carried out photophysical measurements. J. H. T performed the computational calculation under the

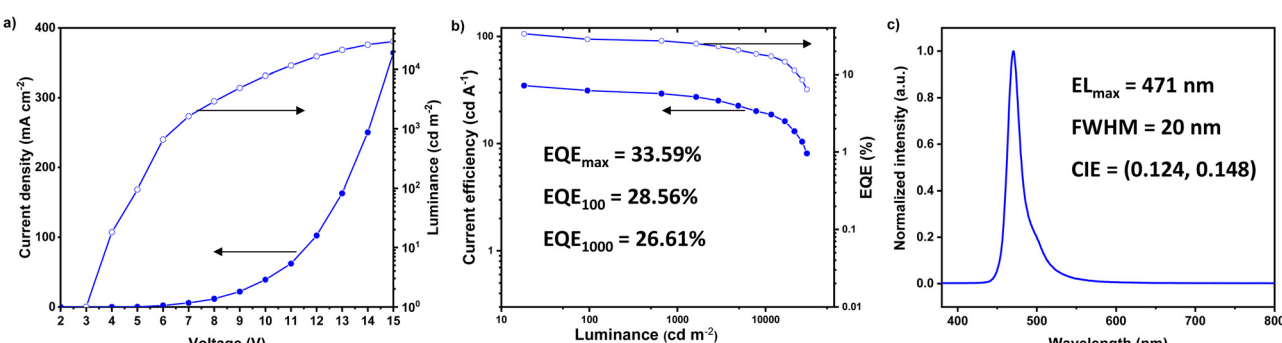


Fig. 5 (a) Current density–voltage–luminance plots, (b) current efficiency–luminance–EQE curves and (c) EL spectrum of an IPrtBuPt based hyper-OLED.

supervision of Z. L. Z. J. Y. and Z. L. Z. contributed to crystal growth and crystal data analysis. Z. L. Z. fabricated the OLEDs and measured the device performance. Y. J. and Z.L. Z. wrote the first draft. Z. L. Z., Y. C. and C.S. L. provided suggestions on experiments, revised manuscript and supervised the project. All authors discussed the experimental result and reviewed the manuscript.

## Conflicts of interest

There are no conflicts to declare.

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