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Induction of Circularly Polarized Electroluminescence from an Achiral Light Emitting Polymer via a Chiral Small Molecule Dopant.

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Organic light emitting diodes (OLEDs) are devices that utilize an emissive electroluminescent organic semiconductor (OSC) thin film sandwiched between two electrodes. Employing polymers as the OSC is a highly attractive approach due to their easy solution processibility. This results in low cost, large area device fabrication possibilities using printing techniques. Circularly polarized (CP) light is central to a large range of current and future display and photonic technologies, including highly efficient LCD backlights^[1], optical quantum information processing and communication^[2,3], and optical spintronics^[4]. There is therefore high interest in constructing CP light emitting devices. Whilst the use of wide-band reflective polarizers as passive components in PLED devices is one means to engineer a CP light output, this results in a relatively complex and thick device architecture, requiring an additional liquid crystal cell^[5]. The *direct* generation of CP light from a conventional PLED would be far more favorable in terms of simplicity, compactness, energy efficiency and product cost, and thus there has been significant interest in their development.

The principle strategy to date to fabricate direct CP-light emitting PLEDs has been the attachment of pendent chiral side-chains onto achiral conjugated polymer backbones. The degree of circular dichroism or CP-emission obtained from such systems is defined by the dissymmetry factor (g factor, where $|g| \le 2$), with g_{abs} describing circular dichroism, g_{PL} describing CP-photoluminescence and g_{EL} describing CP-electroluminescence. Whilst early successful attempts at using this approach, for example with a chiral-substituted poly(p-phenylene vinylene) (PPV), only obtained g_{EL} in the region of $10^{-3[6]}$, considerable synthesis and device optimization effort has resulted in polymers with g_{EL} up to 0.35 (see **Table S1** for comparative data). Subsequent analysis has established the role of the chiral pendent side chain, in aligning the polymer into a chiral arrangement $^{[7,8]}$. An elegantly simple and highly translatable alternative approach should be the use of a chiral small molecule dopant to induce this effect by blending it with a conventional achiral light emitting polymer (LEP). This would avoid the need for bespoke polymer synthesis and could potentially allow the use of a



wide range of device-optimized co-polymers emitting across the full visual spectrum. However, whilst a small number of reports have emerged on using relatively large chiral molecular architectures such as polysaccharides^[9], biaryl compounds^[10], or chiral solvent mixtures^[11] in order to induce CP-*photo* luminesence (CP-PL) from conjugated polymers, these approaches or dopants have significant limitations for translation into the solid state. Therefore the fabrication of direct CP-EL emitting devices has not been previously demonstrated by this method. Furthermore, the level of CP-PL observed from these studies was only very low ($g_{PL} = 10^{-2}$ to 10^{-3} , see Table S1), thus being far from competitive with the polymers bearing chiral side-chains.

Helicenes are intrinsically helical (and therefore chiral) conjugated molecules based on a spiral of fused carbocyclic or heterocyclic rings^[12] (for example **Figure 1a**). They can be separated into their right-handed and left-handed enantiomeric forms, which are well known to exhibit strong chiroptical properties, such as high optical rotary power and strong circular dichroism^[12,13]. This paper concerns our preliminary studies to investigate whether helicenes would act as a chiral dopant and confer their helical shape properties to a conventional achiral LEP, while using their organic semiconducting properties to ensure effective device operation of a direct CP-emitting PLED (structure shown in **Figure 1c**). 1-Aza[6]helicene (**Figure 1a**) was initially chosen for this study, in light of the fact that it can be scalably synthetically assembled and separated into its enantiomeric forms^[14]. While certain helicenes have been shown to directly emit low levels of CP photoluminescence in solution^[15] and racemic helicenes used as the sole organic semi-conductor component of non-CP emitting OLEDs^[16], the approach of using dopant quantities of these compounds to control the CP-emission from a conventional LEPs has not been previously reported. Furthermore, to the best of our knowledge, this particular helicene has not been previously studied with regard to organic semiconductor devices.

Each 1-aza[6]helicene enantiomer was blended into the conventional PLED material poly[9,9dioctylfluorene-co-benzothiadiazole] (F8BT, **Figure 1b**)^[17]. A variety of blending ratios (up to 55%) were employed to explore the impact of the helicene additive on the morphology and spectroscopic characteristics of the F8BT thin film. Whilst thin films of pure F8BT were relatively smooth, uniform and featureless, blend ratios of 7% and above resulted in a granular morphology with crystallite sizes ranging from 50 nm to 200 nm (see **Figure S2**). Preliminary grazing incidence wide angle X-ray scattering (GIWAXS) showed that the blend films contain a novel and highly orientated crystalline or co-crystalline phase, which is crystallographically distinct from that of the pure F8BT or pure helicene thin films, and has a much larger unit cell (see **Figure S3**). This previously unknown structure may resemble that of the intercalated or co-crystalline phase of PBTTT and PCBM recently reported. [18] Two absorption peaks at 325 nm and 450 nm were observed for the F8BT:helicene blend films, with a gradual increase in the 325 nm peak, attributed to 1-aza[6]helicene^[14] (see **Figure S4**), with increasing amounts of dopant (Figure 2a). An emission peak was observed at 580nm (Figure 2a), regardless of the percentage of helicene present, solely representing emission from F8BT. As expected, analogous spectra were obtained when the enantiomeric (-)-1aza[6]helicene was employed (data not shown).

Circular dichroism measurements of our blend films revealed that thin films of pure achiral F8BT have no CD response (**Figure 2b**), as expected. Fascinatingly, when just 7% of an enantiopure 1-aza[6]helicene dopant was added, a strong CD signal was observed around 450 nm, which by comparison with the spectra in **Figure 2a** can be assigned to absorption by F8BT. The strength of the CD response increases with increasing amounts of the helicene additive, with 7% helicene giving a g_{abs} value of 0.03 and 53% helicene giving a significantly



large g_{abs} value of 0.20. This value is similar to those obtained using the chiral side-chain approach (see **Table S1**), and is a >25-fold increase over using a polysaccharide dopant^[9]. Similar to the absorption spectra in **Figure 2a**, another CD peak was observed at 350 nm, which is associated with the known CD response of the helicene additive (see **Figure S4**).

Circularly polarized PL (CP-PL) spectra were then obtained for the blend films (**Figure 2c**). Unsurprisingly, no CP-PL signal was detected from the thin film consisting of pure achiral F8BT. A small amount (7%) of enantiopure 1-aza[6]helicene dopant leads to a big CP-PL response of the F8BT film, with the g_{PL} value exceeding 0.2. The CP-PL spectra spans a broad wavelength range from 520 nm to 675 nm and overlaps well with the PL spectra. Increasing the 1-aza[6]helicene blending ratio results in improvements of the g_{PL} factor, up to a significantly high value of 0.5 for the 53% helicene blend. Such a value is comparable or better than the g_{PL} factors achieved in other studies, which take dramatically different approaches to those disclosed herein (**Table S1**). Taken together, the high g_{PL} and g_{abs} factors, and the doped and undoped PL spectra (**Figure 2a**), all strongly suggest that the helicene dopant preorganizes the polymer into a chiral structure.

The significant chiroptical response of conductive polymers bearing pendent chiral side chains has been reported to originate from a liquid crystalline (LC) cholesteric arrangement of the polymer chains in the film $[^{19}]$, and this results in a g factor which is strongly dependent on film thickness (up to the LC helical pitch length of typically 100 to 200 nm). The film thickness of our blend was therefore varied to determine the effect it would have on the observed chiroptical response. Using an F8BT blend containing 6-7% of 1-aza[6]helicene and altering the spin coating speed, gave films of final thicknesses from 15 nm to 280 nm. The impact of film thickness on the g_{PL} factor for thicknesses between 90 nm and 280 nm is negligible (see **Figure S5**). At thicknesses below 90 nm, the PL response was too weak to be accurately quantified. Optical microscopy using crossed polarizers was carried out on all film thickness between 15 nm to 280 nm. In all cases, isotropic films were observed (see Figure S1). AFM was conducted, and a similar granular nanoscale morphology was observed for all films regardless of thickness (see **Figure S2**). The fact that the observed isotropic structure, nanoscale morphology, and g_{PL} factor are independent of film thickness likely negates a standard cholesteric origin of the chiroptical response. Instead we suggest we have a unique chiroptical co-crystalline phase, as supported by the GIWAXS data (see Figure S3). It has been previously proposed that a cholestric origin is required to give g factors above 10⁻³ in conjugated LC polymers^[19]. The fact we have g_{PL} factors of 0.2-0.5 indicates that appropriate chain packing on much smaller scales (2.36 to 4.7 nm [see Figure S3] as opposed to 100 to 200 nm) may also lead to strong chiroptical effects in conjugated polymers^[16]. Furthermore, as shown in Figures 2b and 2d, the CD and CP-PL spectra of independent blends prepared from either the left-handed (-)-1-aza[6]helicene or the right-handed (+)-1-aza[6]helicene give comparable but opposite responses. This confirms that the origin of the chiroptical response is the handedness of the helicene dopant employed.

Although this method has high practicality for the conversion of a conventional achiral conjugated polymer into a CP-emitting one, it was unclear whether the observed effect was specific to 1-aza[6]-helicene, or indeed more general. A thin film blend of F8BT with an unrelated enantiopure helicene, [7]helicene^[20] was therefore prepared (**Figure 3d**). Pleasingly, the CP-PL spectra from the 52% [7]helicene-F8BT blend, shown in **Figure 3e**, were comparable to those obtained with 1-aza[6]helicene at an analogous blending ratio (53%, see **Figure 2c**), with high g_{PL} factors up to 0.5.



The most important aspect of our approach was the ability to fabricate the F8BT:helicene blends into functional devices to directly emit CP-EL. Single-layer proof-of-concept PLEDs based on the F8BT blends containing 7% of either the left-handed (-)-1-aza[6]helicene or the right-handed (+)-1-aza[6]helicene were fabricated. Typical output J-V and luminescence curves obtained for these devices are shown in Figure 4a. To allow comparison, an undoped F8BT device was also fabricated (see **Figure 4d**). A bright emission achieving 3000 cd/m² was measured for both enantiomeric devices, with an efficiency of 1.1 lm/W. Although there is a slight increase in the turn-on voltage and a decrease in the maximum brightness compared to the reference device, the J-V-L characteristics show that the helicene dopant does not dramatically impact PLED operation or performance. Future synthesis should also allow the helicene to become an integral part of the device (eg. adjusting the position of the HOMO to assist hole injection and transport). The two vibronic peaks at 550 nm and 575 nm in the EL spectrum (Figure 4b) correspond to the emission observed in the PL spectra (Figure 2a). The EL spectrum is also very similar to the reference device (Figure 4d insert). The CP-EL spectra are shown in **Figure 4c**. These resemble the profiles of the CP-PL spectra (**Figure 2c**) and are equal and opposite depending on the enantiomer of the helicene used. A $g_{\rm EL}$ factor as high as 0.2 was observed, which also corresponds well to that obtained for CP-PL. Indeed, despite very little optimization, this CP-EL value is comparable or better than that obtained with liquid crystalline chiral side-chain polymers (**Table S1**). While it could be envisaged that increasing the amount of the helicene additive would increase this g_{EL} factor further, we generally found this to have a detrimental effect on overall device performance, possibly due to the increase domain height of the resultant thin films (see Figure S2).

In summary, we have successfully demonstrated a unique and effective approach to directly generate high levels of CP-EL based on doping a conventional achiral polymer with a chiral aromatic molecule. The results disclosed herein represent proof-of-concept, unoptimized devices, and it therefore remains highly probable that further study should result in the development of efficient PLED devices, which emit practically useful levels of CP light directly. The helicenes, being OSCs in their own right, can further be developed as an integral part of the device structure, for example as optimized hole transport / electron blocking materials. This strategy should be highly translatable, allowing the production of full-color CP-PLEDs from conventional, well-established PLED materials, using helicene dopants. We believe this will have a significant impact on current display technologies and benefit other disciplines from optical communication to quantum computing.

Experimental

Aza[6]helicene^[14] and [7]-helicene^[20] were prepared as previously reported and separated using preparative chiral HPLC.

Solution preparation and thin film deposition. Varying proportions of (+)-1-aza[6]helicene and (-)-1-aza[6]helicene (0% to 53% by weight) were blended into a 15 mg/ml F8BT solution (in toluene). Thin films of the blends were deposited by spin coating at 1200 r.p.m. for 60 seconds onto silica substrates for photophysics or AFM measurements. This gave an average film thickness of 160 nm for all blending ratios (between 6% and 53%1-aza[6]helicene), measured by Dektak [Veeco]. Varying film thickness from 90 nm to 280 nm were prepared by spin coating at different spin speeds from a 15 mg/ml, 6% (by w.t.) doped (+)-1-aza[6]helicene and (-)-1-aza[6]helicene solutions in order to study the effect on CP-PL. Film thickness below 90 nm were prepared using a 10 mg/ml solution at different spin speeds.

Photophysical and morphology characterization. Absorption and PL spectra of the blends at various blending ratios were measured by a Cary 300 UV-Vis spectrometer (Agilent



Technologies) and a FluoroMax-3 (Horiba Jobin Yvon), respectively. Thin film morphology studies were carried out using tapping mode AFM (Veeco multimode).

PLED fabrication and characterization. Pre-patterned ITO substrates were rinsed in an ultrasonic bath with acetone, IPA and deionized water before the deposition of PEDOT:PSS (H.C. Starck GmbH.) (50 nm). Thin films of 7% 1-aza[6]helicene: F8BT blend were then spin coated (1200 r.p.m. for 60 seconds) onto the PEDOT:PSS coated ITO substrates. Finally, a 20 nm Ca layer, capped by a 100 nm Al layer was thermally evaporated onto the organic layer to complete the PLED structure. JVL characterization was performed with a Keithley 2410 and a Topcon BM-9 luminance meter. PLED emission was assumed to be Lambertian. EL spectra were measured using an Ocean Optics USB 2000 charge-coupled device spectrophotometer.

Cross-polarized microscopy. An Olympus BX51 microscope was used to performed crossed-polarized microscopy on F8BT: (+)-1-aza[6]helicene blends. The film thickness was varied between 15 nm and 280 nm, fabricated as described above. 533 nm filter was inserted in between the two linear polarizers to improve the visibility of the images.

CD, CP-PL and CP-EL characterization. The circular dichroism spectra were obtained on a ChirascanTM-plus CD spectrometer. Left-handed and right-handed CP emission spectra from the blended thin film were collected using a linear polarizer and quarter-wave plate prior to a FluoroMax-3 spectrometer. The background introduced by the polarizer, the quarter-wave plate and the silica substrates were corrected by using the CP-PL results from a blank sample. The dissymmetry factor g in the CP-PL spectra were calculated from the equation $g = 2(I_L - I_R)/(I_L + I_R)$, $|g| \le 2$. I_R and I_L are the right-handed and left-handed emission intensities respectively. A similar method was used to analyze the CP-EL spectra. Instead of using the FluoroMax-3 as the spectrometer, the left-handed and right-handed EL spectra from the PLED was recorded using an Ocean Optics USB 2000 charge-coupled spectrophotometer.

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- [1] M. Schadt, Annu. Rev. Mater. Sci. 1997, 27, 305.
- [2] J. F. Sherson, H. Krauter, R. K. Olsson, B. Julsgaard, K. Hammerer, I. Cirac, E. S. Polzik, *Nature* **2006**, *443*, 557.
- [3] C. Wagenknecht, C. –M. Li, A. Reingruber, X. –H. Bao, A. Goebel, Y. –A. Chen, Q. Zhang, K. Chen, J. –W. Pan, *Nat. Photonics* **2010**, *4*, 549.
- [4] R. Farshchi, M. Ramsteiner, J. Herfort, A. Tahraoui, H. T. Grahn, *Appl. Phys. Lett.* **2011**, *98*, 162508.
- [5] M. Grell, M. Oda, K. S. Whitehead, A. Asimakis, D. Neher, D. D. C. Bradley, *Adv*.

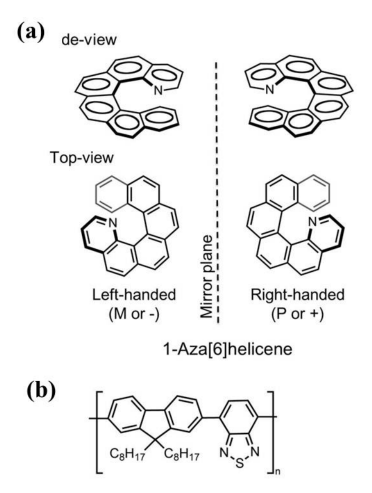


- Mater. 2001, 13, 577.
- [6] E. Peeters, M. P. T. Christiaans, R. A. J. Janssen, H. F. M. Schoo, H. P. J. M. Dekkers, E. W. Meijer, *J. Am. Chem. Soc.* **1997**, *119*, 9909.
- [7] Y. H. Geng, A. Trajkovska, S. W. Culligan, J. J. Ou, H. M. Chen, D. Katsis, S. H. Chen, *J. Am. Chem. Soc.* **2003**, *125*, 14032.
- [8] Y. H. Geng, A. Trajkovska, D. Katsis, J. J. Ou, S. W. Culligan, S. H. Chen, *J. Am. Chem. Soc.* **2002**, *124*, 8337.
- [9] S. Haraguchi, M. Numata, C. Li, Y. Nakano, M. Fujiki, S. Shinkai, *Chem. Lett.* **2009**, 38, 254.
- [10] K. Watanabe, I. Osaka, S. Yorozuya, K. Akagi, *Chem. Mater.* **2012**, *24*, 1011.
- [11] Y. Nakano, Y. Liu, M. Fujiki, *Polym. Chem-Uk* **2010**, *1*, 460.
- [12] Y. Shen, C. F. Chen, Chem. Rev. 2012, 112, 1463.
- [13] R. Hassey, E. J. Swain, N. I. Hammer, D. Venkataraman, M. D. Barnes, *Science* **2006**, *314*, 1437.
- [14] N. Takenaka, R. S. Sarangthem, B. Captain, Angew. Chem. Int. Ed. 2008, 47, 9708.
- [15] J. E. Field, G. Muller, J. P. Riehl, D. Venkataraman, J. Am. Chem. Soc. 2003, 125, 11808.
- [16] L. Shi, Z. Liu, G. Dong, L. Duan, Y. Qiu, J. Jia, W. Guo, D. Zhao, D. Cui, X. Tao, Chem. Eur. J. 2012, 18, 8092.
- [17] D. Kabra, L. P. Lu, M. H. Song, H. J. Snaith, R. H. Friend, *Adv. Mater.* **2010**, 22, 3194.
- [18] N. C. Miller, E. Cho, M. J. N. Junk, R. Gysel, C. Risko, D. Kim, S. Sweetnam, C. E. Miller, L. J. Richter, R. J. Kline, M. Heeney, I. McCulloch, A. Amassian, D. Acevedo-Feliz, □C. Knox, M. R. Hansen, D. Dudenko, B. F. Chmelka, M. F. Toney, J.-L. Brédas, M. D. McGehee, *Adv. Mater.* **2012**, *24*, 6071.
- [19] M. R. Craig, P. Jonkheijm, S. C. J. Meskers, A. P. H. J. Schenning, E. W. Meijer, *Adv. Mater.* **2003**, *15*, 1435.
- [20] R. El Abed, B. Ben Hassine, J. P. Genet, M. Gorsane, A. Marinetti, *Eur. J. Org. Chem.* **2004**, 1517.



Figure Legends:

Figure 1. (a) Molecular structures of 1-aza[6]helicene right- (P or (+)) and left- (M or (-)) handed enantiomers (b) molecular structure of the light emitting polymer F8BT; (c) single layer PLED device structure consisting of a 1-aza[6]helicene doped F8BT thin film sandwiched between a PEDOT:PSS coated ITO anode and a Ca/Al cathode; right- and left-handed σ^+ and σ^- circularly polarized electroluminescent polymer emission was respectively generated from (+)- and (-)-1-aza[6]helicene doped devices.



poly[9,9-dioctylfluorene-co-benzothiadiazole] (F8BT)

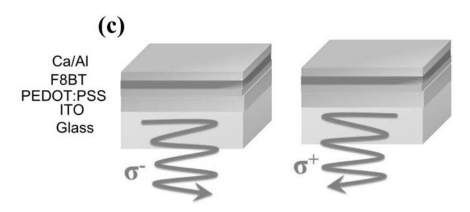
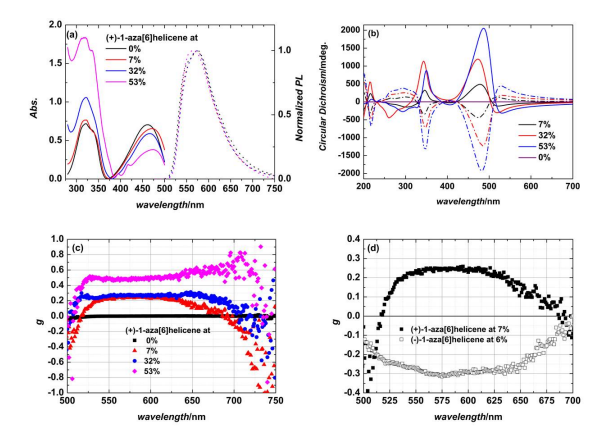
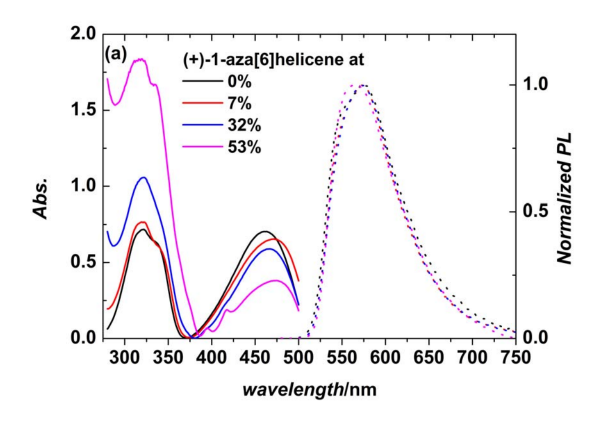




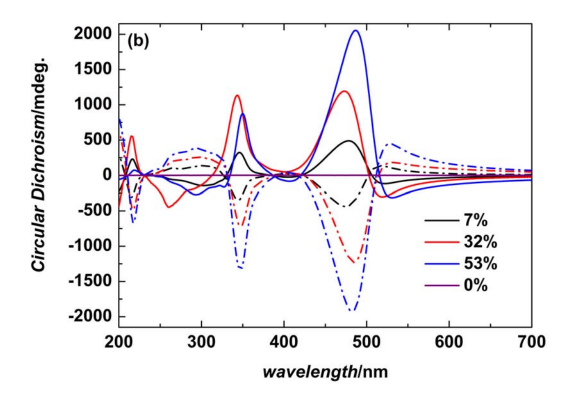
Figure 2. F8BT doped with varying ratios (% by weight) of 1-aza[6]helicene: (a) absorption (solid curves) and photoluminescence (PL) (dashed curves) spectra of F8BT doped with (+)-1-aza[6]helicene; (b) CD spectra of F8BT doped with (+)-1-aza[6]helicene (solid curves) and (-)-1-aza[6]helicene (dashed curves); (c) CP-PL spectra of F8BT doped with (+)-1-aza[6]helicene; (d) CP-PL spectra of F8BT doped with 7% (by weight) (+)-1-aza[6]helicene (filled symbols) and 6% (by weight) (-)-1-aza[6]helicene (open symbols).



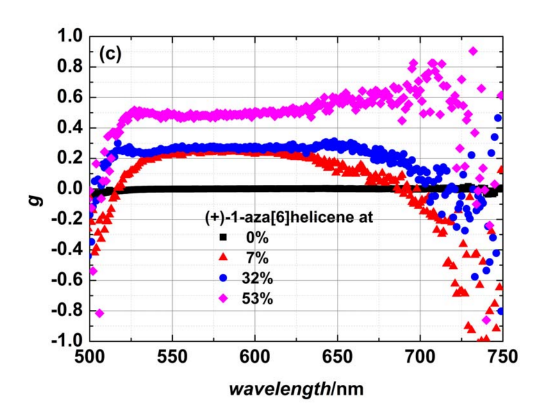
(a.)



(b.)



(c.)



(d.)

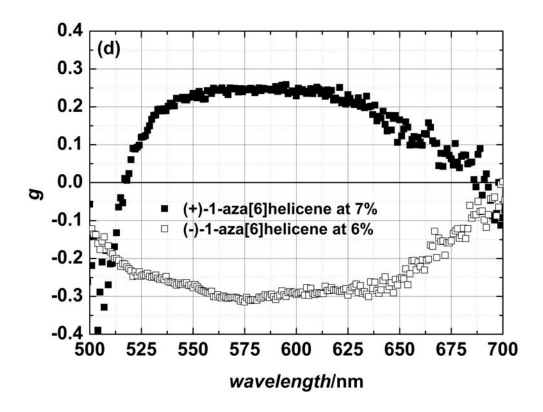
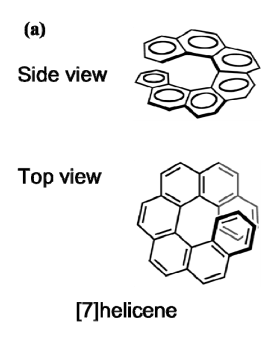




Figure 3. (a) Molecular structure of *P* (or +) [7]helicene; (b) CP-PL spectra of F8BT films doped with 52% (by weight) (+)- and (-)-[7]helicene.



(b)

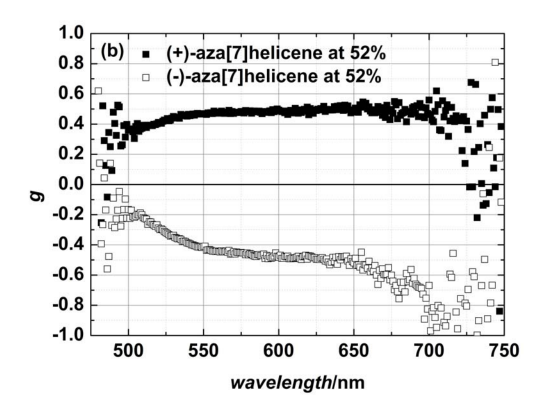
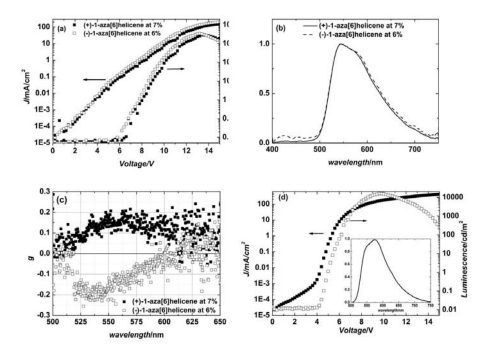
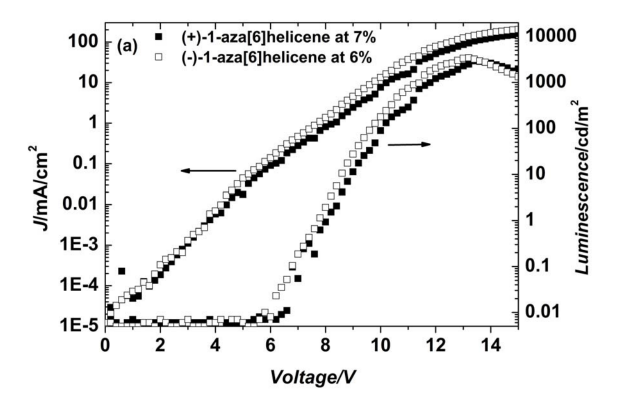


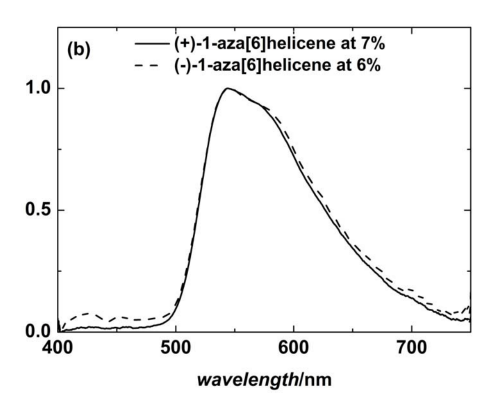


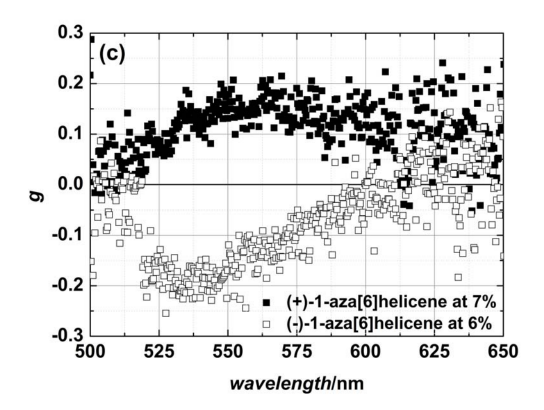
Figure 4. (a) Variation of current J (circles) and luminosity L (squares) with applied voltage V, (b) EL spectra and (c) CP-EL spectra of PLED device (see **Figure 1c**) containing a light emitting layer of F8BT doped with 7% (by weight) of (+)-1-aza[6]helicene (solid symbols) and 6% (by weight) (-)-1-aza[6]helicene (open symbols). (d) J-V-L curves of undoped a F8BT PLED with EL spectrum as the inset.













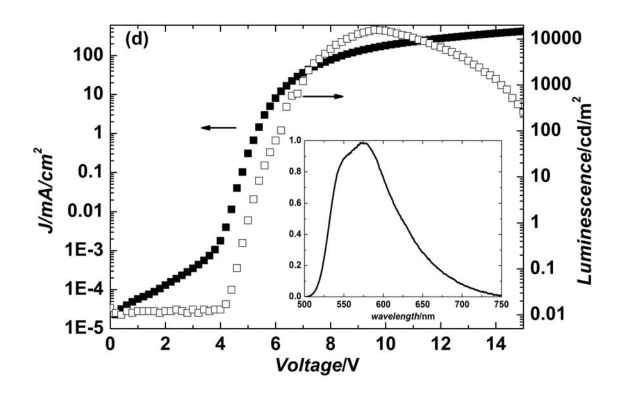


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There is high interest in the construction of organic light emitting devices, where the light output is circularly polarized (CP). Here we show that by simply doping the conventional light emitting polymer F8BT with a helically chiral aromatic molecule, we are able to generate substantial levels of CP-electroluminescence directly. Both photoluminescent and electroluminescent emission from the polymer is observed to become circularly polarized, with the sign of the CP emission directly determined by the handedness of the dopant.