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Polymer Nanowires with Highly Precise Internal Morphology and Topography

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ABSTRACT: The construction of precise soft matter nanostructures in solution presents a challenge. A key focus remains on the rational design of functionalities to achieve the high morphological complexity typically found in biological systems. Advances in controlled polymerisations and self-assembly increasingly allow approaches towards complex hierarchical nanomaterials. By combining tailor-made cylindrical polymer brushes, block copolymers and interpolyelectrolyte complexation-driven self-assembly, we herein demonstrate a facile construction of uniformly compartmentalised and topographically structured polymeric nanowires in aqueous media. The approach offers a modular avenue in programming the internal morphology of polymer nanowires by varying the block copolymers composition and topology.

Many systems in nature reveal a superior ability to compartmentalise functionality and arrange chemistries in three dimensions. Similarly, precise solution-based synthetic soft-matter nanostructures are being pursued; however, their build-up is challenging as it requires a detailed control of the structure formation and the underlying processes. While DNA nanotechnology already allows rationally engineered nanostructures with high precision, Multiblock polymers, particularly diblock copolymers (DB) still lack comparable structuring capabilities despite their widespread use in many applications. Taken that the underlying design principles for complex structures are mastered for DB, they could offer robust and cost-effective alternatives. Their self-assembly is driven by relatively weak competing interactions between the polymer blocks and the solvent, additionally allowing control *via* external stimuli (e.g. solvent changes, pH, temperature) or by incorporating supramolecular interactions. The immense potential of DBs for the construction of complex solvent-based nanostructures has been demonstrated by their assembly to multicompartment nanoparticles and their subsequent assembly to higher orders. A different approach uses crystallisation-driven self-assembly of DBs from seed particles to produce dispersions of complex, compartmentalised 1D and 2D assemblies. A simple, yet powerful, method for hierarchical supramolecular polymer self-assembly builds on the electrostatic complexation between polyanions and polycations, where interpolyelectrolyte complexation (IPEC) enables the formation of nanostructured and multicompartment polymer particles in solution. While IPECs were originally used to produce micelles, 12-16 vesicles, 17-18 or multilayered architectures, 19-22 they have recently exemplified

their potential as a powerful tool to interact with polymer particles — leading to polymer architectures that are otherwise difficult to realise.²³ For example, homopolymer complexation onto polymer micelles has yielded multicompartment particles with collapsed and patchy shells.²⁴ Patchy morphology can also be induced *via* intramolecular complexation on a core-shell cylinder carrying both positive and negative charges in its shell.²⁵ Using diblock copolymers containing a neutral and charged block, oppositely charged micelles can be switched charge-neutral, and subsequently used in photodynamic therapy.²⁶ Recently, IPECs have also shown potential in the design of non-equilibrium polymer micelles capable of reversibly switching their assembly upon change of temperature or salt concentration.¹⁶ Elegant structures *via* complexation of DBs onto soft templates have been predicted – ranging from spherical micelles to fibrillar (1D) and discoidal/planar (2D) particulates.²⁷

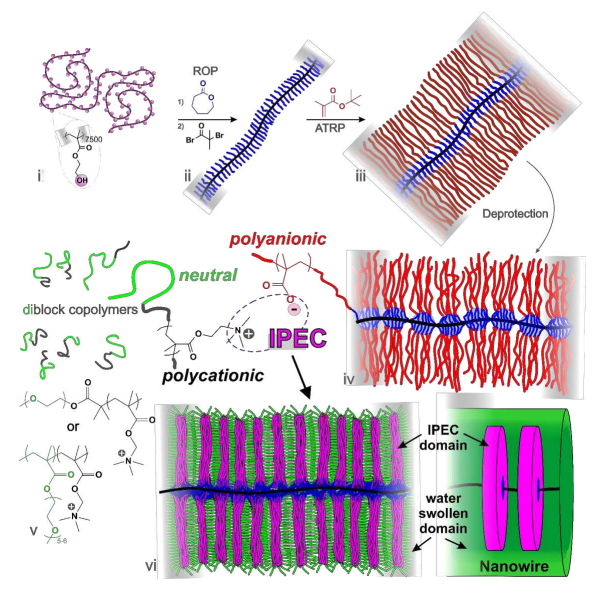
An intriguing 1D polymer nanomaterial (i.e. template material) are cylindrical polymer brushes (CPBs) – also known as molecular brushes, bottlebrush polymers, or comb-shaped polymers – allowing a high level of control over backbone and side chain lengths, and their distribution.²⁸ CPBs are densely grafted polymers, avoiding chain entanglement and existing as unimolecular entities. Given their covalent construction and unique architecture, they find increasing use in self-assembly,²⁹ nanomedicine³⁰ or as polymer scaffolds to produce tailor-made organic and inorganic (nano)materials.³¹⁻³⁶

In this work, we designed core-shell polyelectrolyte CPBs as 1D templates and used IPECs to produce centipede-like polymer nanowires with uniform topographical features and internal compartmentalisation along the CPB backbone (Scheme 1).

The template was synthesised by first grafting polycaprolactone (PCL) from a high-molecular weight poly(hydroxyethyl methacrylate) (PHEMA₇₅₀₀) backbone via ring-opening polymerisation. The terminal hydroxy groups of the PHEMA₇₅₀₀-g-PCL₁₄ homopolymer side chains were modified with α -bromoisobutyrate initiators,³³ to allow the grafting of a poly(tert-butyl methacrylate) (PtBMA) shell using atom transfer radical polymerisation (ATRP) (Scheme 1, Figure S1). Monomer conversion was monitored by proton nuclear magnetic resonance (¹H NMR) spectroscopy, which was used to calculate the side chain length, based on grafting efficiencies from PHEMA polyinitiator backbones³⁷ and PCL CPBs³³ (Figure S2).

Removal of the *tert*-butyl groups with trifluoroacetic acid, followed by dispersion in an alkaline buffer solution (pH 10) yielded negatively charged polyelectrolyte core-shell CPBs, namely PHEMA₇₅₀₀-*g*-[PCL₁₄-*b*-PMAA₃₀₀]. The complete removal of the *tert*-butyl groups was confirmed by the disappearance of the characteristic signal at 1.4 ppm in the ¹H NMR spectrum of the CPB (Figure S3-1), combined with Fourier-transform infrared spectroscopic (FTIR) analysis (Figure S3-2).

Scheme 1. (i) PHEMA is sequentially grafted-from with ε -caprolactone (ii) and tBMA to yield (iii) coreshell CPBs which after deprotection forms (iv) a 1D polyelectrolyte template with a pearl-necklace core structure. IPEC formation with (v) oppositely charged DBs yields (vi) uniformly structured nanowires featuring IPEC discs along the long-axis.



The worm-like conformation of the core-shell brushes was studied by atomic force microscopy (AFM). Before deprotection, the hydrophobic CPB templates formed 'monolayer islands' on mica upon solvent evaporation (THF), with core and shell clearly visible (Figure S4-1). After deprotection, the morphology of the polyelectrolyte template brushes in aqueous solution revealed individual cylindrical brush particles (Figure 1A, Figure S4-2). AFM height analysis indicated significant roughness due to CPB core segmentation. AFM analysis in the dry state (Figure 1D) revealed cross-sectional heights from 5 nm (between two beads, blue plot) to 8 nm (on top of beads, red plot), with apparent flattening due to solvent evaporation. Cryogenic transmission electron microscopy (cryo-TEM) better highlighted the distinct pearl-necklace structure, where the hydrophobic PCL core minimises its interaction with water to form spherical 'pearls' or 'beads' along the brush backbone (Figure 1B, C and Figure S5) – suggesting a distinct core compartmentalisation along the brush long-axis. A grey-scale analysis revealed a constant bead-to-bead distances $(8.1 \pm 2.6 \text{ nm}, \text{Figure 1E})$ and a narrow distribution of the bead diameters $(10.9 \pm 2.2 \text{ nm}, \text{Figure 1F})$.

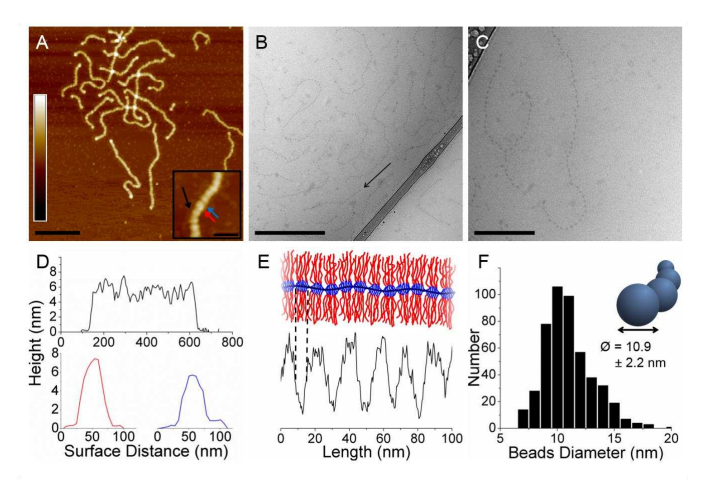


Figure 1. AFM height image of CPBs drop-cast from pH 10 buffer onto freshly-cleaved mica (A). Cryo-TEM images in pH 10 buffer (B, C). AFM cross-sectional analysis on mica across several beads (black), across a single bead (red) and between two beads (blue) (D). Grey-scale analysis across the beads (arrow in (B)) (E). Statistical analysis of bead diameters (data extracted by counting 500 beads on several images) (F). Scale bars: (A, B) 500 nm, (C) 200 nm, (A inset) 100 nm. AFM z-scale: 20 nm.

Next, we synthesised bis-hydrophilic diblock copolymers containing a poly(2-dimethylamino ethyl methacrylate) (PDMAEMA) block as well as either a linear polyethylene oxide (PEO) or comb-like poly(ethylene glycol) methyl ether methacrylate (PEGMA) blocks (Figure S1). The DB syntheses are described in the Supporting Information (Figure S6, S7 and Table S2). PDMAEMA was quaternized with methyl iodide following a published procedure³³ to yield copolymers with a positively charged poly[2-(methacryloyloxy)ethyl trimethylammonium iodide] (PMETAI) block and a non-ionic (PEO or PEGMA) block. The quaternization enabled IPEC formation at alkaline pH, at which PDMAEMA would otherwise be waterinsoluble.

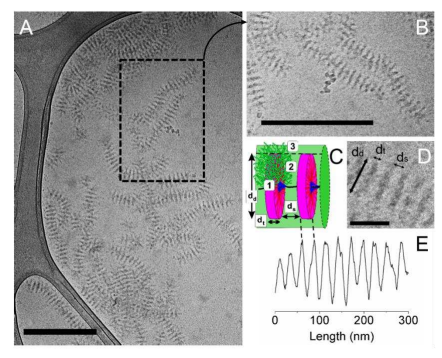


Figure 2. Cryo-TEM image of nanowire-A displaying centipede-like architectures in pH 10 buffer (A, B). Scheme illustrating various features within a nanowire (C) and an exemplary micrograph (D) and grey-scale analysis (E) of one nanowire extracted from (B). Scale bars: (A, B) 500 nm, (C) 100 nm.

We next explored the formation of IPECs between the polyelectrolyte brush and oppositely charged DBs. The IPEC-structured polymer nanowires are denoted as nanowire-X (X represents the DB used for the formation). To produce the nanowires, the CPB template and DBs were mixed in alkaline suspensions using nominally stoichiometric ratios of the negative and positive charges to compensate all charges. For nanowire-A, formed by complexing PEO114-b-PMETAI45 (DB-A) onto the CPB, 6-7 chains of DB-A vs. one PMAA₃₀₀ brush side chain were required, to achieve charge neutrality. After IPEC formation, AFM revealed a significant surface roughness along the nanowire. While the cross-sectional height showed only a slight increase of ca. 25 % (Figure S8-1), the nanowires showed a more pronounced pearl-necklace structure in the dry state upon deposition onto mica (Figure S8-2) compared to the pristine brush template. Cryo-TEM on the other hand revealed remarkably regular patterns along the backbone (Figure 2). This suggested directed DB self-assembly along the CPB templates with the hydrophobic IPEC domains (i.e. PMAA/PMETAI-domains, complexed by stoichiometric charges of the polyanion and polycation; indicated as 1 in Figure 2C) and hydrophilic PEO domains (indicated as 2 in Figure 2C) leading to their mutual repulsion. IPECs showed enhanced contrast in TEM, while PEO between or beyond the IPEC (Figure 2D; regions 2 and 3 in Figure 2C) was highly swollen with water and therefore not resolved. The IPEC appeared to be a lamellar disc, with disc diameter (dd IPEC-A) of 59.3 ± 9.2 nm, sandwiched between PEO domains and perpendicular to the backbone of the template brushes. Each disc within nanowire-A was separated by regular distances (disc-to-disc spacing, $d_{\text{SIPEC-A}} = 14.0 \pm 2.7 \text{ nm}$), which is dictated by the stabilising PEO block that is sandwiched between the IPEC discs (Figure 1C). Similarly, the average disc thickness ($d_{\text{tIPEC-A}}$) of 11.0 ± 1.8 nm was very uniform (Table 1). A trade-off between minimising the IPEC/water interface and generating sufficient space for the PEO chains ultimately drives the formation of this morphology. The volumes occupied by the IPEC (~44 vol%) and the water-swollen PEO (~56 vol%) confined within the nanowire, suggested a lamellar arrangement of the hydrophobic and hydrophilic domains.

Table 1. Overview of DB composition and IPEC-induced nanowire features.

	DBa			IPEC ^b		
	Composition	charged / uncharged	M _n (kDa)	Disc diameter, d_d (nm)	Disc-to-disc spacing, d_s (nm)	Disc thickness, $d_t(nm) \\$
A	PMETAI ₄₅ -b-PEO ₁₁₄ (DB-A)	73 / 27	18	59.3 ± 9.2	14.0 ± 2.7	11.0 ± 1.8
В	PMETAI ₄₈ -b-PEGMA ₁₀₃ (DB-B)	32 / 68	45	100.7 ± 20.7	22.9 ± 5.4	-
C	PMETAI ₂₅ -b-PEGMA ₇₆ (DB-C)	25 /75	30	76.7 ± 16.3	11.6 ± 2.9	8.6 ± 1.5

^aDetermined via ¹HNMR; ^bDetermined via grey-scale analysis of cryo-TEM images.

To investigate the effect of the DB composition, we next used PMETAI₄₈-b-PEGMA₁₀₃ (DB-B) for the complexation. DB-B had a similar number of repeat units to DB-A, but due to the different molecular weights of the repeat units of the non-ionic blocks (PEGMA = 300 g·mol⁻¹ vs. PEO = 44 g·mol⁻¹), the fraction of the non-ionic block was much higher. The similar PMETAI block length meant however that 6-7 copolymer chains per brush side chain were still needed to compensate the charges during nanowire-B formation. Thus, we could directly compare the effect of a larger non-ionic block fraction on the IPEC formation and resulting nanowire morphology (Figure 3D, E). In comparison to nanowire-A, this resulted in a less organized assembly, with less well-defined internal IPEC structures. The bulkiness of PEGMA led to more chain crowding and the increasing volume demand could not accommodate the formation of regular discs but forced the IPEC domains to develop distinct protrusions (Figure 3D, E). The higher steric demand was further reflected in the increased feature spacing ($d_{s \text{ IPEC-B}} = 22.9 \pm 5.4 \text{ nm}$).

Finally, we used PMETAI $_{25}$ -b-PEGMA $_{76}$ (DB-C) to form nanowire-C. The block ratio was comparable to DB-B, but due to the relatively shorter PMETAI block, double the number of polymer chains (~12) of DB-C are needed for charge compensation. The reduced molecular weight allowed nanowire-C to again develop a lamellar IPEC pattern (Figure 3G, H), albeit thinner than in nanowire-A due to the shorter PMETAI block length ($d_{t \, IPEC-C} = 8.6 \pm 1.5 \, nm$). Comparable occupied volumes (~43 vol% IPEC, ~57 vol% PEGMA) yielded IPEC discs featuring uniform disc-to-disc spacings ($d_{s \, IPEC-C} = 11.6 \pm 2.9 \, nm$) and overall diameters ($d_{d \, IPEC-C} = 76.7 \pm 16.3 \, nm$). Considering the regularity of the bead distance (centre-to-centre, 19.0 $\pm 2.5 \, nm$) in the template brush, the centre-to-centre distance between each disc (Table S4) matched that of nanowire-A (25.0 $\pm 4.5 \, nm$) and nanowire-C (20.2 $\pm 4.4 \, nm$). We propose that the pre-orientation of the side chains in the pearl-necklace structure of the template brush directs the regular distances of the forming IPEC structures. The formation of lamellar discs is also likely to result from our flexible brush template, as a previous study on stiff cellulose nanocrystals has shown IPECs would energetically favour the formation of IPEC helices along the long axis.³⁸

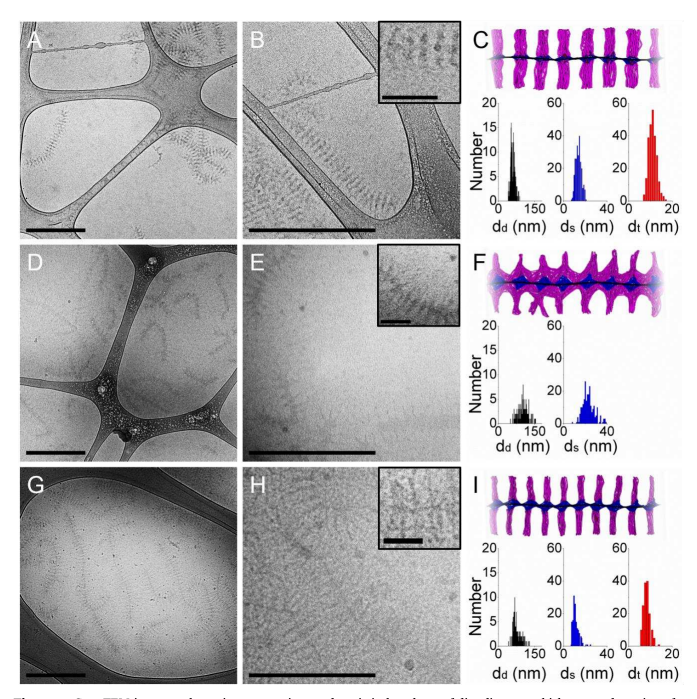


Figure 3. Cryo-TEM images, schematic cross-sections and statistical analyses of disc diameter, thickness, and spacing of nanowire-A (A-C), nanowire-B (D-F) and nanowire-C (G-I). Scale bars: (A, D, G) 500 nm, (B, E, H) 200 nm, (insets) 100 nm. Data extracted from various cryo-TEM images on >150 discs.

In conclusion, nanostructured 1D soft matter can be accessed from a stoichiometric complexation of core-shell polyelectrolyte brush templates and bis-hydrophilic DB containing a neutral- and oppositely-charged block. A regular disc-like lamellar self-assembly directed along the cylindrical polymer brush axis was observed. The well-defined and uniform patterns, i.e. compartments along the 1D polymer backbone, can be tailored by varying DB molecular weight, composition and topology. Altering the number of complexing polymer chains can be a means to alter the overall structure and disc thickness, while changing the molecular weight and block ratio of the DBs showed a significant effect on nanowire morphology and developed features. Combining the highly modular brush particle synthesis and the tuneable structuring approach using IPEC formation may open further avenues towards nanoarchitectures featuring increasing complexity and topographical features, and lead to

opportunities to arrange surface chemistries in ways that even mimic bioactive nanoparticles (e.g. viruses) or brush-on-brush structures (e.g. proteoglycan aggregates).

ASSOCIATED CONTENT

The Supporting Information is available free of charge on the ACS Publications website at DOI:10.1021/jacs.8bo8870. Materials, methods, experimental details, and additional characterisation data (NMR, SEC, cryo-TEM, AFM).

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Notes

The authors declare no competing financial interests.

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