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# High-performance solution-processed red hyperfluorescent OLEDs based on cibalackrot†

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Hyperfluorescent organic light-emitting diodes (OLEDs) have allowed remarkable device performances to be achieved using fluorescent emitters. Superior device performance has been realised using thermally evaporated emissive layers. However, for future large-scale commercialisation, it is essential to obtain similar device performances using low-cost solution-processing techniques. In the case of hyperfluorescent OLEDs, there remains a delicate interplay of molecular interactions and spacing between the three active components: a host, a thermally activated delayed fluorescent (TADF) assistant host, and a fluorescent emitter. Dispersion of the materials is a dominating factor towards the device efficiencies, making efficient solution-processed devices all the more difficult to achieve. Herein, we have demonstrated solution-processed hyperfluorescent devices with an external quantum efficiency (EQE) of 15.3% using cibalackrot as the fluorescent emitter and 4CzIPN-tBu as the TADF assistant host in CBP. By studying the use of either 4CzIPN or 4CzIPN-<sup>t</sup>Bu as the TADF assistant host in both ternary and host-free binary blends, we found that the addition of tert-butyl groups to the TADF material made a significant contribution to the device performance. These sterically hindered groups effectively reduced losses caused by triplet diffusion between the TADF assistant host and the fluorescent emitter by spatially separating adjacent molecules and making a concurrent frontier molecular orbital (FMO) less likely.

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## Introduction

Since the introduction of thermally activated delayed fluorescence (TADF), organic light-emitting diodes (OLEDs) have exhibited significant advancement in performance, achieving emission along the entirety of the visible spectrum. The use of TADF emitters in OLED devices allows efficient harvesting of the otherwise non-emissive triplet excitons which make up 75% of the generated excitons according to the spin-statistics rule of a 1:3 ratio for singlets and triplets. This is achieved by efficient up-conversion of triplets from the triplet state  $(T_1)$  to

the singlet state (S<sub>1</sub>) via a process called reverse intersystem crossing (RISC), allowing these devices to have up to 100% of the generated excitons as emissive species. Efficient RISC is achieved through reduction of the singlet-triplet energy gap  $(\Delta E_{\rm ST})$ , which can be accomplished by the separation of the highest occupied molecular orbital (HOMO) and the lowest unoccupied molecular orbital (LUMO) on the respective donor and acceptor parts of the emissive material. This orbital separation leads to charge-transfer (CT) character, and it is known that the radiative decay from TADF materials is spectrally broad, limiting the scope of TADF-OLEDs in display applications. Furthermore, due to the RISC process, TADF materials show considerably longer emission lifetimes compared to conventional fluorescence emission. As such, TADF devices commonly experience a strong efficiency roll-off during operation at high current densities due to the dominating non-radiative Dexter-type (DET) losses such as singlet-triplet annihilation (STA) and triplet-triplet annihilation (TTA).<sup>2,3</sup>

In 2014, Nakanotani et al. demonstrated the utilisation of TADF emitters as assistant hosts for fluorescent end-emitters to give hyperfluorescence.4 In this design, the TADF assistant host has upconversion of triplet excitons to singlet excitons via the RISC process, and these singlet excitons can be transferred to

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the S<sub>1</sub> of the lower energy fluorescent end-emitter via Förster resonance energy transfer (FRET), allowing conventional fluorescent emitters to achieve up to 100% internal quantum efficiency (IQE). This system effectively reduces excited-state lifetimes and produces a spectrally narrowed emission (reduction of the full-width at half maximum (FWHM)), as the radiative decay occurs from the fluorescent end-emitter rather than the TADF material. As a result, hyperfluorescent OLEDs show improved operation stability with the device performance comparable to conventional fluorescent or TADF devices, 5,6 clearly representing an exciting new opportunity for OLED device design.<sup>7-13</sup> Unfortunately, like TADF devices, hyperfluorescent OLEDs continue to suffer from DET losses and efficiency roll-off due to triplet diffusion between the TADF and fluorescent end-emitters due to close proximity to each other in the film. 11,14,15 This can be suppressed by using relatively low doping concentrations of the fluorescent endemitter (< 2%), and dispersing the TADF assistant in a high energy host (e.g., mCP, CBP) to create a ternary blend. Recently, Lemmer and Howard et al. proposed a plausible approach for suppressing triplet diffusion by the introduction of bulky peripheral functional groups to increase steric effects. 16 Namely, in theory, by increasing the intermolecular distances, the overlap of concurrent molecular orbitals (i.e., HOMO  $\rightarrow$ HOMO, and LUMO  $\rightarrow$  LUMO for TADF  $\rightarrow$  fluorescent emitter) can be minimised.

For the fabrication of OLEDs, thermal deposition is a common technique that permits the assembly of complicated device architectures and facilitates the inclusion of charge injection and charge blocking layers to help improve the charge balance within the device. Additionally, the thermal deposition of materials consistently produces a superior film quality, essential for achieving improved light-out coupling and outstanding device performances.<sup>17</sup> This technique is perfect for small-area devices and research purposes. However, for future large-scale applications, it is essential to develop high-efficiency devices using simple and low-cost fabrication methods such as solution processing.<sup>18</sup> Unfortunately, solution deposition is restrictive towards the device design, as there are limited choices of interlayer materials that are solution compatible (e.g., the issue of the dissolution of bottom layer materials while spin-coating on the top). Furthermore, solution-processing often generates an inferior film quality, where the resulting non-uniform morphology and molecular orientation lower the light out-coupling efficiencies, thus decreasing the external quantum efficiencies (EQEs).19 In the case of hyperfluorescent OLEDs, there remains a delicate interplay of molecular interactions and spacing between the three active components: the host, the TADF-assistant host, and the fluorescent end-emitter. Dispersion of the materials is a dominating factor towards hyperfluorescent device efficiencies, making high-efficiency solution-processed devices more challenging to achieve. Unsurprisingly, to date, there has been few reports of solutionprocessed hyperfluorescent OLEDs. A study by Chen and Cai et al. demonstrated devices with EQEs up to 8% using novel TADF emitters as the assistant hosts and DBP as the red

fluorescent end-emitter.20 Similar EQEs of 8% were also demonstrated by Shukla et al. using CzDBA as the TADF assistant host and DT-DPP as the fluorescent end-emitter.21

We recently reported a new family of solution-processable semiconductors based on cibalackrot, establishing their use as efficient organic laser dyes.<sup>22</sup> The attachment of solubilising peripheral groups to the cibalackrot core endows it with high solubility in a variety of common organic solvents (e.g., toluene, chloroform, etc.) and forms high-quality thin films.<sup>22</sup> These soluble cibalackrot materials were found to possess high thermal stability and exhibit reversible redox profiles. The rigid and flat structure of the cibalackrot core also leads to strong absorption and emission properties with low nonradiative decay rates to give a high solution photoluminescence quantum yield (PLOY) (96%) and a narrow red emission (FWHM = 37 nm), making them ideal candidates for further study in emissive optoelectronic applications, in particular as fluorescent end-emitters in solution-processed hyperfluorescent OLEDs. However, like its non-emissive parent indigo, the use of cibalackrot materials in the literature is limited to their use in organic photodetectors (OPDs),<sup>23</sup> organic solar cells (OSCs), 24,25 organic field-effect transistors (OFETs), 26 and electrochromic devices, 27 and as such, electroluminescence (EL) properties have yet to be realised.

Herein, we demonstrate solution-processed red hyperfluorescent OLEDs with an EQE of ≈15 at% 100 cd m<sup>-2</sup> The materials used in this study are summarised in Fig. 1. Cibalackrot B (referred to as cibalackrot throughout in this work)<sup>22</sup> was chosen as our fluorescent end-emitter, and two similar TADF materials, 4CzIPN and 4CzIPN-<sup>t</sup>Bu, <sup>28,29</sup> were compared as the assistant dopants. CBP was used as the host material in ternary blends. By comparing 4CzIPN and 4CzIPN-tBu in both ternary (CBP host) and binary (host-free) blends, we attempt to develop a better understanding of how increased molecular size and steric interactions (resulting from the additional tert-butyl surface groups) may influence the FRET efficiency and the DET losses in OLED devices.

## Results and discussion

### Energy transfer study of the host and TADF dopants

To better understand the energy level interactions and potential for FRET and DET between CBP, 4CzIPN, 4CzIPN-<sup>t</sup>Bu, and cibalackrot, the S<sub>1</sub> and T<sub>1</sub> energy levels were determined experimentally in solution (toluene) and the solid state (neat thin films). The calculated energy levels are summarised in Tables S1 and S2 (ESI†). Fig. S1 (ESI†) shows the absorption, fluorescence, and phosphorescence (77 K) of the materials in toluene solution as well as in neat films. Only the solution measurements for cibalackrot are shown as they are analogous to the solid-state properties when dispersed at low concentrations (below 2 mol%) throughout a host matrix. The S<sub>1</sub> of cibalackrot in solution was found to agree well with the literature;<sup>30</sup> however, while the PLQY is close to unity, the T<sub>1</sub> yield was relatively weak and could not be detected. It is

Fig. 1 Molecular structures of the materials used in this study.

expected that cibalackrot will have a similarly very low T<sub>1</sub> energy level ( $\approx 1.27$  eV) with phosphorescence at  $\approx 970$  nm, as reported for the unfunctionalised parental core.30 The S1 and T<sub>1</sub> energy levels of 4CzIPN and 4CzIPN-<sup>t</sup>Bu were comparable, and both were found to have an  $S_1$ - $T_1$  energy gap  $(\Delta E_{ST})$  of 0.06 eV in neat films. In solution, 4CzIPN-tBu is slightly redshifted relative to 4CzIPN as the additional tert-butyl groups improve the donor strength of the carbazole units, subtly altering the donor-acceptor interaction (Fig. S1, ESI†). However, it is important to note that the emission of 4CzIPN shows a strong bathochromic shift from solution to neat films ( $\lambda_{PL}$  = 510 nm  $\rightarrow$  568 nm,  $\Delta \lambda_{PL}$  = 58 nm), while the resulted shift for 4CzIPN-<sup>t</sup>Bu is comparably smaller ( $\lambda_{PL} = 525 \text{ nm} \rightarrow 549 \text{ nm}$ ,  $\Delta \lambda_{PL}$  = 24 nm). This suggests that 4CzIPN-<sup>t</sup>Bu experiences weaker intermolecular interactions in the solid-state compared to 4CzIPN due to the bulkier peripheral tert-butyl groups

increasing steric effects as expected. As shown in Fig. 2a, the energy levels of CBP, 4CzIPN, and 4CzIPN-tBu have a good alignment for energy transfer to the S<sub>1</sub> of cibalackrot. There is also a good spectral overlap between the emission from CBP and the absorption of 4CzIPN/4CzIPN-tBu, as well as the emission from 4CzIPN/4CzIPN-<sup>t</sup>Bu and the absorption of cibalackrot in solution (as shown in Fig. 2b). This spectral overlap implies good FRET under optical excitation for the chosen system. We further investigated the FRET exchange radius  $(R_0)$ , and the rate of FRET  $(k_{\text{FRET}})$  as calculated by eqn (1) and (2) below:

$$R_0 = \frac{9000 \ln \kappa^2 \phi_{\rm PL}}{128\pi^5 n_0^4 N_{\rm A}} \int_0^\infty F_{\rm D}(\lambda) \varepsilon_{\rm A}(\lambda) \lambda^4 d\lambda \tag{1}$$

$$k_{\text{FRET}} = \left(\frac{1}{\tau_{\text{D}}}\right) \left(\frac{R_0}{r}\right)^6 \tag{2}$$

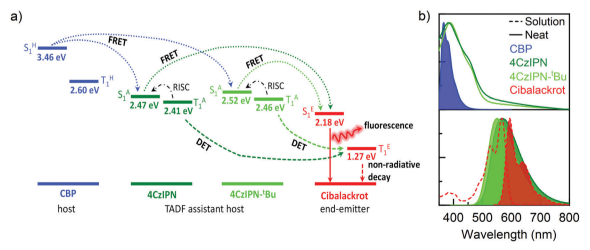


Fig. 2 (a) Schematic illustration of energy levels and energy transfer processes in the ternary blend. The energy levels of CBP, 4CzIPN, and 4CzIPN-¹Bu were determined from neat films, and cibalackrot from toluene solution. Dotted arrows represent FRET, dashed arrows represent DET, dot-dash arrows represent RISC, and solid arrows represent radiative emission. (b) Absorption and photoluminescence (PL) (filled) of the CBP host in neat films (solid), and TADF assistant hosts, showing a good overlap of CBP PL with 4CzIPN/4CzIPN-<sup>t</sup>Bu, and 4CzIPN/4CzIPN-<sup>t</sup>Bu PL and cibalackrot absorption (dashed).

where  $\lambda$  is the wavelength,  $F_D(\lambda)$  is the fluorescence distribution of the donor,  $\varepsilon_A(\lambda)$  is the molar extinction coefficient of the acceptor,  $\phi_{\rm PL}$  is the PLQY of the donor,  $\kappa$  is the orientation factor,  $n_0$  is the refractive index,  $N_A$  is the Avogadro number, and  $\tau_{\rm D}$  is the excited-state lifetime of the donor.

The  $R_0$  values were found to be 3.4 nm for CBP  $\rightarrow$  cibalackrot, 4.7 nm for 4CzIPN  $\rightarrow$  cibalackrot, and 4.9 nm for 4CzIPN- ${}^{t}$ Bu  $\rightarrow$ cibalackrot. Due to the larger exchange radius, energy transfer is expected to occur in a cascading manner from CBP -> 4CzIPN/  $4\text{CzIPN-}^t\text{Bu} \rightarrow \text{cibalackrot.}$  Interestingly, the estimated  $R_0$  for 4CzIPN-<sup>t</sup>Bu → cibalackrot is slightly higher than that for 4CzIPN → cibalackrot, resulting from a better overlap between the PL of 4CzIPN-<sup>t</sup>Bu and the absorption of cibalackrot (integral overlaps of  $1.45 \times 10^{15}$  for 4CzIPN-<sup>t</sup>Bu, and  $1.11 \times 10^{15}$  for 4CzIPN). Due to the non-radiative nature of triplet excitons in the fluorescent compounds, it is challenging to estimate the DET exchange radius, although previous studies have found it to be within a domain of 1-2 nm.<sup>21</sup>

As previously discussed, we aimed to study the  $k_{\text{FRET}}$  and DET loss efficiencies ( $k_{DET}$ ) on device performances using two systems. In the first system, we used ternary blends of cibalackrot in TADF: CBP (≈30:70 mol%) with varying mol% of cibalackrot. In order to optimise the device performance,<sup>31</sup> and realise efficient energy transfer between the CBP host and TADF-assistant dopant, high concentrations of TADF (i.e. 50:50 wt%, equivalent to a 30:70 molar ratio to CBP) have been employed in this work. By implementing a ternary blend with low doping concetrations, we specifically aimed to increase the intermolecular spacing (r) between the TADF and cibalackrot molecules to assist the supression of short-range DET, and simultaneously inhibit concentration quenching. The increased r is expected to give a reduction of  $k_{DET}$  but with little impact on  $k_{\text{FRET}}$  as the long-range FRET processes can still efficiently occur in the cascading manner as described previously. In the second system, we use a binary blend (host-free), composed of a neat TADF material and cibalackrot of varying mol%. By removing the CBP host, we attempt to increase  $k_{\text{FRET}}$ through promoting interactions between the adjacent TADF and Cibalackrot molecules as  $k_{\text{FRET}} \propto 1/r$  (eqn (2)). However, while  $k_{\text{FRET}}$  should increase with decreasing r, the removal of CBP will also lead to a simultaneous increase in  $k_{DET}$  as a result of the increased concentration of the TADF material and its long triplet lifetime. Therefore, additional consideration of triplet diffusion and the resulting DET loss efficiencies was explored by comparing the 4CzIPN TADF material with its more sterically encumbered derivative, 4CzIPN-<sup>t</sup>Bu (Fig. 1).

#### Photophysical properties

Photophysical properties of ternary blends. The photophysical properties of cibalackrot at varying mol% were studied in ternary blends, using a ≈30:70 mol% blend ratio of either 4CzIPN-<sup>t</sup>Bu:CBP or 4CzIPN:CBP. Fig. S2a (ESI†) shows the normalised absorption and PL of 0.5-1.5 mol% cibalackrot doped in 4CzIPN-tBu:CBP, and 1.5 mol% cibalackrot/ 4CzIPN:CBP blend films (spin-coated from 0.5 wt% chloroform solution). As predicted, the absorption of the tertiary blends

was dominated by both CBP and the TADF assistant dopant (4CzIPN or 4CzIPN-tBu) with a minimal contribution from cibalackrot. In contrast, PL was dominated by emission from cibalackrot, with little contribution from the other components. This implies that FRET is indeed occurring efficiently from CBP → 4CzIPN/4CzIPN-<sup>t</sup>Bu → cibalackrot. The excitedstate lifetimes of these blends were estimated from TCSPC prompt and delayed fluorescence decays as shown in Fig. S3a and c (ESI†). The PLQYs along with prompt  $(\tau_p)$  and delayed  $(\tau_d)$ lifetimes of the various blends are further summarised in Table S3 (ESI†). Increasing mol% of cibalackrot resulted in a decreased emission contribution from the TADF-dopant as FRET between the two materials increased. This is noted in the PL by a significant reduction of the emission peak at  $\approx$  540 nm, and the decreasing values of  $\tau_p$  and  $\tau_d$ , as there is progressively less delayed fluorescence from 4CzIPN-<sup>t</sup>Bu. Additionally, the relative  $\tau_p$  and  $\tau_d$  of the ternary blends were found to be much faster than non-doped TADF:host blends, especially  $\tau_{\rm d}$ , which was reduced by a factor of 10, another indication that FRET is efficiently taking place from  $4\text{CzIPN}^{-t}\text{Bu} \rightarrow \text{cibalackrot}$ . The PLQYs of these blends showed a slight reduction with increasing cibalackrot mol% but were generally comparable varying between 74% and 79%. The PL spectra,  $\tau_p$ , and  $\tau_d$  were comparable between 1.5 mol\% 4CzIPN-\textit{t}Bu:CBP and 4CzIPN:CBP blend films (Fig. S2a and S3a, c, ESI†). The PLQY of 1.5 mol% cibalackrot blend films was slightly higher for 4CzIPN-<sup>t</sup>Bu:CBP (74%) compared with 4CzIPN:CBP (69%). This is in agreement with the prediction resulting from the comparison of  $R_0$  where FRET should be more efficient between 4CzIPN-<sup>t</sup>Bu → cibalackrot resulting in higher PLQYs.

Photophysical properties of binary blends. The photophysical properties of binary blends were studied using 1-2 mol% cibalackrot in either 4CzIPN-<sup>t</sup>Bu or 4CzIPN. Fig. S2b (ESI†) shows the normalised absorption and PL of blend films spin-coated from 0.5 wt% chloroform solutions. Similar to the tertiary blends, the absorption was dominated by 4CzIPN/4CzIPN-tBu, with minor contribution from cibalackrot. Again, PL was dominated by emission from cibalackrot, with little contribution from 4CzIPN/ 4CzIPN-<sup>t</sup>Bu, implying a similar cascading FRET from 4CzIPN/ 4CzIPN-<sup>t</sup>Bu → cibalackrot. The excited-state lifetimes of these blends were estimated from TCSPC prompt and delayed fluorescence decays as shown in Fig. S3b and d (ESI†). Similarly, the relative  $\tau_{\rm p}$  and  $\tau_{\rm d}$  were found to be faster than 4CzIPN and 4CzIPN-<sup>t</sup>Bu neat films. In line with observations from the ternary blends and previous predictions resulting from lower  $R_0$ , the binary system also showed reduced PLQYs for blends containing 4CzIPN (47-58%) in comparison with those with 4CzIPN-tBu (65-70%). Additionally, resulting from the substantial increase of TADF mol% in the binary films, the increased intermolecular interactions between TADF - TADF may lead to non-radiative losses and further reduction of PLQYs in comparison to the CBP ternary blends.

### **Device characteristics**

To investigate the EL properties of these blends, OLEDs were fabricated using a solution-processed emissive layer (EML) with

the following device architecture: ITO (100 nm)/PEDOT:PSS (30 nm)/ EML ( $\approx$ 35 nm)/TPBi (65 nm)/Liq (2 nm)/Al (100 nm), where ITO, PEDOT:PSS, TPBi and Liq are indium tin oxide, poly(3,4-ethylenedioxythiophene):poly(styrenesulfonate), 2,2',2"-(1,3,5-benzinetriyl)-tris(1-phenyl-1H-benzimidazole) and lithium 8quinolate, respectively. EML is the emissive layer, containing cibalackrot in TADF:host blends as summarised in Table 1. Fig. 3a shows the corresponding energy level diagram for the device architecture.

Device characteristics using ternary blend EMLs. The current density voltage-luminance (J-V-L) and EQE-current density curves for the OLEDs using ternary blends are shown in Fig. 3b and d. Device performances are summarised in Table 1, and the EL spectra at maximum EQEs (EQEmax) are shown in Fig. 3c. The highest EQEs were achieved using 0.5 mol% cibalackrot in 4CzIPN-<sup>t</sup>Bu:CBP (≈29.5:70 mol%) as the EML, producing a maximum EQE of 15.3% and an EQE of 14.9% at 100 cd m<sup>-2</sup>. The achieved device performances are comparable with thermally evaporated devices (Fig. S4, ESI†) and are higher than previously reported solution processed devices.21,22 When the concentration of cibalackrot was increased from 0.5 to 1.5 mol% in 4CzIPN-tBu:CBP devices, EQE<sub>max</sub> decreases from 15.3% to 8.7%, although other device characteristics such as current densities, brightness, and light turn on voltage  $(V_{on})$  are improved. By comparison of the EL spectra in Fig. 3c, less emission from 4CzIPN-<sup>t</sup>Bu was observed while increasing the cibalackrot dopant concentrations, similar to those observed in the PL spectra (Fig. S2a, ESI†). This is because by increasing the doping concentrations, the effective intermolecular spacing (r) is reduced, leading to more efficient  $k_{\text{FRET}}$  from 4CzIPN-<sup>t</sup>Bu  $\rightarrow$  cibalackrot and hence more emission from cibalackrot instead of 4CzIPN-tBu.

However, as the relative EL contribution from 4CzIPN-<sup>t</sup>Bu is not insignificant, we approximated the relative emission contributions from 4CzIPN-<sup>t</sup>Bu at EQE<sub>max</sub> by comparing the total integrated area of the EL curve to an overlapping area of emission from 4CzIPN-tBu; this gave the relative EL contributions from 4CzIPN-<sup>t</sup>Bu to be 28, 18, and 10% for the respective 0.5, 1.0, and 1.5 mol% based devices (Fig. 3c). By applying this, we further estimated the EQE contribution from cibalackrot to be  $\approx 11.0\%$ , 9.1% and 7.8% for the 0.5, 1.0, and 1.5 mol% blend devices, respectively, agreeing with the decreasing trend observed previously. As the PLQYs of these two blends were similar, the achieved EQEs are far below what may theoretically be achieved (a max EQE of about 15% based on the film PLQY

of 75%). Hence, the considerable reduction in EQEs is likely due to increased losses from DET as short-range interactions between 4CzIPN-<sup>t</sup>Bu → cibalackrot increase with the higher doping concentrations of cibalackrot. Competing  $k_{\text{DET}}$  may also contribute to a reduction of the  $k_{\text{FRET}}$  efficiency under electrical injections. Conversely, the reduction in  $V_{\rm on}$  and the increasing current densities may be due to better ambipolar character within the blend films. Consequently, direct carrier recombination and charge trapping on cibalackrot can also occur more rapidly, leading to non-emissive triplet states which reduce the overall electroluminescence from the device upon decaying to the ground state.

Comparison of 4CzIPN-<sup>t</sup>Bu:CBP and 4CzIPN:CBP is observed in OLED devices containing 1.5 mol% cibalackrot. As the TADF materials are dispersed in CBP, short-range interactions such as DET and aggregate quenching effects are expected to be suppressed, and both blends should show similar device performances. This was indeed observed, and both devices show a relatively similar current density of > 100 mA cm<sup>-2</sup>, and a light turn on voltage,  $V_{\rm on}$ , of 4.4 V. The EL spectra at EQE<sub>max</sub> were also found to be comparable (Fig. 3c). However, while there is significant quenching of electroluminescence in both devices, the losses from DET are more substantial in the 4CzIPN:CBP blend. Similar to the thin film photophysical properties, the observed EQEs at 100 cd m<sup>-2</sup> are much lower in 4CzIPN:CBP at 5.6%, in comparison with 4CzIPN-<sup>t</sup>Bu:CBP at 8.0%. As there is a significant difference between the electroluminescent performances, it is clear that the attachment of tert-butyl surface groups plays an essential role in reducing the overall emission loss from the device.

Device characteristics using binary blend EMLs. The J-V-L and EQE-current density curves for the OLEDs using binary blends are shown in Fig. 4a and b. Device performances are summarised in Table 2, and the EL spectra at maximum EQEs are shown in Fig. 4c. As predicted, the host-free binary blend devices showed comparatively lower performances than the ternary blends, with stronger roll-off and luminescent quenching. The EL spectra of the binary host blends (Fig. 4c), matched the trends observed during PL studies. Interestingly, the contribution of electroluminescence from TADF-dopants varied widely between different doping concentrations and also between 4CzIPN and 4CzIPN-Bu. The binary blend devices showed comparatively higher Von and lower current densities, due to the less alignment of energy levels with TPBi by the removal of CBP (see Fig. 3a). Interestingly, the 4CzIPN-<sup>t</sup>Bu

Table 1 Summary of ternary blend device characteristics

Blend ratio (mol%)						EQE (%)		
Cibalackrot	TADF		Host	$V_{\mathrm{on}}\left(\mathbf{V}\right)$	Max brightness (cd m <sup>-2</sup> )	Max	$100~cd~m^{-2}$	1000 cd m <sup>-2</sup>
0.5	29.5	4CzIPN- <sup>t</sup> Bu	CBP	5.0	4595	15.3	14.9	8.4
1.0	29.0	4CzIPN- <sup>t</sup> Bu	CBP	4.8	5782	11.2	11.0	7.2
1.5	28.5	4CzIPN- <sup>t</sup> Bu	CBP	4.4	5277	8.7	8.0	4.5
1.5	28.5	4CzIPN	CBP	4.4	4680	5.6	5.6	4.4

 $V_{\rm on}$  is the turn on voltage at 1 cd cm<sup>-2</sup>.

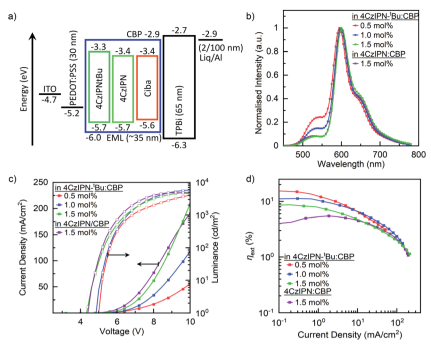


Fig. 3 Device characteristics for OLEDs containing cibalackrot of varying mol% when blended in TADF: CBP ( $\approx 30:70$  mol%). (a) Energy level diagram of the hyperfluorescent device architecture used in this study, (b) EL at EQE<sub>max</sub>, (c) current density-voltage-luminance (JVL) characteristics, and (d) EQE versus the current density.

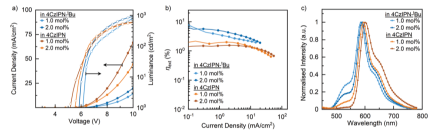


Fig. 4 (a) Comparison of current density-voltage-luminance (J-V-L) characteristics, (b) comparison of EQEs versus current densities, and (c) comparison of EL at EQE<sub>max</sub> for OLED with binary blend EML.

Table 2 Summary of binary blend device characteristics

Blend ratio (mo	ol%)				EQE (%)		
Cibalackrot	TADF		$V_{\mathrm{on}}\left(\mathbf{V}\right)$	Max brightness (cd m <sup>-2</sup> )	Max	$100~{\rm cd}~{\rm m}^{-2}$	1000 cd m <sup>-2</sup>
1.0	99.0	4CzIPN- <sup>t</sup> Bu	6.2	1462	7.7	4.0	2.1
2.0	98.0	4CzIPN- <sup>t</sup> Bu	6.0	1343	5.7	5.0	1.9
1.0	99.0	4CzIPN	5.6	615	2.5	2.2	_
2.0	98.0	4CzIPN	5.2	606	1.5	1.5	_

devices show lower device efficiencies with a higher  $V_{\rm on}$  of < 6.2 V, and a lower current density of <30 mA cm<sup>-2</sup>, in comparison with the 4CzIPN devices (a  $V_{\rm on}$  of < 5.6 V, and a current density of >100 mA cm<sup>-2</sup>), which is likely due to the less close packing of the 4CzIPN-<sup>t</sup>Bu TADF molecules arising from the additional tert-butyl groups. Despite this, the 4CzIPN-<sup>t</sup>Bu devices possess superior EQEs at 100 cd m<sup>-2</sup> of 5%, in contrast to the 4CzIPN devices with EQEs of <2.2%. Like the ternary blend system, this reduction in electroluminescence is likely caused by losses through DET, and the steric effect from the additional tert-butyl groups seems more influential in this second binary system. There are two benefits

related to the attachment of these steric groups. 16 First, the steric groups form a protective shell surrounding the 4CzIPN active core effectively reducing the triplet transport between 4CzIPN-<sup>t</sup>Bu → cibalackrot. Of course, this "shell" is relatively thin (<0.1 nm)<sup>30</sup> and unlikely to result in such a significant effect on the overall performance, unlike the previously reported simulations conducted by Lemmer and Howard et al., which used a protective shell radius of 1.75 nm. <sup>17</sup> For similar reasons, the long-range FRET is unlikely to be affected by the additional tert-butyl groups. Secondly, to suppress efficient triplet energy transfer, less overlap of both the HOMOs and LUMOs between two adjacent molecules is critical.<sup>17</sup> It is expected that the bulkier geometry of 4CzIPN-<sup>t</sup>Bu reduces the intermolecular interaction in the solid-state compared with 4CzIPN. Hence, it is more likely that 4CzIPN-<sup>t</sup>Bu possesses higher electroluminescence performance as a result of the reduced triplet transfer.

As noted earlier, the charge carrier properties in the binary blends seem better for the 4CzIPN devices. Interestingly, the 4CzIPN device efficiencies reflect more fluorescent-type emission with a brightness of <600 cd m<sup>-2</sup> and an EQE of <2.5%, so perhaps all the excitons are funnelled straight to cibalackrot, without efficient RISC taking place. The triplet transport within the emissive layer would also be increased with a better frontier molecular orbital (FMO) overlap, and close packing between 4CzIPN → cibalackrot, making triplet excitons more likely to be lost through non-radiative DET, which may explain the stronger roll-off effect and reduced electroluminescence performances.

## Conclusion

In summary, we have demonstrated solution-processed red hyperfluorescent devices with an unparalleled EQE of 15.3% using a cibalackrot:4CzIPN-<sup>t</sup>Bu:CBP blended EML. By studying the use of either 4CzIPN or 4CzIPN-<sup>t</sup>Bu as a TADF assistant dopant in both ternary and host-free binary blends, we found that the addition of tert-butyl surface groups gave a significant influence towards the device performance. These steric groups effectively suppressed triplet diffusion between the adjacent molecules by reducing the concurrent overlap of the HOMOs and LUMOs to reduce losses through DET. Overall, the device performances are comparable to thermally evaporated devices, and significantly higher than previously reported solution processed red hyperfluorescent OLEDs. These results show that cibalackrot is a promising new family for electroluminescence semiconductors.

# **Experimental**

#### General information

The organic compounds 4CzIPN-tBu and cibalackrot were synthesised according to previously reported procedures. 22,32

### Photophysical measurements

Organic thin-films for optical measurements were fabricated on clean quartz by spin-coating. The substrates were cleaned with acetone and isopropanol and then treated with UV/ozone to remove the adsorbed organic species before deposition. Electronic absorption spectra were recorded using a PerkinElmer Lambda 950-PKA UV-vis spectrophotometer. Fluorescence spectra were recorded using a JASCO FP-8600 fluorometer. Absolute fluorescence quantum yields were measured using a Quantaurus-QY C11347-01 calibrated integrating-sphere system. Transient photoluminescence decay was recorded using a Hamamatsu Photonics Quantaurus-Tau C11367-03 system.

### Device fabrication and measurements

Glass substrates with a pre-patterned, 100 nm thick, 100  $\Omega$  sq<sup>-1</sup> ITO coating were used as anodes. The substrates were washed by sequential ultrasonication in acetone, Semico clean, pure water, and isopropanol and then exposed to UV/ozone for 15 min to remove the adsorbed organic species. The PED-OT:PSS (Heraeus CleviosTM P VP.Al 4083) layer was spincoated at a rotation speed of 500 rpm for 3 s and then 3000 rpm for 60 s on the top of ITO, and annealed at 200 °C for 10 min. The various EMLs were deposited by spin-coating from a chloroform solution (0.25 wt%) at a rotation speed of 1000 rpm for 60 s in a glove box, giving films with thicknesses of ca. 35 nm. The substrates were transferred into the evaporation chamber. Organic layers and the Al electrode were formed by thermal evaporation at a pressure lower than  $1 \times 10^{-4}$  Pa. The deposition rates of the organic layers were  $< 1.0 \text{ Å s}^{-1}$ , while Liq and Al were deposited by the rates of 0.08 and 1.0-2.0  $\mathring{A}$  s<sup>-1</sup>, respectively. The *J-V-L* characteristics of the OLEDs were evaluated using a source meter (Keysight B2911A, Keysight 534 Technologies) and a luminance meter (CS-2000, Konica 535 Minolta, Japan) at a constant DC current at room temperature. The reproducibility of the device performance of the presented devices was confirmed by measuring at least four samples.

### Conflicts of interest

The authors declare no competing financial interest.

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