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## Transition-metal phosphors with emission peak maximum on and beyond the visible spectral boundaries

Guowei Ni,<sup>a</sup> Jie Yan,<sup>b</sup> Yixin Wu,<sup>b</sup> Fan Zhou,<sup>b</sup> Pi-Tai Chou <sup>\*c</sup> and Yun Chi <sup>\*a,b</sup>

Third-row transition-metal complexes remain a key component in the preparation of light-emitting layers for future OLED technology. Hence, it is of utmost importance to expand their emission peak wavelengths into the true-blue (~460–470 nm) and near-infrared (NIR) regions (~700–1000 nm). Stable true-blue phosphors are expected to excel in commercially available lighting luminaries and visual devices, such as displays and monitors, while efficient NIR emitters will enable new applications such as sensing, imaging, and optical communication. In theory, these advanced emitters can be assembled using transition metals such as iridium and platinum and judiciously designed chelates, as elaborated in this chemistry frontier article.

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### Introduction

Organic light-emitting diodes (OLED) are particularly useful in the fabrication of full color display panels and various lighting luminaries. Typical OLED is expected to generate 25% singlet exciton and 75% triplet exciton during operation. According to this 1 : 3 spin statistics, fluorescent materials are only capable of achieving the internal quantum efficiency ( $\eta_{\text{int}}$ ) of 25%, which is defined as the ratio of emitted photons that can be extracted from the total injected carriers (or theoretical electron spin statistics) and, hence, cannot actively serve as adequate OLED emitters. To circumvent this deficiency, both phosphorescent and thermally activated delayed fluorescence (TADF) emitters were employed to replace the inferior fluorescent emitters. On the one hand, the third-row transition metal complexes, due to their better stability and efficient spin-orbit coupling induced by their central metal ion, to a large extent, would effectively remove the spin-forbidden nature of the intersystem crossing between singlet and triplet excitons, resulting in an essentially unitary  $\eta_{\text{int}}$  and bright electroluminescence.<sup>1</sup> On the other hand, TADF emitters take advantage of the significantly reduced energy gap,  $\Delta E_{\text{ST}}$ , between singlet and triplet excited states; hence, the faster

forward (ISC) and reverse intersystem crossing (RISC) result in efficient utilization of inaccessible triplet exciton back to the emitting singlet excited state. In theory,  $\eta_{\text{int}}$  of TADF emitters may also reach as high as 100%. Nowadays, both classes of emitters compete for practical industrial applications.

As for the phosphorescence emitters, the Ir(III) based metal complexes have been employed as the commercially viable red and green OLED phosphors. However, currently, virtually no true-blue Ir(III) phosphors are capable of affording both high efficiency and stability in devices, retarding their application *en route* to commercial applications.<sup>2</sup> These deficiencies encourage intensive and wide investigations on the blue emissive Ir(III) phosphors, particularly the carbene-based Ir(III) complexes,<sup>3</sup> as this class of emitters is expected to be more versatile in chemical designs and superior physical properties and efficiencies in comparison to those with Pt(II), Au(I), and Au(III) metal atoms.

Recently, both technologies termed “hyperfluorescence”<sup>4,5</sup> and “hyperphosphorescence”<sup>6</sup> (also known as the phosphor sensitized fluorescence)<sup>7</sup> have gained particular attention for the fabrication of efficient and durable blue emissive OLED devices. Their basic principle is to effectively harvest triplet excitons by TADF emitters or transition metal phosphors, which then undergo rapid Förster resonance energy transfer to blue organoboron TADF emitters having fast fluorescent lifetimes and narrow emission bandwidths mainly due to the multi-resonance effect.<sup>8</sup> This process partly solves the deadlock on inadequate efficiency and poor durability of true-blue OLED devices, with emission peak maximum located at ~460–470 nm. Concurrently, their narrowband blue emission matches well the BT.2020 color standard for ultra-HD

<sup>a</sup>Department of Chemistry, City University of Hong Kong, Kowloon Tong 999077, Hong Kong SAR, China

<sup>b</sup>Department of Materials Sciences and Engineering, and Center of Super-Diamond and Advanced Films (COSDAF), City University of Hong Kong, Kowloon Tong 999077, Hong Kong SAR, China. E-mail: yunchi@cityu.edu.hk

<sup>c</sup>Department of Chemistry, Taiwan University, Taipei 10617, Taiwan, China. E-mail: chop@ntu.edu.tw



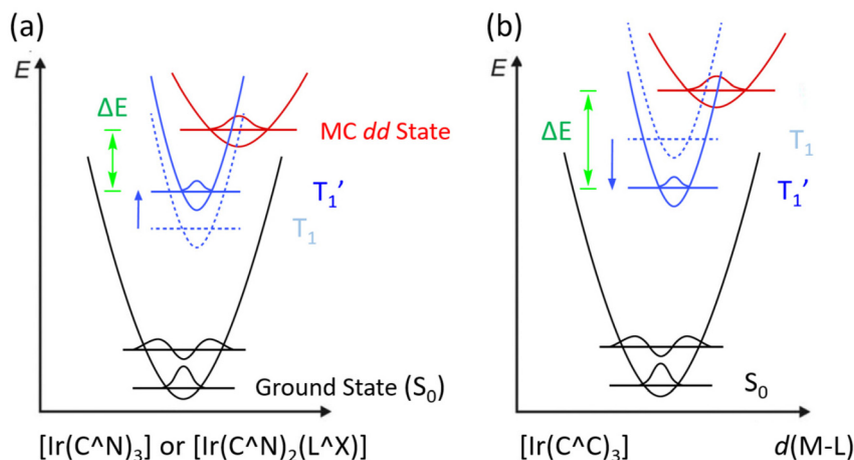
projectors and televisions. Relevant endeavor is timely, which will surely impact the future development of blue OLED devices, emphasizing both efficiency and durability. With appropriate structural modification, Ir(III) sensitized hyperphosphorescence may outperform the TADF sensitized hyperfluorescence due to the partially allowed intersystem crossing and thus better utilization of triplet excitons.

In yet another approach, near-infrared (NIR) irradiation with a peak wavelength spanning 700–1000 nm region is invisible to human eyes and cannot be employed for the fabrication of visual accessories.<sup>9</sup> Alternatively, it possesses an increased penetration depth into the human flesh, and the corresponding OLED devices worn by patients are expected to be pivotal for emerging applications such as sensing, imaging, phototherapy, and hence the fabrication of pulse oximeter and photoplethysmography.<sup>10,11</sup> As efficient NIR phosphors, Pt(II) complexes possess a  $d^8$ -electron configuration and exhibit square-planar geometries. They are in sharp contrast to Ir(III) and Os(II) metal complexes, where they possess a distinctive  $d^6$ -electron configuration with octahedral geometry.<sup>12</sup> It should be noted that in the octahedral complexes, the surrounding chelates provide effective steric shielding against the formation of any direct metal–metal contact. Differently, the square-planar Pt(II) complexes would facilitate face-to-face stacking between adjacent Pt(II) metal units, leading to intimate Pt...Pt and/or  $\pi$ ... $\pi$  interactions.<sup>13,14</sup> This linearly arranged aggregate would exert a strong influence on their photophysical properties, *i.e.*, giving metal–metal-to-ligand charge transfer (MMLCT) or even excimeric  $\pi\pi^*$  transition, rather than the combined ligand-centered  $\pi\pi^*$  and metal-to-ligand charge transfer (MLCT) characteristics. Emission originating from the MMLCT transition process is typically much red shifted with respect to the “isolated” monomers that exist in dilute solution, dispersed polymer matrix, or co-deposited thin film. Hence, the monomeric Pt(II) complexes with sky-blue and green emissions may become NIR emitters when self-assembled, thus opening the gateway to efficient NIR OLEDs.

Hence, we will use this frontier article to discuss the fundamental insights into these true-blue and NIR emissive Ir(III) and Pt(II) phosphors, together with their potential impacts on the future development of relevant OLED devices. Moreover, both classes of emitters, synthetically, were constructed using distinctive chelates and metal elements, which are judiciously selected for achieving optimal structural and photophysical properties. Thus, critical comments will also be provided on the pros and cons of their designs and perspectives so that readers can gain an in-depth understanding, beneficial for future development.

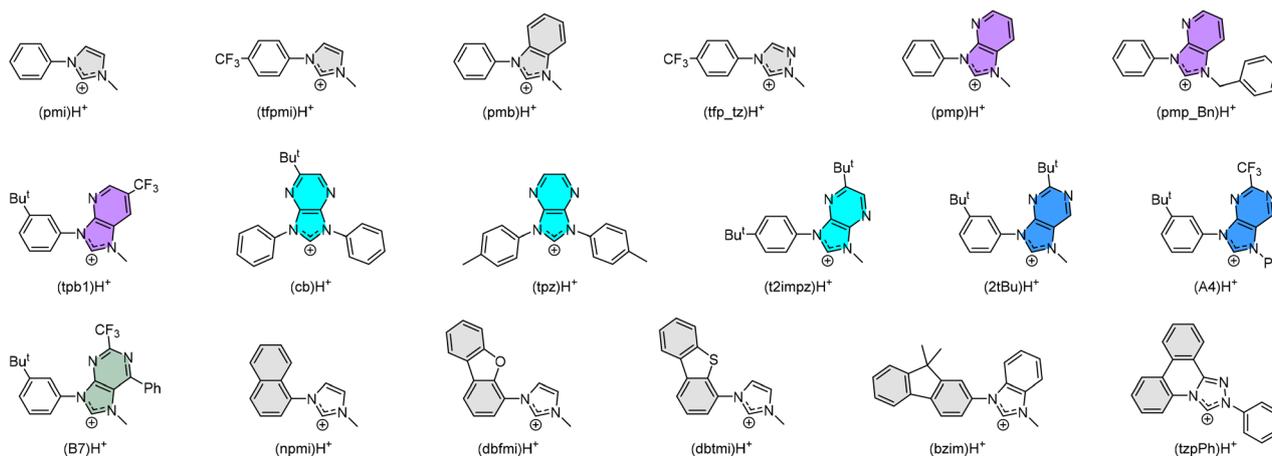
## True-blue emitters

There are two possible classes of Ir(III) complexes suitable for efficient blue emission. One involves functional cyclometalates linked to the N-donor fragment, such as pyridine, pyrazole, or imidazole, in the form of either homoleptic  $\text{Ir}(\text{C}^{\wedge}\text{N})_3$  or heteroleptic  $\text{Ir}(\text{C}^{\wedge}\text{N})_2(\text{L}^{\wedge}\text{X})$  architectures, where  $\text{C}^{\wedge}\text{N}$  denotes the N-containing aromatics and  $\text{L}^{\wedge}\text{X}$  specifies the anionic ancillary.<sup>15–17</sup> The alternatives are carbene complexes  $f$ -Ir( $\text{C}^{\wedge}\text{C}$ )<sub>3</sub> and  $m$ -Ir( $\text{C}^{\wedge}\text{C}$ )<sub>3</sub>,<sup>18–21</sup> where  $f$ - and  $m$ - indicate facial and meridional orientation, respectively. Their proposed energy diagrams are depicted in Scheme 1. The most well-known examples of Ir(III) based pyridine and carbene-containing complexes are sky-blue bis[2-(4,6-difluorophenyl)pyridinato- $\text{C}^2, \text{N}$ ] (picolinato)iridium(III) (**FIrpic**)<sup>22</sup> and purple emitting tris(1-phenyl-3-methylbenzimidazolin-2-ylidene- $\text{C}, \text{C}'$ )iridium(III), [Ir(pmb)<sub>3</sub>],<sup>23</sup> for which the ground state ( $S_0$ ), lowest energy triplet excited state ( $T_1$ ), and upper lying metal-centered (MC) dd excited state are indicated in black, dotted blue, and red traces, respectively. The  $T_1$  excited state of both complexes constitutes a mixed ligand-centered  $\pi\pi^*$  and MLCT transition, while the high-lying MC dd state involves a  $T_1$  state geometry with metal–ligand distances lengthened, *i.e.*, an activated radiationless state close to the  $T_1$  excited state.<sup>24</sup> The Ir( $\text{C}^{\wedge}\text{C}$ )<sub>3</sub>



**Scheme 1** Potential energy surface diagrams for hypothetical Ir(III) complexes with either pyridine (a) or carbene-based cyclometalates (b) and strategic approach in obtaining blue emission; definition of  $T_1$  and  $T_1'$  states is elaborated in the text.





**Scheme 2** Some representative carbene cyclometalates employed in the preparation of homoleptic Ir(III) metal complexes.

complexes exhibit more destabilized  $T_1$  and MC dd excited states than those of  $\text{Ir}(\text{C}^{\wedge}\text{N})_3$  and  $\text{Ir}(\text{C}^{\wedge}\text{N})_2(\text{L}^{\wedge}\text{X})$  due to the high-lying  $\pi^*$  orbital of carbene fragments and stronger Ir–C bonding interaction. With this in mind, tuning emission from sky-blue **Irpic** and analogues to true-blue color can be done by destabilizing the  $T_1$  state, *i.e.*, the addition of electron-withdrawing (or donating) group at the HOMO (or LUMO) segment of Ir(III) complexes, to which the newly obtained  $T_1$  state, *i.e.*,  $T_1'$  state, is now indicated in blue solid line (Scheme 1(a)). Accordingly, this manipulation is expected to reduce the  $T_1'$ -MC dd energy gap ( $\Delta E$ ), inducing a faster emission quenching. Conversely, tuning emission from purple  $[\text{Ir}(\text{pmb})_3]$  and derivatives to true-blue required stabilization of the  $T_1$  state *via* the addition of electron-deficient substituent(s) at the carbene entity, resulting in the increase of  $\Delta E$  (see Scheme 1(b)). The enlarged  $T_1'$ -MC dd energy gap would retard the non-radiative decay process, making them one of the best candidates for showing both efficient and durable blue emission.

Practically, tuning emission to blue can be easily achieved according to the aforementioned strategies. Scheme 2 depicts the structures of parent carbene pro-chelates  $(\text{pmi})\text{H}^+$ ,<sup>23</sup>  $(\text{tfpmi})\text{H}^+$ ,<sup>25</sup> and  $(\text{pmb})\text{H}^+$ ,<sup>23</sup> together with a series of functional derivatives with an aim to reduce the  $T_1'$ -MC dd energy gap. Chemically, this can be done *via* the addition of electron-deficient substituent(s) to the carbene fragment or, alternatively, extending the  $\pi$ -conjugation of carbene at the *N*-aryl site as the less effective methodology.

The positive effects on Ir(III) complexes with electron-deficient carbene can be comprehended as follows: (i) Carbene complexes  $[\text{Ir}(\text{pmb})_3]$  with *m*- and *f*-arranged benzo[*d*]imidazolylidene chelates exhibit ultraviolet emission at  $\lambda_{\text{max}} = 389$  nm and 395 nm, and radiative lifetimes of 5.5 and 7.5  $\mu\text{s}$  in degassed solution at RT,<sup>23,26</sup> whereas the corresponding imidazo[4,5-*b*]pyridin-2-ylidene complexes *m*- $[\text{Ir}(\text{pmp})_3]$  and *f*- $[\text{Ir}(\text{pmp})_3]$ <sup>27</sup> gave better emission at  $\lambda_{\text{max}} = 465$  nm and 418 nm and much shorter radiative lifetimes of 1.58 and 1.03  $\mu\text{s}$ , respectively. (ii) Carbene complex *f*- $[\text{Ir}(\text{cb})_3]$  with imidazo[4,5-*b*]pyrazin-2-ylidene chelates displays emission at 456 nm in

toluene with a high quantum yield of 88% and radiative lifetime of 0.47  $\mu\text{s}$  in polystyrene matrix at 1 wt%.<sup>28</sup> Despite being a mixture of isomers, *f*- $[\text{Ir}(\text{cb})_3]$  has been successfully used in the fabrication of blue OLED devices with high efficiency and steadily improved device lifespans.<sup>29,30</sup> This is in agreement with the high chemical stability of an analogue *f*- $[\text{Ir}(\text{tpz})_3]$  in the solid state.<sup>31</sup> (iii) In sharp contrast to the Ir(III) complexes with electron-deficient carbene chelates such as  $(\text{pmp\_Bn})\text{H}^+$ ,<sup>32</sup>  $(\text{tpb1})\text{H}^+$ ,<sup>33</sup>  $(\text{t2impz})\text{H}^+$ ,<sup>34</sup>  $(2\text{tBu})\text{H}^+$ ,<sup>35</sup>  $(\text{A4})\text{H}^+$ ,<sup>36</sup> and  $(\text{B7})\text{H}^+$ ,<sup>36</sup> those with extended  $\pi$ -conjugation, *i.e.*,  $(\text{npmi})\text{H}^+$ ,<sup>37</sup>  $(\text{dbfmi})\text{H}^+$ ,<sup>38</sup>  $(\text{dbtmi})\text{H}^+$ ,<sup>39,40</sup>  $(\text{bzim})\text{H}^+$ ,<sup>41</sup> and even  $(\text{tzpPh})\text{H}^+$ ,<sup>42</sup> can only afford Ir(III) complexes having structured emission, long radiative lifetime, and relatively poor quantum yield, the latter of which points out the inferiority of the associated designs. (iv) Homoleptic Ir(III) carbene emitters with electron-deficient carbene chelates, *i.e.*,  $(\text{tfp\_tz})\text{H}^+$ ,<sup>43</sup> and others have all been adequately tested in giving prominent blue hyperphosphorescence, where the device longevity is still at the top of the agenda and needs improvement. So far, one such carbene emitter had already achieved both high efficiency and long device lifetime of  $\text{LT}_{50}$  of 431 h for an initial brightness of 500  $\text{cd m}^{-2}$ ,<sup>44</sup> confirming their potential as desirable blue emitters.

## Near-infrared emitters

Maximizing possible Pt...Pt stacking in the solid state should be one of the most promising strategies in inducing the red-shifted emission, and hence, we proceeded to synthesize representative Pt(II) metal complexes, namely:  $\text{Pt}(\text{fppz})_2$ ,<sup>45,46</sup>  $\text{Pt}(\text{fprpz})_2$ ,<sup>47</sup> **(4H)**,<sup>48</sup> **(DR)**, and **(Per-DR)**,<sup>49</sup> as shown in Scheme 3. It is notable that all of them possess the electron-deficient 3-trifluoromethyl pyrazolyl and pyrimidinyl coordination fragments, to which the pyrimidinyl fragment offered further red-shifted emission attributed to the stronger Pt–C covalent interaction. Also, the uncoordinated N-atom of these fragments is





**Scheme 3** Representative Pt(II) complexes showing strong solid-state aggregation and efficient emissions in saturated red and NIR regions.

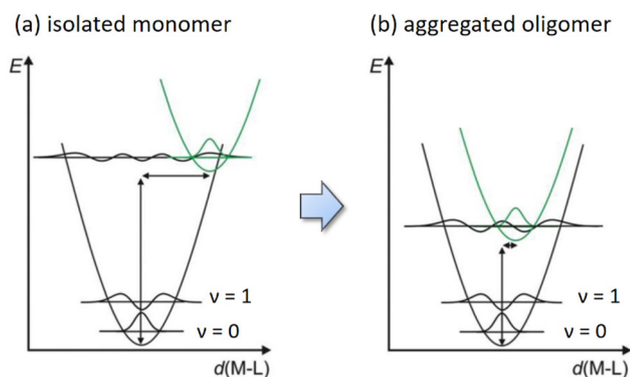
expected to foster two inter-chelate N...H-C hydrogen bonding, giving nearly planar coordination geometry.

Moreover, the electron deficiency of chelates would pass to the central Pt(II) atom, allowing the easy formation of Pt...Pt interaction (for sharing their electrons) in the solid state upon grinding or forming a thin film *via* thermal vapor deposition. In contrast, disruption of Pt...Pt stacking can be achieved by replacement of CF<sub>3</sub> substituents of Pt(fppz)<sub>2</sub> or Pt(fprpz)<sub>2</sub> with electron-donating methyl or *tert*-butyl substituent.<sup>45</sup> The increased electron richness turned corresponding Pt(II) complexes from sparingly soluble to highly soluble in common organic solvents at RT and afforded typical MLCT +  $\pi\pi^*$  transition characteristics. In sharp contrast, the electron-deficient Pt(II) complexes (*i.e.*, those depicted in Scheme 3) exhibited higher energy emission in diluted and degassed solution at RT or upon co-deposition with host material or inert media in a doped thin film,<sup>50</sup> indicating the prohibition of Pt...Pt interaction. Such a remarkable photophysical behavior can be explained using two potential energy diagrams shown in Scheme 4.

Hence, as isolated single molecules, their excited state structures are expected to be significantly different from that of the ground state (*i.e.*, greater deviation in d(M-L)). Consequently, it causes a significant displacement of the equi-

librium distance in the corresponding potential energy curve, as shown in Scheme 4a. However, upon the formation of aggregated oligomers, the excited state characteristics are now dominated by the MMLCT transition, which lowered the potential energy of the excited state in comparison to that of the monomer. The energy gap of aggregated Pt(II) complexes is thus reduced, most likely to the NIR region, depending on the chelate design. Moreover, the generated exciton will be delocalized along the aligned molecules upon excitation, where the associated vibration modes will be evenly partitioned over all effective stacked molecules, *i.e.*, the N number involved in the exciton delocalization. Theoretically, this is equivalent to reducing the vibrational reorganization energy from  $\lambda$  of the monomeric complex to  $\lambda/N$  in the N exciton delocalization, producing a less displaced excited-state potential energy surface (PES) with respect to the PES of the ground state (see Scheme 4b).<sup>48</sup> The delocalized exciton, therefore, results in the exciton-vibration decoupling that reduces the rate of internal conversion, which virtually bypasses the radiationless quenching imposed by the energy gap law. The net result is the generation of efficient NIR emission, leading to all Pt(II) complexes depicted in Scheme 3 exhibiting very high emission efficiencies at peak wavelengths beyond 700 nm and approaching the NIR II region starting from 1000 nm.

Finally, for further minimization of exciton-vibration coupled deactivations, we executed the H-D substitution of Pt(II) derivative DR for reducing the high-frequency C-H vibrations, where the difference in energy between C-H and C-D promoting stretch modes is  $\sim 800\text{ cm}^{-1}$ . Therefore, for the C-D substituted Pt(II) complexes, according to the energy gap law, upon electronic-vibrational (C-D stretching) coupling, the accepting vibrational modes, such as those C=C vibrations associated with  $\pi\pi^*$  transition or d(M-L) changed vibration associated with MLCT, required higher quanta to match the energy gap than that of the C-H complexes. Therefore, the C-D complexes are subject to a lower internal conversion rate than the corresponding C-H analogues, especially in the NIR region. This operation is nontrivial and requires a multi-step synthetic protocol. Worth all the effects, photoluminescence with max. at  $\sim 1000\text{ nm}$  and quantum yield of  $23 \pm 0.3\%$  was obtained for the fully deuterated Pt(II) derivative Per-DR.<sup>49</sup> In addition to the improved photophysical data, the respective



**Scheme 4** Schematic potential energy diagrams for square planar Pt(II) emitters (a) as isolated molecules and (b) after aggregation and formation of intimate Pt...Pt interaction.



NIR OLED device gave a peak wavelength of 995 nm, maximum external quantum efficiency (max. EQE) of 4.31%, and radiance of  $1.55 \text{ W Sr}^{-1} \text{ m}^{-2}$  at  $10 \text{ mA cm}^{-2}$ , marking a new milestone for the studies.

## Challenges and future perspectives

Several challenges need to be solved before efficient and durable true-blue and NIR-emitting OLED devices can be utilized for possible commercial applications. As to the true-blue phosphors, according to our analysis, homoleptic Ir(III) carbene complexes should outperform their N-donor counterparts in terms of operational stability, attributed to the increased ligand–metal bond strength. Moreover, Ir(III) emitters are more stable than the Au(I) and Cu(I) based blue emitters;<sup>51,52</sup> the latter possess only the monodentate ligands, of which their ligand–metal bond strength should be less robust than the bidentate bonding in the Ir(III) complexes. Under similar criteria, the bidentate metal–chelate interaction of Ir(III) based emitters is also less stable than that of the tetradentate coordinative Pt(II) based blue emitters,<sup>53</sup> but the latter suffers from longer radiative lifetimes due to the diminished total zero-field splitting ( $\Delta E(\text{ZFS})$ ) of the emitting triplet state.<sup>1</sup> The higher the  $\Delta E(\text{ZFS})$ , the larger the MLCT character of the emitting triplet state and the more efficient the spin–orbit coupling. Hence, the fast radiative decay of Ir(III) complexes helps to retain their candidacies in the lists of useful blue emitters.

Notably, the majority of these electron-deficient Ir(III) carbene complexes have already shown adequate emission peak maximum at  $\sim 460\text{--}470 \text{ nm}$ , high quantum yield  $\geq 85\%$ , a shortened radiative lifetime of  $\leq 1 \mu\text{s}$ , and reasonably good chemical and thermal stability. The next task is to push up their chemical and photophysical stabilities to meet stringent industry standards. After that, new designs and synthetic protocols should be executed to yield only a single product in high yields out of mixed products such as *f*- and *m*-isomers and other configurational derivatives. This effort should lower the production costs of emitters and relevant OLED devices. Finally, various functional groups need to be added to the emitters for the reduction of unwanted emitter–host interactions, which could deteriorate the overall performance of OLED devices. These groups may include the inert and bulky *tert*-butyl substituents or other functionalities that could improve the injection and transportation of electrons and holes and the balance of charge carriers within the light-emitting layer of OLED devices.

For NIR emitters and OLED devices, one of the most accessible directions is the design and preparation of new and better emitters and the fabrication of NIR OLED devices thereof. Despite that, we have already made Pt(II) emitters and NIR OLED devices showing a peak maximum at  $\sim 1000 \text{ nm}$  and maximum EQE of over 4%; there is still a large plateau for NIR emitters with a peak maximum at  $\sim 800 \text{ nm}$ . Emissions with a peak maximum located in this specified region will be totally

invisible to humans, and devices with a peak maximum of  $850 \text{ nm}$  (also at  $1310$  and  $1550 \text{ nm}$ ) may be useful for applications involving optical fiber communications. Higher efficiency is expected for emissions with high energy (*i.e.*,  $750\text{--}850 \text{ nm}$ ) as predicted by the energy gap law. A priori option is to fabricate NIR emitters with a peak maximum at  $\sim 800 \text{ nm}$  and with higher efficiency, such that the emitters can be tested for industrial applications. Moreover, our NIR OLED devices are currently fabricated using thermal vacuum deposition-induced self-assembly in forming the closely stacked (or aligned) Pt(II) molecules for the non-doped EML. Therefore, both the emission wavelength and efficiency will be critically dependent on deposition parameters. Theoretically, this obstacle can be circumvented using Pt(II) based dimers or even trimers with inherently connected Pt...Pt interaction. If these emitters are accessible, OLED devices can then be fabricated using either doping technology or a solution process. Both methods are expected to reduce wastage and associated higher costs. Moreover, they shall be suitable for bio-imaging and sensing applications, attributed to the anticipated high solubility in organic solvents and aqueous systems.

## Conflicts of interest

The authors have no conflicts to declare.

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