# Chemical Science

ANNIVERSARY

Volume 11 Number 25 7 July 2020 Pages 6341-6592



ISSN 2041-6539



## Chemical Science



## **EDGE ARTICLE**

View Article Online
View Journal | View Issue



Cite this: Chem. Sci., 2020, 11, 6383

dll publication charges for this article have been paid for by the Royal Society of Chemistry

Received 11th February 2020 Accepted 3rd April 2020

DOI: 10.1039/d0sc00806k

rsc.li/chemical-science

## Oxidation promoted self-assembly of $\pi$ -conjugated polymers†

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Self-assembly is an attractive strategy for organizing molecules into ordered structures that can span multiple length scales. Crystallization Driven Self-Assembly (CDSA) involves a block copolymer with a crystallizable core-forming block and an amorphous corona-forming block that aggregate into micelles with a crystalline core in solvents that are selective for the corona block. CDSA requires coreand corona-forming blocks with very different solubilities. This hinders its use for the self-assembly of purely  $\pi$ -conjugated block copolymers since blocks with desirable optoelectronic properties tend to have similar solubilities. Further, this approach is not readily reversible, precluding stimulus-responsive assembly and disassembly. Here, we demonstrate that selective oxidative doping of one block of a fully  $\pi$ -conjugated block copolymer promotes the self-assembly of redox-responsive micelles. Heteroatom substitution in polychalcogenophenes enables the modulation of the intrinsic polymer oxidation potential. We show that oxidized micelles with a narrow size distribution form spontaneously and disassemble in response to a chemical reductant. This method expands the scope of  $\pi$ -conjugated polymers that can undergo controlled self-assembly and introduces reversible, redox-responsive self-assembly of  $\pi$ -conjugated polymers.

## Introduction

The self-assembly of block copolymer nanostructures1 has been explored extensively in recent decades2-4 for a wide range of applications.5-11 Unique properties and functions emerge in self assembled nanostructures, due to precise molecular arrangements over a range of length scales and control over assembly and disassembly in response to distinct stimuli.1-6,10 Crystallization-Driven Self-Assembly (CDSA) is one technique for controlled block copolymer self-assembly in solution.2 It involves the crystallization of a core block into nanostructures, driven by solvent interactions, while an amorphous corona block remains solvated for solution stability. 12,13 A preference for core crystallization drives the formation of micelle morphology.14 These micelles can be fragmented by sonication into shorter 'seed' structures, which have a low dispersity in size. Adding a solvated block copolymer, or a unimer, to the seeds preferentially causes micelle growth to occur epitaxially from the exposed crystal faces at the termini of the micelles. This control allows a range of complex micelle architectures to be formed such as block co-micelles,15 branched micelles,16 and larger hierarchical structures.17

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† Electronic supplementary information (ESI) available. See DOI: 10.1039/d0sc00806k

CDSA has been used with block copolymers containing  $\pi$ -conjugated cores and amorphous coronas.  $^{18-23}$   $\pi$ -conjugated polymers have a wide range of applications in optoelectronic devices,  $^{24-26}$  where their properties are heavily impacted by their nanoscale morphology.  $^{27,28}$  Using CDSA, one-dimensional (1D) nanofibers  $^{18-22}$  and two-dimensional (2D) platelets  $^{23}$  can be prepared with control over length and area, allowing applications in transistors  $^{29}$  and as platforms for photophysical studies.  $^{30}$  Investigating nanostructure formation in  $\pi$ -conjugated polymers is necessary to expand the scope of functional organic materials.

However, nanostructures formed by CDSA have significant limitations. First, micelles formed by CDSA are kinetically trapped by crystallization and only assemble in response to solvent or temperature changes. 12,13 Photo-responsive selfassembly has been demonstrated in specific  $\pi$ -conjugated polymers,31 but is irreversible. Expanding the scope of application requires nanostructures that assemble and disassemble in response to a diverse range of stimuli.5,6,10 Second, the core and corona blocks must possess significant solubility differences to allow selective crystallization. Unfortunately,  $\pi$ -conjugated polymers with desirable properties for optoelectronic applications, such as regioregularity, tend to have similar solubilities (i.e. they are primarily soluble in halogenated solvents) such that selective crystallization is very difficult. 24,32-34 One approach to address, rather than differentiating the blocks, is to eliminate the corona block and instead add bulky aliphatic side chains to

a crystallizable homopolymer.<sup>35,36</sup> However, this approach requires extensive sidechain engineering and has only been demonstrated on a unique homopolymer; a general approach capable of working even with minimal block differentiation would be of greater use. Thus, expanding the range of stimuli to promote the self-assembly and scope of polymers that can be assembled is an important challenge.

Polychalcogenophenes are a well-studied class of  $\pi$ -conjugated polymers37 often utilized in CDSA19,21,38 whose properties can be predictably modified through heteroatom substitution.26,39 S- and Te-containing polychalcogenophenes are called poly(thiophene)s (PThs) and poly(tellurophene)s (PTes), respectively. Relative to PThs, PTes possess slightly reduced solubility in halogenated solvents,40 while their planarity and energetic barriers to backbone rotation are increased.41 In addition, PTes have an oxidation potential ~0.2 eV higher than PThs, allowing the polymer to be oxidized more readily than other polychalcogenophenes. 42 Polychalcogenophenes possess other attractive features; regioregular polychalcogenophenes will crystallize, 43,44 which narrows their electronic band gap and increases their oxidation potential. 45,46 Polychalcogenophene oxidation, or oxidative doping, results in cations delocalized along the polymer backbone, which improves their chargetransport properties in functional materials. 42,47 Importantly, π-conjugated polymer oxidative doping also induces aggregation and precipitation from solution, 48,49 a feature which we realized could be exploited to promote self-assembly.

We hypothesized that in a  $\pi$ -conjugated block copolymer containing blocks with different oxidation potentials, selective oxidation would promote polymer self-assembly (Scheme 1). The oxidized block would aggregate and form a crystalline core, while the unoxidized corona block would promote colloidal stability, a process which we have termed Oxidation-Promoted Self-Assembly (OPSA). We took inspiration from

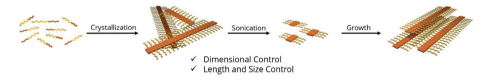
supramolecular polymerization, wherein a chemical input can act as a fuel for assembly or disassembly to allow stimulus-responsive, dynamic, or transient systems.  $^{50-52}$  OPSA uses a chemical oxidant as a fuel for self-assembly and can therefore use a reductant fuel for disassembly. Oxidation has previously been shown to promote self-assembly in non-conjugated systems, but never in  $\pi\text{-conjugated}$  polymers.  $^{53,54}$  This process allows the reversible, stimulus-responsive nanostructure formation promoted by oxidative doping and provides the potential for a general method to promote self-assembly in  $\pi\text{-conjugated}$  block copolymers.

## Results and discussion

To test our hypothesis, poly((3-(3,7-dimethyloctyl)thiophene)-b-(3-(3,7-dimethyloctyl)tellurophene)) (PTh<sub>80</sub>-b-PTe<sub>60</sub>) (Scheme 2) was selected as a model system. A 3,7-dimethyloctyl side chain was chosen to enable the controlled polymerization of the PTe block to a high molecular weight. 41 PTh<sub>80</sub>-b-PTe<sub>60</sub> was prepared by Kumada catalyst-transfer polycondensation (KCTP) using previously reported conditions,55 and characterized by 1H NMR spectroscopy and gel permeation chromatography (GPC) (Scheme 2) (see the ESI, Fig. S1-S4† for characterization and the Experimental section for synthetic details). The PTh and PTe blocks are 80 and 60 repeat units long, respectively, with a block copolymer dispersity (D) of 1.3. Heteroatom substitution allows us to prepare a block copolymer in which the blocks differ only in oxidation potential and solubility, allowing us to sequentially study the effects of solvent environment and oxidation on selfassembly.

Ultraviolet-Visible-Near Infrared Absorption Spectroscopy (UV-Vis-NIR) was used to understand the state of solvation and oxidation of the polymer. Solvated  $PTh_{80}$ -b- $PTe_{60}$  in chloroform displays a major absorption peak at 450 nm characteristic of the

#### Previous Work: Crystallization-Driven Self-Assembly (CDSA)



#### This Work: Oxidation-Promoted Self-Assembly (OPSA)



- ✓ Selective, Reversible Oxidation
- Assembly and Disassembly in Response to Redox Stimulus
- Coupled Electronic and Structural Control

Scheme 1 Strategy for 1D structural control in  $\pi$ -conjugated polymers. 1D CDSA is shown for comparison, in which crystallization drives the formation of a single micelle morphology. Sonication induces fragmentation of the polymer crystals, resulting in much smaller 'seed' micelles. An additional unimer preferentially crystallizes on the exposed crystal faces, resulting in length-controlled structures. OPSA utilizes selective oxidative doping of a single block of a  $\pi$ -conjugated polymer to drive self-assembly. Redox-active micelles with uniform lengths form which are reversibly disassembled upon reduction.

$$\begin{array}{c} Ph_2P_{Ni}Pph_2 \\ CI \end{array} \xrightarrow{CIMg} \begin{array}{c} CIMg \\ THF \end{array} \xrightarrow{CIMg} \begin{array}{c} CIMg \\ R \end{array} \xrightarrow{CIMg} \begin{array}{c} CIMg \\ R \end{array} \xrightarrow{CIMg} \begin{array}{c} CIMg \\ R \end{array} \xrightarrow{R} \begin{array}{$$

Scheme 2 Synthesis of  $PTh_{80}$ -b- $PTe_{60}$ .  $M_n$  and D determined by GPC performed at 1.0 mg mL<sup>-1</sup> in 1,2,4-trichlorobenzene at 140 °C.

 $\pi$ - $\pi$ \* transition of PTh, with a minor absorption shoulder at  $\sim$ 600 nm corresponding to the  $\pi$ - $\pi$ \* transition of PTe (Fig. 1b, black trace). After ageing for 18 h in a 95% dichloromethane (DCM)/5% chloroform solution, a fraction of PTh<sub>80</sub>-b-PTe<sub>60</sub> aggregates into micelles with a PTe core to form a partially aggregated solution. Partially aggregated PTh<sub>80</sub>-b-PTe<sub>60</sub> (blue trace) retains the PTh signal, with new peaks appearing from 650-900 nm which are associated with solid-state PTe that we attribute to PTe aggregation (consistent with solid-state absorption data; see the ESI, Fig. S5†). The observed vibronic structure indicates molecular alignments previously attributed to polymer crystallization. 44,56-58 Once this spectral change was observed, we added the oxidant iron(III) p-toluenesulfonate (FeTs<sub>3</sub>), which has been previously used to oxidatively dope PTe and is soluble in organic solvents.42 Upon the addition of iron(III) p-toluenesulfonate (FeTs<sub>3</sub>) (50 μM), we observe broad absorption in the near-IR region (red trace, >900 nm) indicative of (bi)polarons.<sup>59</sup> Polarons and bipolarons arise from radical cations and dications on the polymer backbone, compensated by the p-toluenesulfonate counter ion, and therefore demonstrate oxidation. The PTh absorption peak at 450 nm is unchanged while the PTe signal at ∼600 nm decreases, indicating that the PTe block is selectively oxidized. A further increase of FeTs<sub>3</sub> concentration (see the ESI, Fig. S6†) is correlated with the decreasing intensity of the crystalline PTe absorption from 650-900 nm and increasing intensity of the (bi) polaron signal. This inverse relationship strongly suggests that the degree of oxidation can be modulated, and that oxidation is selective for PTe.<sup>47</sup> Electron Paramagnetic Resonance (EPR) spectroscopy (Fig. 1d) confirms the presence of a radical cation species in oxidized PTh<sub>80</sub>-b-PTe<sub>60</sub>, with a singlet centered at a *g*-factor of 2.0023 consistent with an organic radical cation and identical to that of a free electron.<sup>60</sup> The evidence demonstrates that the PTe block has been selectively oxidized and suggests it crystallizes as well.

While PTe is selectively crystallized and oxidized, PTh remains fully solvated. Solvated poly(thiophene)s exhibit strong photoluminescence (PL) that is quenched by polymer crystallization.61 By contrast, poly(tellurophene)s are only marginally photoluminescent, displaying emission three orders of magnitude weaker than poly(thiophene)s.62 Emission observed from PTh<sub>80</sub>-b-PTe<sub>60</sub> must be due to the PTh block as any PTe emission is negligible; PL spectroscopy is therefore an excellent tool for exclusively assessing PTh solvation. PL spectroscopy of solvated PTh<sub>80</sub>-b-PTe<sub>60</sub> shows strong emission from 525–700 nm upon excitation at 450 nm (Fig. 1c) as we expect. Photoluminescence is slightly reduced in both partially aggregated and oxidized PTh<sub>80</sub>-b-PTe<sub>60</sub> due to quenching by crystallized PTe. However, the majority retention of photoluminescence is strong evidence for the neutral, solvated state of PTh. Both UV-Vis-NIR and PL spectroscopy indicate PTh solvation in the solvated, partially aggregated, and oxidized samples.

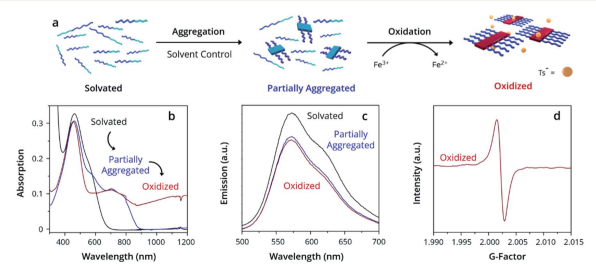


Fig. 1 Oxidation-promoted self-assembly. (a) Schematic depicting the aggregation and oxidation of  $PTh_{80}$ -b- $PTe_{60}$  by  $FeTs_3$  to oxidized micelles. Demonstrated by (b) UV-Vis-NIR spectra, (c) PL spectra, and (d) EPR spectra. All solutions prepared at 0.05 mg mL<sup>-1</sup>  $PTh_{80}$ -b- $PTe_{60}$ .

By examining the structure of the partially aggregated sample, we observe the limits of solvent-driven assembly. First, Dynamic Light Scattering (DLS) was used to measure the apparent hydrodynamic radius ( $R_h^{app}$ ) of polymer aggregates in solution. There are two size distributions in the partially aggregated sample (Fig. 2a, blue trace); a major distribution  $(R_{\rm h}^{\rm app} \sim 20 \text{ nm})$  attributed to an aggregate species and a minor distribution ( $R_{\rm h}^{\rm app} \sim 5$  nm) attributed to a unimer. Partially aggregated PTh<sub>80</sub>-b-PTe<sub>60</sub> can therefore best be described as a mixture of small aggregates and the unimer, which is consistent with the UV-Vis-NIR peak attributed to crystalline PTe (blue trace, Fig. 1b). This characterization is consistent with Scanning Transmission Electron Microscopy (STEM) images of the drop-cast, partially aggregated solution (see the ESI, Fig. S7†). We observe primarily amorphous features attributed to the unimer and a broad length distribution of 1D fibers, including  $\sim$ 10-20 nm long aggregates. The longer fibres likely result from unimer deposition on aggregates during drying.33 This demonstrates the presence of the unimer and small aggregates in the partially aggregated sample, and it highlights that the solvent conditions are insufficient to control the selfassembly of PTh<sub>80</sub>-b-PTe<sub>60</sub>. This may be due to similar solubilities, poor PTe crystallization limiting a crystallization-driven approach, or both.

Oxidation, conversely, promotes complete self-assembly into micelles. DLS indicates a single size distribution

(Fig. 2a, red trace) which is larger ( $R_h^{app} \sim 60 \text{ nm}$ ) than partially aggregated PTh<sub>80</sub>-b-PTe<sub>60</sub>. By Transmission Electron Microscopy (TEM), we observe exclusively 1D nanostructures after oxidation (Fig. 2b) with uniform widths of 20  $\pm$  2 nm. The crystalline core-forming PTe block has an average contour length of approximately 17 nm,63 suggesting that the micelle widths result from fully extended PTe upon crystallization. An average micelle contour length  $(L_n)$  of 53 nm (Fig. 2c) with an  $L_{\rm n}/L_{\rm w}$  of 1.12 is consistent with DLS measurements, corroborating that this is the species we observe in solution. The absence of a detectable unimer in both DLS and TEM suggests that oxidation promotes complete aggregation. To measure the widths of the full polymer micelles, i.e. of both the PTe core and PTh corona, we used Atomic Force Microscopy (AFM). The full micelle width is expected to be equal to the PTe core block width plus a large contribution from PTh,64 which has a contour length of 22 nm.63 We measure micelle widths of 49  $\pm$  4 nm by AFM (Fig. 2d), which is consistent with a crystalline PTe core with the PTh corona being present on both sides of the core (Scheme 3).

To rationalize the narrow oxidized micelle size distribution, we propose this mechanism for OPSA. First, the PTe aggregates formed by solvent conditions become preferentially oxidized due their higher intrinsic oxidation potential (Scheme 3).43,65 Next, growth occurs as the electron-rich unimer is attracted to the oxidized aggregates, while oxidized aggregates are repelled

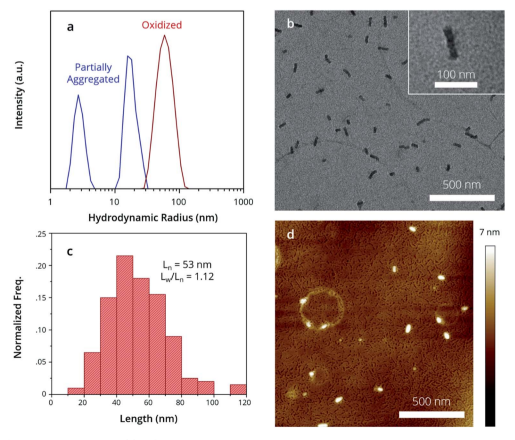


Fig. 2 Structure of oxidized micelle seeds. (a) DLS demonstrates oxidized micelle structure in solution, while morphologies are observed by (b) TEM images, (c) contour length distribution of micelle seeds in (a) and (d) AFM images.

Oxidized Micelles

Scheme 3 Proposed mechanism for oxidation promoted self-assembly

from one another. Finally, larger micelles are formed as the unimer is added epitaxially to the exposed crystal faces and becomes oxidized. 43,48,65 In this way, the process is promoted by both electrochemical induction, which causes the electron-rich unimer to be attracted to the oxidized aggregates, as well as the greater stablization of the radical cations by delocalization over a larger crystalline polymer domain.66,67 This mechanism is consistent with the increase in size and the loss of the unimer observed upon oxidation. The small size and narrow distribution of the oxidized micelles originates from the even smaller size of the initially aggregated PTe nuclei (Fig. 2a) which appear to only add the unimer and do not aggregate with each other.

The micelles in their oxidized form are colloidally stable, contain a crystalline core, and are likely kinetically trapped. Long-term colloidal stability was assessed by aging the PTh<sub>80</sub>-b-PTe<sub>60</sub> oxidized micelles under an inert atmosphere for 2 months in a sealed container. DLS and STEM measurements of the oxidized micelles after 2 months display no change in micelle size or morphology (see the ESI, Fig. S11 and S12†). This indicates that, in the absence of an external stimulus, the micelles remain assembled and colloidally stable. It further suggests

that the micelles are kinetically trapped in their oxidized state much like micelles formed by CDSA. 12,13,19 Kinetic trapping implies that there is a large energetic barrier to dissociation, potentially due to a crystalline core, which we investigated by Wide-Angle X-ray Scattering (WAXS). The oxidized micelles display a broad scattering peak with a d-spacing of 24.2 Å (see the ESI, Fig. S13†), wider than the previously reported homopolymer PTh and PTe d-spacings of 22.8 and 19.0 Å, respectively.68 This wider spacing between scattering planes suggests the intercalation of counter ions between the PTe crystalline lamellae. Given the need to charge balance the positively charged, oxidized PTe, it is expected that the presence of counter ions would increase the d-spacing. This is consistent with the oxidation-induced increase in d-spacing previously observed by our group in PTe homopolymers.42 Importantly, it demonstrates that a change in crystallization is associated with oxidation, and is consistent with micelles that are kinetically trapped in their oxidized form.

Perhaps the most striking feature of this process, however, is its reversible nature. Unique to OPSA, the addition of a chemical reductant causes disassembly of PTh<sub>80</sub>-b-PTe<sub>60</sub> oxidized

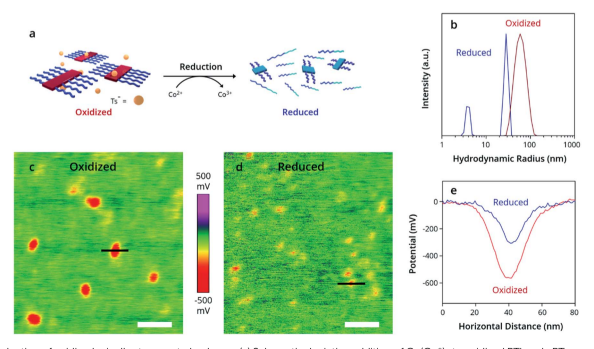


Fig. 3 Reduction of oxidized micelles to a neutral polymer. (a) Schematic depicting addition of  $Co(Cp^*)_2$  to oxidized PTh<sub>80</sub>-b-PTe<sub>60</sub>, resulting in reduced PTh<sub>80</sub>-b-PTe<sub>60</sub>. (b) DLS demonstrates the change in size distributions from oxidized (red trace) to reduced (blue trace) PTh<sub>80</sub>-b-PTe<sub>60</sub>. (c-e) Kelvin probe force microscopy images of the (c) oxidized and (d) reduced polymer, with potential measurements across indicated black lines plotted in (e). White scale bars = 100 nm.

micelles back to neutral aggregates and free unimers. Bis(pentamethyl-cyclopentadienyl)cobalt( $\pi$ ) [Co(Cp\*)<sub>2</sub>], a strong reducing agent with a high oxidation potential of 3.2 eV,<sup>69</sup> was added to the oxidized PTh<sub>80</sub>-b-PTe<sub>60</sub> at 100  $\mu$ M to return the polymer to its neutral state (Fig. 3a). By DLS, the reduced PTh<sub>80</sub>-b-PTe<sub>60</sub> displays two size distributions (Fig. 3b, blue trace) which are nearly identical to the partially aggregated PTh<sub>80</sub>-b-PTe<sub>60</sub> size distributions prior to oxidation (Fig. 2a, blue trace). No (bi)polaron states are observed by UV-Vis-NIR spectroscopy (see the ESI, Fig. S14†), with an absorption spectrum identical to the nucleated PTh<sub>80</sub>-b-PTe<sub>60</sub> pre-oxidation (Fig. 1b), confirming the reduction to a neutral polymer. This evidence demonstrates that oxidized micelles reversibly assemble and disassemble in response to redox stimuli and are only kinetically trapped when oxidized.

Kelvin Probe Force Microscopy (KPFM) (Fig. 3c–e) was used to measure the change in work function and enable the estimation of the doping level. KPFM allows spatial mapping of the chemical potential and work function ( $\varphi$ ) across a surface<sup>70</sup> referenced to the background substrate ( $\varphi_{\rm Si}\sim 4.6$  eV). We measured a lower potential and higher work function in the oxidized micelles (Fig. 3e). For comparison, we prepared PTe films dip-doped at 1, 5, and 10 mM FeTs<sub>3</sub>. The difference in work function between the neutral and oxidized samples is 270  $\pm$  90 meV for the micelles, which agrees with the 240  $\pm$  50 meV difference in work function for the 1 mM dip-doped PTe film (Table S1†). Because a 1 mM dip-doped PTe film has a doping level of 10–30%,  $^{42,47}$  we estimate a 10–30% doping level for the oxidized micelles.  $^{71,72}$ 

## Conclusion

This work demonstrates oxidation-promoted self-assembly. Oxidized micelles of a narrow size distribution reversibly form due to the selective chemical oxidation of a single block of a  $\pi$ -conjugated block copolymer. This selectivity is enabled by the 0.2 eV higher oxidation potential of PTe relative to PTh, such that self-assembly can be achieved with only heteroatom substitution differentiating the polymer blocks. The high similarity of the blocks in this study demonstrates the potential for a general method that can be used in a wide range of  $\pi$ -conjugated block copolymers as long as the blocks differ in oxidation potential. We hypothesized that the growth of the oxidized micelles and their narrow size-distribution are the consequence of a mechanism wherein the PTe nuclei are oxidized first in solution.

Critically, the structural dependence of the micelles on the polymer oxidation state allows oxidized micelles to be responsive to a redox stimulus. Beyond this proof-of-concept, OPSA could be employed to fabricate stimulus-responsive functional materials composed of all-regioregular  $\pi$ -conjugated polymers. Nanostructures formed by OPSA possess further utility as a platform to study the oxidative doping process. The demonstration of OPSA in a larger range of  $\pi$ -conjugated polymers and in a range of solvents for specialized applications is a subject of future work.

## **Experimental section**

#### **Materials**

All reagents were purchased from Sigma-Aldrich and used as received unless otherwise noted. Monomers and block copolymer PTh<sub>80</sub>-b-PTe<sub>60</sub> were synthesized according to the literature procedures. [1,2-Bis(diphenylphosphino)ethane](o-tolyl)chloronickel( $\pi$ ) was synthesized according to a literature procedure. Iron( $\pi$ ) p-toluenesulfonate hexahydrate (462861) was dried under vacuum at 75 °C to eliminate hydration adducts prior to use. DCM and chloroform were dried over calcium choride, distilled from phosphorus pentoxide, then stored over 4 Å molecular sieves in darkness in an atmosphere of dry  $\pi$ 0. THF was dried over sodium metal overnight, then cannulated into a sealed bomb and stored over 4 Å molecular sieves in darkness in an atmosphere of dry  $\pi$ 1.

#### Instrumentation

NMR spectra were recorded on a 500 MHz Agilent DD2. Chemical shifts are reported in ppm at ambient temperature. <sup>1</sup>H chemical shifts are referenced to the residual protonated chloroform peak at 7.26 ppm. Polymer molecular weights were determined with a Viscotek HT-GPC (1,2,4-trichlorobenzene, 140 °C, and 1 mL min<sup>-1</sup> flow rate) using Tosoh Bioscience LLC TSK-GEL GMH<sub>HR</sub>-HT mixed-bed columns and narrow molecular weight distribution polystyrene standards. A UV detector was used to detect the eluted polymer with respect to elution volume. UV-Vis-NIR absorption spectra were recorded on a Varian Cary 5000 UV-vis-NIR spectrophotometer using Hellma quartz cuvettes with a 5 mm path length. Baseline correction was performed by subtracting the spectrum of the pure solvent. Light scattering experiments were carried out using a Malvern Zetasizer Nano-ZS instrument equipped with a He-Ne laser ( $\lambda = 633$  nm). The TEM imaging was performed on a Hitachi HT7700 microscope at an acceleration voltage of 80 kV. Scattered electrons were blocked with an objective aperture in the focal plane of the objective lens. The STEM imaging was performed on a QUANTA FEG 250 microscope at an acceleration voltage of 20 kV. Dimension measurements were obtained by direct measurement using ImageJ of 200 individual structures on different parts of the grid. Carboncoated copper grids were exposed to UV-radiation for 30 minutes prior to use. One drop of the sample solution was dropped onto the grids and allowed to dry overnight. AFM and KPFM were conducted using a Bruker Dimension Icon atomic force microscope. AFM measurements were performed using a Bruker NCHV-A antimony (n) doped silicon tip with a nominal tip radius of 8 nm in tapping mode, with width measurements obtained on >25 different structures. KPFM measurements were performed using PQFNE-AL silicon nitride tips in Peakforce KPFM mode; micelle samples were prepared by immersing a silicon wafer in the sample for 30 s, then washing twice in pure DCM for 1 min. Films of PTe were spincoated from 10 mg mL<sup>-1</sup> chlorobenzene solutions onto indium tin oxide (ITO) on glass. Films were dip-doped into acetonitrile solutions of FeTs<sub>3</sub> for 3 minutes then washed with neat **Edge Article** 

acetonitrile to remove any residual FeTs3. WAXS measurements were performed using an Anton Paar SAXSpace. All selfassembly was performed under a dry N2 atmosphere.

#### Synthesis of PTh<sub>80</sub>-b-PTe<sub>60</sub>

Isopropyl magnesium chloride (0.97 eq.) was added to a solution of 2,5-dibromo-3-(3,7-dimethyloctyl)thiophene (173 mg, 0.45 mmol) in dry THF, and the mixture was stirred for 15 min at rt. External initiator Ni(dppe)(o-tolyl)Cl (2.57 mg, 4.4 µmol) was added in THF, and the mixture was stirred at rt for 30 min, after which an aliquot (1.0 mL) was removed and quenched with dilute HCl followed by extraction with chloroform. The organic layer was dried over MgSO4 and concentrated under reduced pressure. In a separate flask, isopropyl magnesium chloride (0.97 eq.) was added to a solution of 2,5-diiodo-2-(3,7dimethyloctyl)tellurophene (194 mg, 0.34 mmol), which was then added to the polymerization flask. The polymerization mixture was immersed in an oil bath at 40  $^{\circ}\mathrm{C}$  and allowed to stir for an additional 1 h before quenching with dilute HCl. The polymer was precipitated into methanol and purified by sequential Soxhlet extraction (methanol, hexane, and chloroform). The chloroform fraction was further purified by column chromatography in chloroform before being concentrated under reduced pressure to yield a dark purple solid (54.2 mg, 27%). GPC profiles (see the ESI, Fig. S4†) and <sup>1</sup>H NMR spectra of the monomers and block copolymer (see the ESI, Fig. S1-S3) are provided in the ESI.†

#### General procedure for self-assembly of PTe-PTh

1.0 mg of the block copolymer poly((3-(3,7-dimethyloctyl) thiophene)-b-(3-(3,7-dimethyloctyl)tellurophene)) was dissolved in 20.0 mL of 95% DCM and 5% chloroform in a vial to form a 0.05 mg mL<sup>-1</sup> red solution. The polymer solution was then allowed to age for 18 h, becoming orange-yellow in colour. Iron(III) p-toluenesulfonate (0.57 mg, 1.0 μmol) was added in a 0.5 mM standard solution in THF to a separate vial, and the solvent was removed under reduced pressure. The polymer solution was added to iron(III) p-toluenesulfonate (50 μM), after which the solution immediately turned bright yellow. This solution was allowed to age 18 h prior to characterization.

#### Procedure for disassembly of PTe-PTh

20.0 mL of a 0.05 mg mL<sup>-1</sup> solution of the block copolymer in 95% DCM and 5% chloroform containing iron(III) p-toluenesulfonate (50 µM) was prepared as described above. Bis(pentamethyl-cyclopentadienyl)cobalt(II) (0.66 mg, 2.0 µmol) was added in 0.1 mL DCM, after which the solution changed from bright yellow to orange. The solution was allowed to age 1 h prior to characterization.

### Conflicts of interest

The authors declare no competing financial interest.

## Acknowledgements

This work was supported by the University of Toronto, the NSERC, the CFI, and the Ontario Research Fund. D. S. S. is grateful to the Connaught Foundation for the McLean Award and the NSERC for the E. W. R. Steacie Memorial Fellowship. We are grateful to Nimrat K. Obhi and Mitchell A. Winnik for editing and advice.

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