Chemical Science

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Cite this: Chem. Sci., 2020, 11, 2640

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Efficient white polymer light-emitting diodes (WPLEDs) based on covalent-grafting of [Zn₂(MP)₃(OAc)] into PVK⁺

Guorui Fu,‡^a Yani He,‡^a Wentao Li,^a Tiezheng Miao,^a Xingqiang Lü, ^b*^a Hongshan He, ^b^b Li Liu ^b*^c and Wai-Yeung Wong ^b*^d

Thanks to the straightforward white light of single grafting-type polymers based on earth-abundant Zn(II)complexes, producing cost-effective flexible WOLEDs/WPLEDs with good device performance remains a formidable challenge. Herein, by using the polymer **Poly(NVK-co-[Zn₂(MP)₃(OAc)])** with excellent physical properties for single-layer WPLEDs, record-high efficiencies ($\eta_c^{Max} = 13.0 \text{ cd A}^{-1}$, $\eta_p^{Max} = 6.1 \text{ lm}$ W^{-1} and $\eta_{EQE}^{Max} = 9.2\%$) and low (*ca.* 25%) efficiency roll-off compared to previous organo-Zn²⁺-based WOLEDs/WPLEDs are realized. This finding renders single Zn(II)-complex-grafted polymers a new route to low-cost and large-area flexible WPLEDs for potential full-colour flat displays.

Received 19th October 2019 Accepted 29th January 2020

DOI: 10.1039/c9sc05288g

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Introduction

Since the pioneering work¹ on white organic light-emitting diodes (WOLEDs) by Kido et al. through a typical trichromatic integration approach, WOLEDs have attracted great interest due to their potential applications² in full-colour displays, ambient lighting and backlights for liquid crystal displays. Especially with the aim of cost-effective flexible devices for industry-scale production, a solution-processable technology for WOLEDs³ based on small molecule emitters and polymer-matrix largearea white polymer light-emitting diodes (WPLEDs)⁴ is preferred. During the pursuit of high performance, although emitting materials, especially those based on iridium(III)⁵ or platinum(II) complexes⁶ are highly praised and some have already been demonstrated to offer industrial applications,7 the intrinsic high cost associated with these rare metals impedes their popularity. Toward a solution, earth-abundant metal resources⁸ including zinc(II) should be considered as an alternative. On one hand, the price9 of Zn(II)-precursors is

^bDepartment of Chemistry, Eastern Illinois University, Charleston, IL 61920, USA

^cHubei Collaborative Innovation Center for Advanced Organic Chemical Materials, Ministry of Education Key Laboratory for the Synthesis and Application of Organic Functional Molecules, School of Chemistry and Chemical Engineering, Hubei University, Wuhan 430062, China. E-mail: liulihubei@hubu.edu.cn

‡ These authors contributed equally and should be considered co-first authors.

approximately 1000–1400 times cheaper than K_2PtCl_4 and $IrCl_3 \cdot xH_2O$. Moreover, in terms of synthesis, Zn(n)-complexes are advantageous with straightforward conversion and significantly higher yields, unlike the typical two-step procedure¹⁰ for both Ir(m)- and Pt(n)-complexes. More importantly, through the smooth structure modification of stable Zn(n)-complexes, their desirable high efficiency and colour-tunable capabilities can also be realized.

Despite concerted efforts toward monochromatic (blue-,¹¹ green-,¹² yellow-,¹³ orange-¹⁴ or red-light¹⁵) electroluminescent devices from Zn(II)-complexes, their reliable panchromatic devices (WOLEDs/WPLEDs) greatly underperform. In this regard, benefiting from the volatility of most Zn(II)-complexes, a vacuum-deposition procedure for their WOLEDs was employed. One approach is through the mixing of Zn(II)complex-based primary colours in a single emitting layer, where the high-energy-state Zn(II)-complex as the host¹⁶ and/or sensitizer¹⁷ and the low-energy-state complex as the guest¹⁸ can be adopted to produce electrically-driven white light with the desired energy transfer between multiple small-molecule chromophores. Another alternative relies on a multilayer structure consisting of two or more separate emitting layers for WOLEDs, where the superposition of a Zn(II)-complex within different primary colours,¹⁹ and/or an additional colour-compensation layer by exciplex formation,²⁰ is also capable of white-light modulation. In comparison, multi-layered WOLEDs,19,20 especially those with facilitated carrier-transport abilities, exhibit significantly higher efficiencies ($\eta_c = 1.39$ –1.65 cd A⁻¹ to 5.2 cd A^{-1} and η_p up to 1.43 lm W^{-1}) at high luminance (10^3-10^4 cd) m $^{-2}$) than those ($\eta_{\rm c}=0.12$ –0.28 cd A $^{-1}$ and $\eta_{\rm p}=0.018$ –0.082 lm W^{-1} at $10^2 - 10^3$ cd m⁻²) of single-unit devices.¹⁶⁻¹⁸ Nonetheless, besides the inevitable high cost of the vacuum-deposition unsatisfactory performance-based procedure, device

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[&]quot;School of Chemical Engineering, Northwest University, Xi'an 710069, Shaanxi, China. E-mail: lvxq@nwu.edu.cn

^dDepartment of Applied Biology and Chemical Technology, The Hong Kong Polytechnic University, Hung Hom, Hong Kong, China. E-mail: wai-yeung.wong@polyu.edu.hk

[†] Electronic supplementary information (ESI) available: Starting materials and characterization methods; detailed synthesis and characterization; XRD; quantum calculations. CCDC 1894830. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/c9sc05288g

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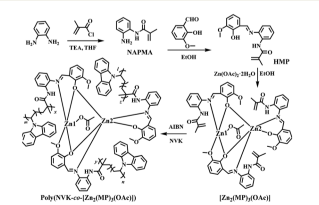
crystallization and discrepant components-induced white-light instability pose problems for the two strategies. Convincingly, the circumvention of these problems, to some extent, cannot depend on Zn(II)-based white-light-emitting coordination polymers²¹ with a lack of inherent processing ability, but can only rely on the doping of Zn(II)-complexes²² and/or sensitizers²³ into a polymeric host with a deep LUMO-HOMO bandgap for solution-processable and large-area WPLEDs. The attractive η_c of 14.67 cd $A^{-1}, \eta_{\rm p}$ of 6.58 $\rm lm \, W^{-1}$ and $\eta_{\rm EQE}$ of 6.88% are traded with detrimental deficiency with the inferior CIE (Commission Internationale de l'Eclairage) chromaticity coordinate of (0.42, 0.44) and unstable white light from a phase separation arises during incompatible multi-component integration. As a further solution, a conceptual strategy to use covalent-grafting of a Zn(II)-complex into a polymeric host is considered, where through the molecular dispersion of organo-Zn(II) guests into the colour-compensatory polymer host for high homogeneity, colour-tunable white light should also be realized for the single Zn(II)-complex-grafted polymeric emitter with motivated hostto-guest energy transfer. Herein, based on the copolymerization of NVK with efficient yellow-light tris-vinyl-functionalized complex monomer $[Zn_2(MP)_3(OAc)]$ for the newly-formed polymer Poly(NVK-co-[Zn₂(MP)₃(OAc)]) (Scheme 1) with a blue-light PVK support, photo-luminescent colour-tuning to white light after PVK-to-[Zn₂(MP)₃(OAc)] energy transfer is expected through judicious adjustments of both the grafting concentration and the excitation wavelength. Moreover, desirable WPLEDs fabricated from that single Zn(II)-complex-grafted polymeric emitter are also pursued.

Results and discussion

Synthesis and characterization of the ligand HMP and its complex monomer [Zn₂(MP)₃(OAc)]

Also as shown in Scheme 1, condensation of the vinyl-modified organic precursor **NAPMA** with 2-hydroxy-3-methoxybenzaldehyde gave rise to the vinyl-modified Schiff-base ligand **HMP** in a yield of 75%. By further treating it with

 $Zn(OAc)_2 \cdot 2H_2O$, a yellow microcrystalline product of the trisvinyl-functionalized complex monomer $[Zn_2(MP)_3(OAc)]$, soluble in common solvents except water, was isolated with a good yield of 60%. HMP and its complex monomer [Zn₂(-MP)₃(OAc)] were well-characterized by elemental analysis, FT-IR, ¹H NMR and ESI-MS. Especially in the ¹H NMR spectrum (Fig. S1[†]) of [Zn₂(MP)₃(OAc)], besides the stipulated proton molar ratio of 3 : 1 between the deprotonated (MP)⁻ ligands and the OAc⁻ anion, almost no shifted vinyl-characteristic proton resonances ($\delta = 5.75$ and 5.52 ppm) are observed in comparison with those ($\delta = 5.77$ and 5.53 ppm) of the free **HMP** ligand, despite the coordination of two Zn²⁺ ions. The molecular structure of [Zn₂(MP)₃(OAc)]·EtOH·H₂O was determined by Xray single-crystal diffraction (XRD) analysis with the crystallographic data collected in Tables S1 and S2.[†] [Zn₂(MP)₃(OAc)]. EtOH \cdot H₂O crystallizes in the monoclinic space group of P2(1)/c, where the structure unit is composed of one neutral molecule $[Zn_2(MP)_3(OAc)]$, one solvate EtOH and one solvate H₂O. As shown in Fig. 1, for the host part, two [Zn(MP)]⁺ portions with similar N^{\wedge}O-chelation (N2^{\wedge}O2 or N5^{\wedge}O8) modes for each (MP)⁻ ligand are bridged by the third $(MP)^-$ ligand with a N^{\O\O}chelation (N3^{\cap}O5^{\cap}O4) mode, resulting in the formation of} a homoleptic binuclear framework. Despite the similar fivecoordinate character of each Zn²⁺ center (Zn1 or Zn2), the distorted square pyramidal geometry of Zn1 consists of one cis- N_2O_2 core ($N2^{\wedge}O2$ and $N3^{\wedge}O5$) from two (MP)⁻ ligands as the base plane and one phenoxide-O (O8) from the third (MP)⁻ ligand at the apical position. In contrast, one cis-NO₃ core $(N5^{\circ}O8 \text{ and } O4^{\circ}O5)$ from two $(MP)^{-}$ ligands as the base plane and the axial occupation by one O atom (O10) from the monodentate OAc⁻ anion contribute to the five-coordinate environment of the other Zn^{2+} center (Zn2). The two Zn^{2+} centers (Zn1 and Zn2) are bridged by two phenoxide-O atoms (O5 from the N^{O} O-chelating (MP)⁻ ligand and O8 from the N^{O} -chelating $(MP)^{-}$ ligand), giving a Zn…Zn separation of 3.206(2) Å. It is worth noting that each of the terminal methacrylamide groups of the three (MP)⁻ ligands do not participate in the coordination, while their tris-vinyl functionality (1.290(20)–1.357(13) Å of



Scheme 1 Reaction scheme for the synthesis of the precursor NAPMA, the vinyl-modified Schiff-base ligand HMP, the tris-vinyl-functionalized complex monomer $[Zn_2(MP)_3(OAc)]$ and the grafting polymer Poly(NVK-co- $[Zn_2(MP)_3(OAc)]$).

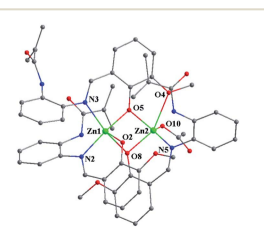
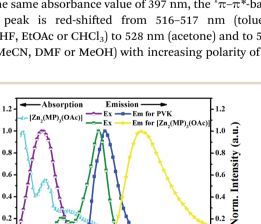


Fig. 1 Schematic of the homoleptic binuclear framework in the trisvinyl-functionalized complex monomer $[Zn_2(MP)_3(OAc)]$ ·EtOH·H₂O. All H atoms and solvates are omitted for clarity.

the typical C=C bond lengths) renders $[Zn_2(MP)_3(OAc)]$ active in the following copolymerization. Thermogravimetric (TG) analysis (Fig. S2[†]) of [Zn₂(MP)₃(OAc)] shows its favourable thermal stability at a decomposition temperature $(T_d, \text{ corre-}$ sponding to 5% weight loss) up to 290 °C.

Photophysical property and electronic structure calculation of the complex monomer $[Zn_2(MP)_3(OAc)]$

The photophysical property of [Zn₂(MP)₃(OAc)] was examined in different solutions or in the solid-state at RT or 77 K, and summarized in Table S3[†], Fig. 2, S3 and S4.[†] As shown in Fig. 2, [Zn₂(MP)₃(OAc)] in MeCN solution at RT shows a distinctively broadened absorption compared to that (Fig. S3[†]) of the free **HMP** ligand; the intense absorption bands ($\lambda_{ab} = 230$ and 290 nm) within the high-energy range similar to those present in the free HMP ligand can be assigned to the intraligand π - π * transition, and the other relatively weak bands ($\lambda_{ab} = 404, 414$ and 448 nm) extended to 500 nm should be attributed to a Zn²⁺perturbed intraligand $\pi - \pi^*$ transition. Upon excitation ($\lambda_{ex} =$ 397 nm), an intense and broad emission with $\lambda_{em} = 538$ nm is observed for [Zn₂(MP)₃(OAc)] in MeCN solution, giving rise to a bright yellow light with CIE chromaticity coordinate x = 0.487and y = 0.506. The outstanding luminescence property of $[Zn_2(MP)_3(OAc)]$ is further validated by an attractive quantum efficiency ($\Phi_{\rm em}$) of 63%, which is top-level among the reported Zn(II)-complexes.²⁴ The emission at 538 nm for [Zn₂(MP)₃(OAc)] decays biexponentially with lifetimes of 0.26 and 1.74 ns, indicating that it is characteristic of two Zn(II)-centered species arising from the $\pi^{-}\pi^{+}$ -based RT fluorescence. By contrast, a distinctively red-shifted emission at 590 nm (also Fig. S3⁺) for $[Zn_2(MP)_3(OAc)]$ in MeCN solution at 77 K is observed, and its phosphorescent nature can be further confirmed by the biexponentially time-decayed lifetimes of 0.63 and 2.12 µs. Notably, the RT fluorescence behaviours of [Zn₂(MP)₃(OAc)] in different solvents show the solvatochromism effect (Fig. S4[†]), in which, apart from a positive influence on the fluorescence enhancement at the same absorbance value of 397 nm, the $\pi^{-}\pi^{+}$ -based emission peak is red-shifted from 516-517 nm (toluene, CH₂Cl₂, THF, EtOAc or CHCl₃) to 528 nm (acetone) and to 536-538 nm (MeCN, DMF or MeOH) with increasing polarity of the



0.4

0.2

450 500 550 600 650

Wavelength (nm)

400

solvent. Moreover, contributing to the intermolecular interactions, the aggregation-induced bathochromic shift (also Fig. S4[†]) at 552 nm for [Zn₂(MP)₃(OAc)] in the solid state relative to that ($\lambda_{em} = 538$ nm) in MeCN solution at RT is also observed.

To explore the real origin of the photophysical properties of [Zn₂(MP)₃(OAc)], time-dependent density functional theory (TD-DFT) calculations based on the optimized S₀ geometry were performed, and summarized in Table S4[†] and Fig. 3. As shown in Fig. 3, the frontier molecular orbitals (FMOs) of the S₀ state are primarily dominated by the π orbitals originating from the three coordinated (MP)⁻ ligands, and the contribution from the two Zn^{2+} ions (d_{π}) or the coordinated $(OAc)^{-}$ anion appears to be distinctly small. The HOMO and LUMO are mainly (82.76% and 80.59%) localized on the MP1 (N2[^]O2-MP)⁻; from the obvious spatial overlap, the strong optical absorption should correspond to the transition from HOMO to LUMO. However, the LUMO+1 or the LUMO+2 is predominantly (95.44% or 80.21%) located at MP2 (N5^{\circ}O8-MP)⁻ or MP3 (N3^{\circ}O5^{\circ}O4-MP)⁻, respectively, while the HOMO-1 and HOMO-2 are the dominant combinations (54.25% and 36.30% versus 38.68% and 52.00%) of the MP2 or the MP3. By further checking Table S4,† the calculated $S_0 \rightarrow S_n$ (n = 1-3) transition absorption wavelengths of [Zn₂(MP)₃(OAc)] are predicted at 450, 416 and 406 nm. For the $S_0 \rightarrow S_1$ transition absorption at 450 nm, a population analysis of HOMO \rightarrow LUMO (91.06%) and HOMO \rightarrow LUMO+2 (5.93%) transitions verifies the dominant ${}^{1}\pi$ - π * transition associated with MP1 and the partial ligand-to-ligand charge transfer (LLCT) feature from the π orbitals of MP1 to the π^* orbitals of MP3. The calculated absorption peak at 417 nm or 407 nm mainly results from the corresponding HOMO \rightarrow LUMO+2 (76.36%) or HOMO \rightarrow LUMO+1 (89.74%) transition, respectively. All the calculated absorptions featuring intraligand $\pi^{-}\pi^{*}$ transitions are in good agreement with the experimental data ($\lambda_{ab} = 404$, 414 and 448 nm) of [**Zn**₂(**MP**)₃(**OAc**)] in solution. In order to definitively elucidate its emissive property, natural transition orbitals (NTOs; Table S5 and Fig. S5[†]) were further obtained from the $S_0 \rightarrow T_1$ excitation with optimized T_1 geometry, whereas the prevailing MP1-centered contribution to both particles (96.96%) and holes (95.21%) is also apparent, and thus nearly 100% (99.5%) of hole \rightarrow particle transitions show that the ${}^{3}\pi$ - π *-dominant transition by the MP1 of

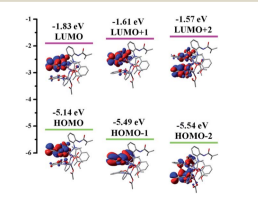


Fig. 3 The HOMO and LUMO patterns for the complex monomer [Zn₂(MP)₃(OAc)] based on its optimized S₀ geometry.

Norm. Absorption (a.u.)

0.2

250 300 $[\mathbf{Zn}_2(\mathbf{MP})_3(\mathbf{OAc})]$ is responsible for its visible phosphorescence. Hence, based on the FMO distribution (also Table S4†) for the T₁ state of $[\mathbf{Zn}_2(\mathbf{MP})_3(\mathbf{OAc})]$, the MO shape also favours the emissive LUMO \rightarrow HOMO (83.56%) transition, and the calculated phosphorescence is located at 593 nm, which is in good agreement with the experimental value of 590 nm (also Fig. S3†) for $[\mathbf{Zn}_2(\mathbf{MP})_3(\mathbf{OAc})]$ in solution at 77 K.

Synthesis, characterization and photophysical property of the grafting-type polymeric film Poly(NVK-*co*-[Zn₂(MP)₃(OAc)])

In consideration of the excellent physical properties²⁵ of semiconducting PVK as a popular matrix, grafting-type polymeric films Poly(NVK-co-[Zn2(MP)3(OAc)]) with different feedings were obtained from the AIBN-initiated copolymerization (also Scheme 1) of NVK and [Zn₂(MP)₃(OAc)]. As a matter of fact, not only does the blue-light PVK matrix act as colour-compensation for the yellow-light-emissive [Zn₂(MP)₃(OAc)], it also functions as an effective energy donor with a large absorption crosssection to transfer energy via Förster mechanism²⁶ to the Zn(II)-complex-acceptor. To address the AIBN-initiated radical copolymerization, all the isolated grafting-type polymers Poly(NVK-co-[Zn₂(MP)₃(OAc)]) were characterized by FT-IR, ¹H NMR and GPC methods. On one hand, besides the combined absorption and proton resonances (also Fig. S1[†]) of the polymerized $[Zn_2(MP)_3(OAc)]$ and PVK, the absence of the original vinyl-characteristic proton signals of [Zn₂(MP)₃(OAc)] or NVK also confirms the covalent-grafting²⁷ of [Zn₂(MP)₃(OAc)] into the PVK backbone. To quantify the actual content of the $Zn(\pi)$ complex segment for each polymer, XPS quantitative analyses (Table S6^{\dagger}) reveal that the Zn(π)-complex-grafted concentration is found to be slightly higher than the initial feeding one, which arises from the loss of oligomeric PVK during isolation. Moreover, GPC results (also Table S6^{\dagger}) of their feeding-related $M_{\rm n}$ values of 17 496-82 512 g mol⁻¹ with a relatively narrow PDI (polydispersity index) range of 1.13-1.29 should result from the AIBN-initiated radical polymerization. Furthermore, the similarity of the amorphous PXRD pattern (Fig. S6[†]) of the representative polymer (200:1) to that of PVK also validates the lowconcentration homogeneous distribution of the complex monomers in the film-forming PVK matrix. TG and DSC analyses (also Fig. S2[†]) for the representative polymer Poly(NVK-co-[Zn₂(MP)₃(OAc)]) (200 : 1) show distinctively enhanced thermal stability (ca. 410 °C T_d , like PVK) over that (290 °C) of [Zn₂(-**MP**)₃(**OAc**)], and desirable T_g up to 202 °C.

The photophysical properties of all the polymers **Poly(NVK***co*-[**Zn**₂(**MP**)₃(**OAc**)]) (100 : 1, 200 : 1, 300 : 1, 400 : 1 or 500 : 1) were investigated in the solid-state film at RT, and summarized in Fig. 4 and S7.† Considering the crossover of the absorption of yellow-light ($\lambda_{em} = 538 \text{ nm}$) [**Zn**₂(**MP**)₃(**OAc**)] with excitation at 345 nm and the excitation at 315 nm of bluelight-emissive ($\lambda_{em} = 418 \text{ nm}$) PVK, also shown in Fig. 2, $\lambda_{ex} =$ 315–345 nm with 10 nm step-sizes should be used as a suitable excitation regime to realize both their simultaneous emissions and the effective PVK-to-[**Zn**₂(**MP**)₃(**OAc**)] energy transfer²⁶ for efficient dichromatic-modulated white light.²⁸ Convincingly, the integrated emissions of PVK-based blue light and

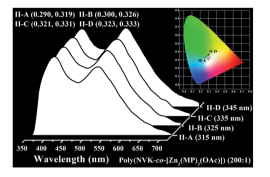


Fig. 4 Normalized emission spectra and corresponding CIE chromatic coordinates (inset) of the polymeric film Poly(NVK-co-[Zn₂(-MP)₃(OAc)]) (200 : 1) upon excitation ($\lambda_{ex} = 315-345$ nm) at RT.

[Zn₂(MP)₃(OAc)]-centered yellow light also depend on the feeding adjustment. Under feeding of 100:1, as shown in Fig. S7(a),† photo-excitation with $\lambda_{ex} = 315-345$ nm gives rise to the dominant yellow light (I-A-D; x = 0.396-0.441, y =0.410-0.458; CCTs = 4553-5337 K and CRIs = 62-66) of [Zn₂(MP)₃(OAc)]. The deficiency of the PVK-based blue light is due to the effective PVK-to-[Zn₂(MP)₃(OAc)] energy transfer.²⁶ However, the residual peak emission at 419 nm confirms the allowance of PVK-based blue light after that preferential energy transfer.²⁶ Further decreasing the $[Zn_2(MP)_3(OAc)]$ grafted content of the polymer Poly(NVK-co-[Zn₂(MP)₃(OAc)]) (200:1), the emissive intensity (Fig. 4) of either the PVKincorporated blue light ($\lambda_{em} = 419$ nm) or the [Zn₂(MP)₃(-**OAc**)]-centered yellow light ($\lambda_{em} = 540 \text{ nm}$) is also λ_{ex} -dependent within the whole 315-345 nm range. Through comparing their combinations, all the resultant dichromatic-integration colour-coordinates (II-A–D (x = 0.290-0.323, y = 0.319-0.333)) are located within the desirable white-light regime, covering a broad range of 350-750 nm emissions with CCTs of 8533-9494 K and CRIs of 74-75. The obtained quantum yield of up to 18.4% for the ideal white-light point II-D (x = 0.323, y = 0.333; CCT of 8533 K and CRI of 75) under excitation at 345 nm is the highest among all reported Zn²⁺-complexdoping systems.^{22,23} The 2.8 ns lifetime of the PVKincorporated blue light ($\lambda_{em} = 419 \text{ nm}$) and the [Zn₂(MP)₃(-**OAc)]**-centered yellow light ($\lambda_{em} = 540 \text{ nm}$) lifetime of 77 ns confirm that the optimal dichromatic white light should have unambiguous fluorescence characteristics. Interestingly, as for the polymer Poly(NVK-co-[Zn₂(MP)₃(OAc)]) (300:1), although all the integrated points III-A-D (x = 0.258-0.265, y =0.296-0.305) also fall into the blue-white-light regime (Fig. S7(b)[†]), their inferior qualities characteristic of supercold white light (CCTs = 10790-12099 K) contribute to the relative inefficiency of [Zn₂(MP)₃(OAc)]-centered yellow light. If more PVK-based blue light is provided for the polymer Poly(NVK-co-[Zn₂(MP)₃(OAc)]) (400:1 or 500:1), the excess PVK renders the integrated colours (Fig. S7(c and d)[†]) significantly deviated from the white-light region, exhibiting bluewhite (IV-A–D: x = 0.226-0.230, y = 0.229-0.249) and blue light (V-A-D: x = 0.205-0.208, y = 0.180-0.282), respectively. It is worth noting that the [Zn₂(MP)₃(OAc)]-centered yellow-light

Chemical Science

species decays with an almost identical lifetime (75-80 ns) to that (77 ns) of Poly(NVK-co-[Zn₂(MP)₃(OAc)]), which is independent of both the feeding and the excitation wavelength, and should be assigned to the excess amount of PVK with the saturated PVK-to-[Zn₂(MP)₃(OAc)] energy transfer.²⁶ Moreover, for each of the polymers Poly(NVK-co-[Zn₂(MP)₃(OAc)]), the facilitated separation of [Zn₂(MP)₃(OAc)] chromophores within the PVK backbone to avoid unexpected self-quenching²⁹ also occurs upon low-concentration grafting. With the aim of knowing the distribution of [Zn₂(MP)₃(OAc)] chromophores within the PVK backbone, AFM images of the polymeric films (40 nm) Poly(NVK-co-[Zn₂(MP)₃(OAc)]) obtained by spin-coating on ITO glass slides were investigated. As shown in Fig. S8,† all the polymeric films with different grafting contents (100:1-500:1) show smooth surfaces with a low root-mean-square (RMS) roughness less than 1.7 nm (1.684 nm (100:1); 1.031 nm (200:1); 1.022 nm (300:1); 1.018 nm (400 : 1); 1.013 nm (500 : 1)), which is also indicative of their homogeneous nature without guest aggregation.²⁹ Meanwhile, due to the smooth surface and the stipulated Zn(II)-grafting content reflected in the SEM-EDS analysis (Fig. S9[†]), it seems that the representative polymer film Poly(NVK-co-[Zn₂(MP)₃(OAc)]) (200 : 1) possesses qualities desirable for high-performance devices.

Device performance of WPLEDs-I-II based on the polymeric film Poly(NVK-*co*-[Zn₂(MP)₃(OAc)]) (200 : 1)

Considering the straightforward and highly efficient white light of the polymeric film Poly(NVK-co-[Zn₂(MP)₃(OAc)]) (200:1), its prototype WPLED-I configured with ITO/PEDOT : PSS (40 nm)/ Poly(NVK-co-[Zn₂(MP)₃(OAc)]) (200 : 1; 40 nm)/TPBi (30 nm)/LiF (1 nm)/Al (100 nm) (Fig. 5(a)) was fabricated using a simple solution-processing technique. The CV result (Fig. S10[†]) of $Poly(NVK-co-[Zn_2(MP)_3(OAc)])$ (200:1) shows its electrochemical HOMO and LUMO levels of -2.69 and -5.21 eV, respectively. PEDOT : PSS (poly(3,4-ethylenedioxythiophene) : poly(styrenesulfonate)) acts as the holeinjecting material, and TPBi (1,3,5-tris(2-N-phenylbenzimidazolyl)benzene) was adopted to facilitate the electrontransport. Due to the critical matching of both the LUMO (-2.69eV) and HOMO (-5.21 eV) levels of the polymer with the LUMO level (-2.70 eV) of TPBi and the HOMO (-5.20 eV) of PEDOT : PSS, respectively, the injected electrons and holes can effectively be recombined within the polymer, and subsequently carrier-trapping and colour-compensation should occur. Just as expected, WPLED-I gives simultaneous emissions (Fig. 5(b)) of PVK-based blue light ($\lambda_{em} = 420 \text{ nm}$) and [Zn₂(MP)₃(OAc)]centered yellow light ($\lambda_{em} = 540$ nm) throughout the whole applied bias voltage range (7.0-18.0 V; Fig. 5(c)). Moreover, after the turn-on voltage (V_{on} at 1 cd m⁻²) of 7.0 V, their comparative combination exhibits stable white light (CIE coordinates of x =0.297-0.300, y = 0.332-0.339; CCTs of 7105-7346 K and CRIs of 81-82) within the 7.0-12.0 V range. As shown in Fig. 5(f), with an increase in the applied bias voltage, both the luminance (L, cd m^{-2}) and the current density (J, mA cm⁻²) monotonously increase, and $L^{\text{Max}} = 44.2$ cd m⁻² is achieved at 12.0 V with

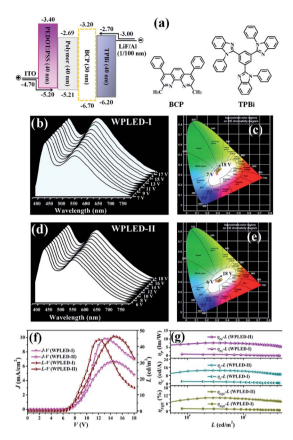


Fig. 5 (a) Device structures for the WPLED-I–II with energy level diagrams; (b) electroluminescence spectra and (c) CIE chromaticity coordinates for the WPLED-I; (d) electroluminescence spectra and (e) CIE chromaticity coordinates for the WPLED-II; (f) J-V-L and (g) η_c-L , η_p-L and $\eta_{EOE}-L$ for the WPLED-I-II.

a current density of 9.8 mA cm⁻². Under the stable premise (7.0– 12.0 V) for the WPLED-I, with an increase in luminance or applied bias voltage, as shown in Fig. 5(g), all the efficiencies $(\eta_c, \eta_p \text{ and } \eta_{EQE})$ instantly decrease with $\eta_c^{Max} = 2.2$ cd A⁻¹, $\eta_p^{Max} = 1.0 \text{ lm W}^{-1}$ and $\eta_{EQE}^{Max} = 1.7\%$. To our dismay, upon using an applied bias voltage up to 12.0 V, both the η_c and η_p efficiencies decrease to 0.5 cd A⁻¹ and 0.15 lm W⁻¹, respectively, and severe (*ca*. 73%) efficiency roll-off ($\eta_{EQE} = 0.45\%$) also takes place. Especially when the applied bias voltage is further increased (>12.0 V), the significantly inferior performance caused by the carrier imbalance³⁰ should be attributed to the ageing of the WPLED-I.

Based on optimization with the additional BCP layer for carrier blocking,³¹ the BCP-incorporated (30 nm) WPLED-II was fabricated. Convincingly, because charge carriers can be confined within the broadened recombination zone, the WPLED-II exhibits improved electroluminescence properties as desired. First, besides a lower $V_{\rm on}$ of 6.0 V, the dichromatic white light (Fig. 5(d and e); CIE coordinates of x = 0.325-0.329, y = 0.300-0.310; CCTs of 7050-7318 K and CRIs of 82-84) can be stable within an extended applied bias voltage range of 6.0-15.0 V, and $L^{\rm Max}$ (46.2 cd m⁻²) is slightly increased with a lower *J* of 6.7 mA cm⁻² at 15.0 V. More importantly, its maximum

efficiencies (η_c^{Max} = 13.0 cd $A^{-1}\text{,}~\eta_p^{Max}$ = 6.1 lm W^{-1} and $\eta_{\rm EOE}^{\rm Max} = 9.2\%$) reformed almost 5–6 times more than those of the WPLED-I and are the best (Table S7[†]) among reported Zn(II)complex-based WOLEDs¹⁶⁻²⁰/WPLEDs,^{22,23} to our knowledge. Furthermore, within the stable illuminating range, the high efficiencies ($\eta_c = 9.1 \text{ cd } A^{-1}$, $\eta_p = 4.4 \text{ lm } W^{-1}$ and $\eta_{EQE} = 6.8\%$) are maintained, with distinctively weaker efficiency roll-off (ca. 25%) than that (ca. 73%) of the WPLED-I. The WPLED-II is distinctively superior to the ubiquitous WOLEDs/WPLEDs (>40%) based on Ir(III)-5 or Pt(II)-complexes6 with large phosphorescence-decayed lifetimes. Inspiringly, the recordhigh performance (high efficiencies and weak efficiency rolloff) of the WPLED-II renders the single Zn(u)-complex-grafted polymer a new platform for cost-effective and large-area flexible WPLEDs, and could be further improved through material modification and device optimization.

Conclusions

In summary, for the first time, tris-vinyl-functionalized yellowlight complex monomer $[Zn_2(MP)_3(OAc)]$ was copolymerized with NVK to obtain the single grafted-type polymers Poly(NVKco-[Zn2(MP)3(OAc)]) for WPLEDs. Especially, using Poly(NVK-co-[Zn₂(MP)₃(OAc)]) (200:1) with straightforward white light as the emitting layer, a reliable WPLED-II with optimized carrier balance gives record-high device performance (both high efficiencies (13.0 cd A⁻¹ of η_c^{Max} , 6.1 lm W⁻¹ of η_p^{Max} and 9.2% of $\eta_{\rm EOE}^{\rm Max}$) and weak (ca. 25%) efficiency roll-off) compared with organo-Zn²⁺-based previous WOLEDs/WPLEDs. This outstanding result renders the single Zn(II)-complex-grafting polymer a new platform for cost-effective and large-area flexible WPLEDs for potential full-colour flat displays.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

Prof. X. Q. Lü thanks the National Natural Science Foundation (21373160, 21173165), the Program for New Century Excellent Talents in University from the Ministry of Education of China (NCET-10-0936), the State Key Laboratory of Structure Chemistry, and the Wisteria Scientific Research Cooperation Special Project of Northwest University in P. R. of China for the support of this work. Prof. H. S. He thanks the National Science Foundation (1507871) in the United States for the support of this work. Prof. L. Liu thanks the National Natural Science Foundation of China (21671061) and the Application Foundation Frontier Special Project by Wuhan Science and Technology Bureau (2019010701011414) for the support of this work. Prof. W.-Y. Wong thanks the Hong Kong Research Grants Council (PolyU153062/18P), the Hong Kong Polytechnic University (1-ZE1C) and the Endowed Professorship in Energy from Ms. Clarea Au (847S) for the financial support.

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