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### Post-Polymerization C-H Borylation of Donor-Acceptor Materials Gives Highly Efficient Solid State Near-Infrared Emitters for NIR-OLEDs and Effective Biological Imaging

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# Post-Polymerization C-H Borylation of Donor-Acceptor Materials Gives Highly Efficient Solid State Near-Infrared Emitters for NIR-OLEDs and Effective Biological Imaging

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KEYWORDS: near-infrared emission, bioimaging, post-polymerization modification, C-H borylation, low band-gap.

Post-polymerization modification of the donor-acceptor polymer, poly(9,9-dioctylfluorene-alt-ABSTRACT: benzothiadiazole), PF8-BT by electrophilic C-H borylation is a simple method to introduce controllable quantities of near-infrared (NIR) emitting chromophore units into the backbone of a conjugated polymer. The highly stable borylated unit possesses a significantly lower LUMO energy than the pristine polymer resulting in a reduction in the band-gap of the polymer by up to 0.63 eV and a red shift in emission of more than 150 nm. Extensively borylated polymers absorb strongly in the deep red / NIR and are highly emissive in the NIR region of the spectrum in solution and solid state. Photoluminescence quantum yields (PLQY) values are extremely high in the solid state for materials with emission maxima  $\geq$ 700 nm with PLQY values of 44% at 700 nm and 11% at 757 nm for PF8-BT with different borylation levels. This high brightness enables efficient solution processed NIR emitting OLEDs to be fabricated and highly emissive borylated polymer loaded conjugated polymer nanoparticles (CPNPs) to be prepared. The latter are bright, photo-stable, low toxicity bio-imaging agents that in phantom mouse studies show higher signal to background ratios for emission at 820 nm than the ubiquitous NIR emissive bioimaging agent indocyanine green. This methodology represents a general approach for the post-polymerization functionalization of donor acceptor polymers to reduce the band gap as confirmed by the C-H borylation of poly((9,9-dioctylfluorene)-2,7-diyl-alt-[4,7-bis(3-hexylthien-5-yl)-2,1,3-benzothiadiazole]-2c,2cc-diyl) (PF8TBT) resulting in a red-shift in emission of > 150 nm thereby shifting the emission maximum to 810 nm.

#### INTRODUCTION

The development of organic materials that are effective solid state near-infrared (NIR) emitters is of considerable significance for many applications including nightvision displays, sensors and in-vivo fluorescence imaging.<sup>1,2</sup> The latter is particularly important as it is a non-invasive diagnosis method that provides information not available by in-vitro or ex-vivo imaging.<sup>3-5</sup> However, efficient solid state NIR emission is challenging as organic materials that demonstrate substantial NIR emission (>700 nm) in solution often show severely decreased photoluminescence quantum yield (PLQY) values in the solid state.<sup>6</sup> This is generally due to aggregation caused quenching (ACQ)<sup>7</sup> an effect that is often considerable in NIR emitting materials as the prerequisite low band-gap is typically achieved through the incorporation of extended planar  $\pi$ -systems and/or strongly electron donating and withdrawing units. These structural features generally promote aggregation which can induce alternative nonradiative decay processes.<sup>8</sup> Whilst the majority of NIR emissive materials developed to date are based on small molecules the development of conjugated polymer NIR emitters is highly desirable for facilitating solution processing of organic electronic devices and for generating dye-loaded polymer nanoparticles for bio-imaging applications that are stable to leakage of the NIR emitter.<sup>9</sup> However, effective NIR emissive polymers are rare and those reported to date generally contain transition metal complexes, complex repeat units (undesirable from a cost and/or toxicity perspective) and / or low percentages of the NIR chromophore (to minimize ACQ) thereby reducing the signal to background fluorescence ratio in the NIR region.<sup>1,2,10</sup>

To apply these NIR emitting polymers for *in vivo* imaging they need to be dispersed in water as conjugated

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polymer nanoparticles (CPNPs).<sup>11-17</sup> CPNPs are highly desirable due to their low cost and low toxicity, particularly relative to heavy atom based emitters (e.g., CdSe quantum dots).<sup>18</sup> Fluorophores that have both high absorbance and good emission in the spectral region between 650 and 1500 nm are essential for *in-vivo* imaging as this region allows for greater photon penetration depths in mammalian tissue (millimetre versus micrometre with visible light).<sup>4,19</sup> An ideal agent for biomedical imaging should absorb in the deep red (or longer  $\lambda$ ), have a large Stokes shift, emit in the NIR region of the spectrum with high PLQY values, have good photostability, be non-toxic, and stable in biological fluids.<sup>20,21</sup> In contrast, the most widely used NIR emitters (such as the FDA approved dye indocyanine green (ICG)) have relatively low brightness, small Stokes shifts and limited photostability.<sup>22</sup> Developing improved NIR imaging agents is therefore an important current challenge and herein we report a new family of NIR emitting polymers that meet many of the requirements outlined above. Electrophilic C-H borylation is used as a facile procedure to modify BT containing polymers, such as poly(9,9-dioctylfluorene-alt-benzothiadiazole) (PF8-BT), and this results in a substantial reduction of the polymer bandgap (by over 0.5 eV). The borylated F8-BT polymers have large Stokes shifts, and are highly efficient NIR emitters with emission  $\lambda_{max}$  values of  $\geq$ 700 nm and solid state PLQY values up to 44%. Efficient solution processed NIR OLED devices and highly emissive, photostable CPNPs are constructed from these polymers. In phantom mouse studies the latter show superior signal to background ratios (SBR) for emission at 820 nm relative to ICG.

### **RESULTS AND DISCUSSION**

Synthesis of P1-4. Electrophilic C-H borylation of BT containing small molecules by the addition of BCl<sub>3</sub> and subsequent functionalization at boron using diarylzinc reagents has been previously reported by several of us to produce highly emissive low band-gap small molecules.<sup>23-27</sup> The C-H functionalization methodology can be extended to PF8-BT (Mn 41 kDa, Mw 89 kDa, in THF at  $35^{\circ}$ C) and the percentage of borylated F8-BT NIR chromophore units incorporated into the polymer can be simply controlled by varying the molar equivalents of BCl<sub>2</sub> used (with respect to the repeat unit) followed by the addition of ZnPh<sub>2</sub>. This results in the high yielding synthesis (>95%) of partially borylated P1-3 (10, 15, and 25% borylation, Figure 1, top) and fully borylated PF8-BT, P4, functionalized with C,N-chelated BPh<sub>2</sub> groups. These are all highly stable to protic species and are thermally stable to >300 °C. It should be noted that post-polymerisation borylation produces a regio-irregular polymer (each fluorene unit could contain either 0, 1 or 2 boryl groups, see Scheme S1). However, we have previously shown that both types of borylated units have similar low LUMO energies.<sup>23</sup>

**Electrochemical Properties.** Cyclic voltammograms (CVs) of **P1-3** all show two reduction waves, a major reversible wave associated with the unborylated F8-BT units and an additional minor reduction wave at a significantly less negative reduction potential (by >0.5 eV) associated with the lower LUMO energy borylated F8-BT units (Figures S20 and S21). In the maximum borylated polymer **P4** only a single reduction wave is



**Figure 1**. Top, synthesis of **P1-4**. Bottom, UV-vis absorbance of PF8-BT and borylated polymers A) in toluene ( $1 \times 10^{-5}$  M), B) as thin films, C) encapsulated into CPNPs (PP-CPNPs and Si-CPNPs at 7.5 µg/ml). Emission of PF8-BT and borylated polymers, D) in toluene ( $1 \times 10^{-5}$  M), E) as thin films, F) encapsulated into CPNPs (PP-CPNPs and Si-CPNPs at 7.5 µg/ml).

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observed which is associated with the borylated units (Figure S23).Post-polymerization modification is particularly attractive as it removes the need to synthesize and polymerize borylated monomers,<sup>28,29</sup> and therefore represents a facile method to significantly reduce the band-gap of PF8-BT from ~2.60 eV for PF8-BT to 1.94 eV post C-H borylation by principally altering the LUMO energy.

Photophysical Properties. Photophysical studies confirm a low band-gap for these polymers, which is in the region of 1.9 eV from the onset of absorption for P2-P4 (see tables  $S_2 - S_6$ ). It is noteworthy that the absorbance spectra of P1-3 in toluene solution shows absorption bands (~370-510 nm), associated with the unborylated F8-BT units, the intensity of which decreases as borylation level increases (Figure 1A and table S2). Concomitantly, the absorbance band associated with the borylated F8-BT units (520 to 650 nm) increases in absorption intensity on increasing the proportion of borylated chromophores. Upon exciting P2 in dilute solution intense photoluminescence from the unborylated F8-BT units (emission  $\lambda$ max = 527 nm) is observed with only a minor amount of emission from the borylated units (emission  $\lambda max = 706$ nm). Conversely,  $\mathbf{P_3}$  shows the majority of the emission from the borylated units (emission  $\lambda max = 710$  nm) and minor emission from the F8-BT units. The maximum borylated polymer P4 shows no observable optical features associated with unborylated F8-BT units (Figure 1A) and in toluene P4 shows a considerable Stokes shift of (~100 nm) with a emission  $\lambda$ max at ~702 nm with a high (for fluorophores with emission  $\lambda max > 700$  nm) PLQY value of 12%.

For organic electronic devices and when dispersed as CPNPs in water the solid state absorbance/emission characteristics of the polymers are more relevant. Polymers P1-P4 show very similar trends in absorbance to those in solution but are red shifted (Figure 1A and B). In contrast, the solid state PL spectra of the borylated polymers (Figure 1E) differed drastically from those recorded in toluene solution (Figure 1D). When P1 is excited at 468 nm in the solid state, the emission from the F8-BT units is almost completely absent with the majority of emission observed with an emission  $\lambda$ max of 700 nm and an exceptionally high (for deep red/NIR emitters) solid state PLQY of 44%. Emission from the unborylated F8-BT units is completely quenched in the solid state upon increasing the borylation level above 10% (P2-4) due to the increased concentration of borylated units in the polymer chain leading to an increased probability of energy transfer to the lower band-gap chromophore units. A progressive red shift in emission upon increasing the borylation level was observed with a solid state emission  $\lambda$ max of 700 nm for P1 (PLQY = 44%), 710 nm for P2 (PLQY = 32%), 731 nm for P3 (PLQY = 23%), and the largest red shift is observed in the fully borylated polymer P4 with a emission  $\lambda$ max value of 757 nm (PLOY = 11%). Whilst the solid state PL of all four polymers is high that of P4 is particularly notable as it red shifted by >200 nm relative to PF8-BT, and the PLQY value of 11% at 757 nm is extremely high for organic fluorophores emitting in the solid state with a emission  $\lambda max > 750$  nm.<sup>6</sup> Furthermore, there is a significant Stokes shift ( $\geq 130$  nm) in all these polymers whilst lifetime measurements revealed only prompt fluorescence ( $\tau = <10$  ns).

**OLED Devices.** With an appropriate spectral overlap to realise efficient energy transfer between PF8-BT and the borylated polymers PL studies on polymer blends were performed. This revealed excellent PLQY values, for example, upon excitation at 468 nm (optimal for unborylated PF8-BT) **P2** and **P3** blended with PF8-BT showed excellent PLQY values for solid state emission with  $\lambda$ max >700 nm (emission  $\lambda$ max = 715 nm and PLQY = 32-35%, Figure S18). Based on these promising results, solution processed deep red/NIR emitting OLEDs were fabricated using the borylated polymers (neat or as blends with PF8-BT) as the emissive layer. These devices were fabricated with the following architecture, ITO (45 nm) / Plexcore OC<sup>®</sup> AQ1200 (65 nm) / PF8-TFB (poly[(9,9-dioctylfluorenyl-2,7-diyl)-co-(4,4'-(N-(4-sec-

butylphenyl)diphenyl amine))]) (22 nm) / emission layer (EML) (100 nm) / Ba (4 nm) / Al (100 nm) / Ag (100 nm). Devices with the EML consisting of neat films of **P1** (Device 1) and **P3** (Device 2) showed EQE<sub>max</sub> values of 0.18 and 0.41%, respectively with both devices showing low EQE roll off at high current densities (EQE at 100 mA cm<sup>-2</sup>: Device 1 = 0.17%, Device 2 = 0.26%). OLED devices were not fabricated from **P4** due to its poor surface wetting and film forming properties. The electroluminescence emission  $\lambda$ max values ( $\lambda$ max<sub>EL</sub>) of Devices 1 and 2 were 688 and 716 nm, respectively (Fig. 2). The EQE<sub>max</sub> value for **P3** is amongst the highest reported for non-doped solution processed, fluorescent OLEDs with a  $\lambda$ max<sub>EL</sub> >700 nm, particularly with such a simple device structure.<sup>6</sup>



Figure 2. Electroluminescence spectra of devices 1 and 2.

**CPNP Properties.** The incorporation of a high percentage of NIR chromophore units in **P3** and **P4** leads to an appreciable absorbance in the deep red and excellent solid state emission in the NIR and suggests that these polymers are well-suited for use as in-vivo biological imaging agents. By contrast polymers **P1** and **P2** do not absorb significantly at  $\lambda > 600$  nm and are therefore not capable of excitation using photons in the spectral region required for deep tissue penetration. Recently, green emitting PF8-

BT has been encapsulated within diblock-copolymer CPNPs consisting of PEG-PLGA (PP)<sup>30,31</sup> or silica/Pluronic® F127 (Si) CPNPs.<sup>32,33</sup> Using an analogous approach **P3** and **P4** were encapsulated into either core shell PP or Si-CPNPs in which the conjugated polymer made up 5% and 1.4% of the theoretical CPNP mass, respectively. For both **P3** and **P4** systems, PP-CPNPs were larger than Si-CPNPs (~130 vs 30 nm) and were typically more polydisperse (Figure 3).



**Figure 3**. Top attributes of PP and Si-CPNP containing **P3** or **P4**, left Zeta potential (ZP) and right hydrodynamic diameter (mean  $\pm$  standard deviation (SD), n=3 CPNP replicates). Bottom TEM micrographs of representative CPNPs

The photophysical properties of the P3 and P4 polymers in the core shell CPNPs suspended in aqueous solution were comparable to those observed in the thin films, albeit with a slight blue shift in the emission. Nevertheless, all emission maxima are >710 nm and quantum yields remain excellent for emission in the NIR region of the spectrum (PLQY values (emission maxima) P3-PP = 14% (714 nm), P4-PP = 6% (727 nm), P3-Si = 16% (717 nm) and P4-Si = 12% (721 nm)) with the majority of emission being > 700 nm (Figure 1F). Importantly, all four CPNPs showed excellent photophysical and colloidal stability for up to 20 days in suspension (Figure S27). As intravenous injection is a likely administration route for clinical diagnostic applications, haemocompatibility studies were performed over a concentration of 3-300 µg/mL total CPNP solids concentration (Figure 4). Significant haemolytic activity was observed only for the PP-CPNPs at the highest concentration tested. According to Dobrovolskaia et al.<sup>34</sup> in-vitro haemolysis assay values >50% of the positive control are often associated with possible clinical complications, while values <25% typically denote no clinical concerns, thus all concentrations of P3-Si and P4-Si meet this criteria. Furthermore, nanoparticle-induced platelet aggregation was not observed in this study, suggesting good biocompatibility (Figure 4).



**Figure 4.** Top, Percentage erythrocyte haemolysis and bottom platelet aggregation (mean  $\pm$  standard deviation, n=3 donors, \*P<0.05, \*\*P<0.01). 100% haemolysis was defined as the amount of haemoglobin release following treatment with poly-L-lysine, while 100% aggregation was defined as the turbidity value following treatment with adenosine diphosphate. Vehicle controls consisted of phosphate buffered saline (PBS).

**Phantom Mouse Studies.** To quantitatively compare the performance of the borylated polymer CPNPs with ICG for bioimaging applications, a phantom mouse with autofluorescence and light-scattering properties matching those of mouse muscle tissue was used. A 10 µL sample containing equivalent fluorophore masses (and therefore variable total solids concentrations) was pipetted into a silicon tube and placed inside the phantom at two depths (ventral = 4.0 mm and dorsal = 17.0 mm). The signal:background ratio (SBR) of the fluorescence signal (total radiant efficiency of sample = ROI1) to autofluorescence background (total radiant efficiency of background = ROI<sub>2</sub>) was calculated from identically sized regions of interest (ROI) and defined as (ROI1-ROI2)/ROI2.35 Maximum SBR values for CPNP systems containing P3 and P4 were observed for the excitation and emission wavelengths of 640 / 820 nm, respectively, while ICG was imaged using a wavelength combination of 745 / 820 nm (Figure 5A/B). A linear increase in SBR values was observed for all CPNPs over a fluorophore mass range of 0.125 - 1 µg. CPNPs containing P4 showed substantially higher SBR values compared to those of P3 at all concentrations and tissue thicknesses, while Si-CPNP generally showed a slightly enhanced performance compared to PP-CPNP (Figure 5, right). Solutions of ICG in distilled water were non-linear over the concentration range tested and were outperformed by P4 systems at all concentrations bar one. Thus P4-Si CPNPs combine the desirable properties of a simple high yielding preparation procedure (P4-Si CPNPs are produced in >90% yield), low toxicity, high photostability, a large Stokes shift, good quantum yield, high brightness for emission in the NIR when excited in the deep red region of the spectrum and a signal to background ratio for emission at 820 nm that outperforms ICG in tissue studies.

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**Figure 5.** Imaging performance in a phantom mouse. Determination of maximal SBR over a range of excitation (A) and emission (B) wavelengths. SBR values for systems containing  $0.125 - 1 \mu g$  fluorophore (in 10  $\mu$ L) measured from the ventral (C) and dorsal (D) positions. E = representative images of signal patterns in a phantom mouse viewed from the ventral side. The color scales show total radiant efficiency values ([photon/s/cm<sup>2</sup>/steradian]/[ $\mu$ W/cm<sup>2</sup>]) ranging from 7.44 x10<sup>5</sup> (blue) to 6.34 x10<sup>8</sup> (red).

**Borylation of PF8TBT.** With the utility of postpolymerization C-H borylation as a method to significantly reduce the band gap established on PF8-BT we next sought to demonstrate the extension of this methodology to other polymers. Therefore the C-H borylation of poly((9,9-dioctylfluorene)-2,7-diyl-alt-[4,7-bis(3hexylthien-5-yl)-2,1,3-benzothiadiazole]-2c,2cc-diyl)

(PF8TBT), a donor acceptor polymer which has been studied in a range of optoelectronic devices, was investigated.<sup>36,37</sup> PF8TBT requires C-H borylation of a thiophene moiety instead of fluorene and was chosen as it is a potentially challenging polymer to borylate due to the sterically hindered environment around the borylation position of the thiophene (which has an ortho-hexyl chain).38,39 After the addition of excess BCl3 to PF8TBT (Mn 47 kDa, Mw 85 kDa) NMR spectroscopy (Figure S8-10) confirmed that significant C-H functionalization has proceeded. Subsequent degassing and functionalization with AlMe<sub>3</sub> enabled the resultant borylated polymer (P5, scheme 1) to be isolated with > 95 % C-H borylation level with the polymer stable to ambient atmosphere and wet (non-purified) solvents. The synthesis of P5 demonstrates that post borylation the boron centre can be functionalized with alkyl groups equally readily to aryl (as used with PF8-BT) further increasing the scope of this methodology. Importantly, a significant reduction in the optical bandgap of the borylated polymer P5 relative to PF8TBT was observed (optical band gap reduced from 2.07 to 1.65 eV,  $\Delta E = 0.42$  eV) in addition to a considerable red-shift in the photoluminescence emission maximum which changes from 655 nm for PF8TBT to 810 nm for P5 (measured as dilute DCM solutions see Figure S19). Therefore this post polymerization C-H functionalization method is a simple and general procedure for modulating the band gap and emission properties of common donor acceptor polymers. It is particularly useful for generating low band gap materials that are emissive in the NIR region of the spectrum.



Scheme 1. Synthesis of P5

#### CONCLUSION

In conclusion, post-polymerization modification by C-H electrophilic borylation is a simple and general high yielding method to introduce controlled concentrations of low band-gap chromophores into conjugated polymer main chains and red shift the emission by >150 nm relative to the pristine polymers. These polymers possess large Stokes shifts and exhibit extremely strong photoluminescence in the solid state with absolute PLQY values up to 44% in the far red/NIR region of the electromagnetic spectrum. Unoptimized solution processed OLED devices with the emission layer comprised of neat films of the borylated polymers showed good EQE<sub>max</sub> values for simple non-doped solution processed devices in the NIR region of the electromagnetic spectrum. It is also noteworthy that polymers containing higher borylation levels demonstrated red shifted EL and superior EQE values. Core shell CPNPs with encapsulated borylated polymers exhibited excellent colloidal and photophysical properties that are suitable for in vivo imaging applications. In phantom mouse studies these CPNPs outperformed the ubiquitous NIR dye, indocyanine green, across a range of dye loadings. We are currently investigating a variety of other low band-gap borylated BT-polymers for a range of applications.

### ASSOCIATED CONTENT

**Supporting Information**. The Supporting Information is available free of charge on the ACS Publications website at DOI: xx.xxxx/acsami.xxxxxx.

Experimental details, characterization of nanoparticles, NMR spectra, TGA plots, DSC plots, OLED data, CVs. GPCs (PDF)

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