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Bulk and surface morphologies of ABC miktoarm star terpolymers comprised of PDMS, PI and PMMA arms

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Abstract

DIM miktoarm star copolymers, comprised of polydimethylsiloxane [D], poly(1,4-isoprene) [I] and poly(methyl methacrylate) [M], were synthesized using a newly developed linking methodology with 4-allyl-1,1-diphenylethylene as a linking agent. The equilibrium bulk morphologies of the DIM stars were found to range from [6.6.6] tiling patterns to alternating lamellar and alternating cylindrical morphologies, as determined experimentally by small angle X-ray scattering and transmission electron microscopy and confirmed by dissipative particle dynamics and self-consistent field arguments. The thin film morphologies, which differ from those found in the bulk, were identified by scanning electron microscopy, coupled with oxygen plasma etching. Square arrays of the PDMS nanodots and empty core cylinders were formed on silica after oxygen plasma removal of the poly(1,4-isoprene) and poly(methyl methacrylate) which generated nanostructured substrates decorated with these features readily observable.

Keywords

Polydimethylsiloxane (PDMS), poly(1,4-isoprene) [PI], poly(methyl methacrylate) [PMMA], anionic polymerization, ABC miktoarm star, Flory-Huggins interaction parameter, surface patterning, surface structuring, surface lithography, plasma etching, Archimedean tiling pattern.
Introduction

Etch-resistant patterns formed by certain types of block copolymers, for example, block copolymers containing Si, Sn and/or Fe atoms in one of the blocks, can be used to produce nanopatterned graphene, silicon or polymeric substrates that have applications in semiconductor, water purification and solar cell industries.\textsuperscript{1-4} Lithographic potential of linear di-/triblock copolymers\textsuperscript{5-12} have been exhaustively studied to this end, while there are relatively few reports\textsuperscript{13-15} on the use of ABC miktoarm star terpolymers, which have a much richer diversity of morphologies and nanopatterns that can be produced.

The complexity of the architecture of the terpolymer, in comparison to linear di-, tri-, or multiblock copolymers, comes at the cost of a more detailed synthesis and purification. ABC miktoarm star terpolymers are typically produced by anionic polymerization of each arm of the star, followed by consecutively connecting the arms to a difunctional linking agent. In a typical example, lithium $p$-dimethylsilanolate functionalized 1,1-diphenylethylene was used as the initiator to polymerize a PDMS (polydimethylsiloxane) arm from the lithium silanolate, followed by the addition of a second poly(styryl)lithium arm to the free 1,1-diphenylethylene group and further growth of PTBMA [poly(tert-butyl methacrylate)] from the middle of the resultant PDMS-PS (polystyrene) diblock copolymer.\textsuperscript{16,17} In another example, the consecutive addition of the living PI (polyisoprene), PS and PDMS to methyltrichlorosilane produced a miktoarm star terpolymer.\textsuperscript{18} More recently, a trifunctional capping agent, benzaldehyde with two alkynyl units, was utilized for the synthesis of PS-PI-PFS (ferrocenylethylmethylsilane) star terpolymers.\textsuperscript{19} In the present work we further exploit the area of Si containing ABC miktoarm stars by elaborating on DIM miktoarm stars comprised of PDMS, PI and PMMA (poly(methyl methacrylate)) arms.

Dissipative particle dynamics (DPD) and self-consistent field theory (SCFT) have been used to tackle the complexity of morphologies exhibited by ABC miktoarm stars theoretically.\textsuperscript{20-26} Modeling of ABC terpolymers poses a challenge, due to the large number of parameters required, in comparison to diblock copolymer counterparts.\textsuperscript{26-42} For diblock copolymers, at least two parameters are needed to simulate the morphology: volume fraction of block A, $f_A$, and interaction parameter, $\chi_{AB}N$, where
\( \chi_{AB} \) is the pairwise Flory-Huggins interaction parameter between segment A and B and \( N \) is the total number of Kuhn segments. However, for ABC terpolymers, at least five parameters are required: \( f_A \), \( f_B \), \( \chi_{AB}N \), \( \chi_{BC}N \) and \( \chi_{AC}N \). Some of the modeling studies focused on benchmarking the model phase diagram results against the experimental results in which molecular weight of one of the block was varied, while keeping lengths of the other two arms fixed.\(^29,31,32\) Other groups also followed it up by construction of the full triangular phase diagrams in which the block fractions form the sides of the triangle.\(^30,37,43\) For simplicity, constant interaction parameters have been used though the molecular weights of the blocks were changed. Two different cases of interaction parameters could be identified: 
Case I: all the interaction parameters are the same,\(^27,28,32\) and Case II: two of the interaction parameters are equal and one of them is larger or smaller than the other two.\(^26,29\)

In the idealized Case I, where \( \chi_{AB} = \chi_{AC} = \chi_{BC} \), a sequence of ordered patterns were predicted to form as the composition is altered under the constraint that the A and B arms are kept equal in size, while C is varied. In the balanced case, i.e. where all arms occupy equal volumes, a three-colored hexagonal honeycomb is the equilibrium morphology, as shown in Figure 1. For \( 0.5 < x < 2.5 \) (\( x = C/A = C/B \)) a number of columnar structures with tiling-pattern cross-section are predicted, while for larger \( x \) a hierarchical, alternating lamellar morphology [ALT. LAM] should form (\( 2 < x < 10 \)) and for even larger \( x \) a hierarchical, alternating cylindrical morphology [ALT. CYL] is predicted. In this case, hexagonally packed cylinders with alternating layers of A and B are embedded in a matrix of C. Based on these simulations we would expect a [6.6.6] morphology for \( x \sim 1 \); either [10.6.4; 10.6.6] or [12.6.4] or [ALT. LAM] for \( x \sim 2 \); [ALT. LAM] for \( x \sim 5 \); and [ALT. CYL] for \( x \sim 13 \). Experimentally, the exact location of the phase boundaries as a function of \( x \) will most likely not match Figure 1 precisely, since they are molecular weight dependent\(^44\), to some extent, and, more importantly, the pair-wise interaction parameters are not symmetric, as assumed in the simulations.

Relative incompatibilities between the chains comprised of PDMS-PI-PMMA arms of the miktoarm star terpolymers used in the present work could be assessed from the difference in solubility parameters (\( \delta \)) of the individual homopolymers. The PMMA (\( \delta = 18.9 \text{ MPa}^{1/2} \)) and PDMS (\( \delta = 15.1 \text{ MPa}^{1/2} \)) are the most incompatible blocks, PMMA and PI (\( \delta = 16.4 \text{ MPa}^{1/2} \)) are moderately
incompatible, while PDMS and PI are the least incompatible components. The solubility parameters were used to calculate interaction parameters for performing coarse grained DPD simulations to probe the effect of the asymmetric interactions.

Figure 1. Generic phase diagram based on simulation predictions under the constraint of two arms (green A and blue B) having equal volumes while the third arm (red C) is varied. The parameter $x$ is the fractional length of C relative to A. The tiling patterns are labeled by a set of numbers $[k_1,k_2,k_3]$ indicating that three polygons ($k_1$-gon, $k_2$-gon, $k_3$-gon) meet at every vertex and repeat in a cyclic order. Tilings with more than one topologically distinct vertex are denoted $[k_1,k_2,k_3; k_4,k_5,k_6]$.44

Case II, where $\chi_{AB} \approx \chi_{AC} > \chi_{BC}$ or $\chi_{AB} \approx \chi_{AC} < \chi_{BC}$, qualitatively captures the morphologies of the poly(styrene) [S], poly(isoprene) [I] and poly(2-vinyl pyridine) [V] and PS-PI-PDMS (SID) miktoarm terpolymers. For the former, the interaction parameters are not balanced because $\chi_{SI} \approx \chi_{SV} < \chi_{IV}$ and for the latter, $\chi_{DS} \approx \chi_{DI} > \chi_{SI}$. The SID stars having symmetric composition formed a cross-shaped morphology resembling St. Andrew’s cross which we designate as the [4.8.8] tiling. This morphology is due to high incompatibility between D and the other two arms leading to the tendency of S and I domains to increase their mutual contact area to reduce the less favorable contacts with D so as to minimize the free energy of the system. Changes in the incompatibility among different monomers has been shown to have significant effect on the morphologies in the case of linear terpolymers. By keeping the composition of PI-$b$-PS-$b$-PMMA fixed and varying only the interaction parameter by partial hydrogenation of the PI block different morphology - lamellar or network - was observed, depending on the degree of hydrogenation of PI block.

In this manuscript the bulk morphologies of DIM miktoarm star terpolymers are presented, followed by a discussion of the morphologies in thin films. Both experimental and computational results will be presented for the bulk systems, while for thin films, only experimental characterization was realized, since, computationally, the number of parameters required to quantitatively describe the morphologies is prohibitive. Specifically, lack of information about the film thickness relative to the
like-domain repeat periods and the absolute as well as relative interactions of each block with the substrate and air interfaces makes even qualitative comparisons between simulations and experiments prohibitive. Two of the four systems used for the bulk characterization were used for the thin film studies. The thin film morphologies were found to be different from the bulk, underscoring the importance of surface effects in defining the morphologies.

**Materials and methods**

All chemicals were purchased from Sigma-Aldrich unless otherwise stated. Tetrahydrofuran (THF) was distilled from ketyl radical of benzophenone under argon. Cyclohexane was distilled from living poly(styryl)lithium under argon. Hexamethycyclotrisiloxane (D₃) was consecutively sublimed from calcium hydride (CaH₂) and di-n-butylmagnesium. Isoprene was consecutively distilled from CaH₂ and di-n-butylmagnesium. Methyl methacrylate (MMA) was consecutively distilled from CaH₂ and triethylaluminum (Et₃Al).

The molecular weight was determined by gel permeation chromatography (GPC) using THF with 1% triethylamine as an eluent at 0.5 ml/min flow rate with a column set consisting of a pre-column and two 300 x 8 mm main columns (PLgel Mixed C and Mixed D). For transmission electron microscopy and SAXS ~ 0.5 mm polymer films were produced by solvent casting from 7% THF solutions under N₂ for 1 week in the dark, followed by a thermal annealing for 5 days at 150 °C, staining with 4% OsO₄ aqueous solution for 4 h, microtoming at room temperature to 70 nm thick sections, and an additional staining with 4% OsO₄ aqueous solution on copper grids for 4 h. The TEM was performed on FEI Tecnai T20 G² at 200 kV accelerating voltage in a bright field mode. The SAXS was measured using a Ganesha SAXS-LAB nm with Cu K radiation (λ = 0.154 nm). The sample-to-detector distance was 1041 mm, and the X-ray beam area was 0.04 mm². Thin films were spin coated from 1-2% toluene solutions at speeds ranging from 2-6 krpm and maximum acceleration (6 krpm/s) to achieve thicknesses close to the equilibrium domain spacing (d₀), followed by annealing in saturated acetone vapors for 20 h. The SEM of the microphases on silica substrates was realized
4-Bromo-1,1-diphenylethylene (DPE-Br). As shown on Scheme 1, bromobenzene (24 g, 0.228 mol) was converted to Grignard reagent by reacting it with magnesium turnings (5.7 g, 0.234 mol) in 600 ml of dry diethyl ether followed by slow addition of 4-bromoacetophenone (42 g, 0.211 mol). The reaction mixture was stirred for 0.5 h at room temperature and neutralized by 1.5 L of saturated NH₄Cl aqueous solution. After extraction with diethyl ether organic phase was washed with saturated NH₄Cl and water, then dried and concentrated. Azeotropic dehydration using Dean-Stark apparatus in toluene in the presence of catalytic amount of p-toluenesulfonic acid afforded a crude product which was further concentrated and distilled under reduced pressure, yielding 50 g (91%) of transparent liquid (b.p. 105 °C at 0.025 mbar). 1H NMR (CDCl₃, 400 MHz, in ppm): 7.50 (d, J = 8.6 Hz, 2H), 7.37 (m, 5H), 7.26 (d, J = 8.5 Hz, 2H), 5.51 (dd, J = 8.1, 1.0 Hz, 2H).

4-Allyl-1,1-diphenylethylene (DPE-All). DPE-Br (30 g, 0.116 mol) was converted to Grignard reagent by reacting it with magnesium turnings (3.1 g, 0.128 mol) in dry THF resulting in a dark brown solution to which allyl bromide (11.0 ml, 1.127 mol) was added via syringe leading to immediate reaction as judged by the disappearance of color and the boiling of the mixture. The mixture was warmed to 65 °C for 10 min and cooled to room temperature. Standard work-up consisting of diethyl ether extraction and repetitive water washings followed by concentration on a rotary evaporator and fractional distillation afforded 12 g (43%) of DPE-All as a transparent liquid (b.p. 100 °C at 0.09 mbar). 1H NMR (CDCl₃, 400 MHz, in ppm): 7.39 (m, 9H), 6.06 (ddt, J = 16.8, 10.1, 6.7 Hz, 1H), 5.50 (dd, J = 9.7, 1.3 Hz, 2H), 5.16 (m, 2H), 3.48 (d, J = 6.7 Hz, 2H). Elemental analysis: theory for C₁₇H₁₆ 92.68% C, 7.32% H, found 92.53% C, 7.34% H. Mass spectrometry: m/z theory for C₁₇H₁₆ [M⁺] 220.1, found 220.4. The compound (DPE-All) must be stored at -20 °C since at 4 °C slow decomposition was evidenced by a yellow discoloration.

Anionic polymerization. The experiments were realized using an anionic polymerization set-up described elsewhere. In a typical example (Scheme 1), D₃ was polymerized using sec-butyllithium as an initiator in THF at -5 °C for 16 h plus 6 days at -20 °C followed by end capping with 3 times...
excess of chlorodimethylsilane. The PDMS-H produced was reacted with DPE-All by a hydrosilylation using Karstedt's catalyst in toluene at 40 °C, affording functionalized PDMS-DPE macromonomer. In order to link the second poly(1,4-isoprene) [PI] arm to the macromonomer, isoprene was added to the mixture of sec-butyllithium initiator and cyclohexane in 1L reactor and stirred for 10 h at 40 °C (overpressure!). Such nonpolar polymerization conditions yield poly(isoprene) characterized by 75% cis-1,4, 20% trans-1,4 and 5% 3,4 microstructure. The solution of living poly(1,4-isoprenyl)-Li (1 eq.) was transferred to the second reactor containing excess of dried PDMS-DPE macromonomer (2 eq.) and LiCl (5 eq.) in dry THF resulting in the development of an intense red color. After 2 h at -45 °C the reaction mixture was cooled down to -78 °C and a predetermined amount of MMA was added. The polymerization of the MMA monomer was conducted for 1 h at -78 °C and the resulting product was precipitated into excess of methanol. The purification was effected by extraction of an excess of unreacted PDMS-DPE with hexane and/or fractionation in toluene/methanol mixtures. The product was dried under vacuum at 50 °C for 16 h at 0.001 mbar.

Dissipative particle dynamics. The simulations were performed as described previously, except for the values of the interaction parameters between simulation beads building up the chains.
The coarse-graining takes a balanced $x = 1$ star as reference which is represented with 3 beads along each chain all attached to a central neutral junction bead. The low number is needed to be able to reach $x = 13$ which needs 39 beads along one chain. From the solubility parameters $\delta_i$ and $\delta_j$ one can calculate the $\chi_{ij}$ parameter between species $i$ and $j$ as

$$\chi_{ij} = \frac{v}{k_B T} (\delta_i - \delta_j)^2$$  \hspace{1cm} (1)

where $k_B T$ is the thermal energy and $v$ is the average volume of the two species involved as they are represented in the simulations which is dependent on the chosen coarse-graining. The calculated $\chi_{ij}$ parameters are then converted into a DPD-interaction strength $\alpha_{ij}$ between species $i$ and $j$ controlling a soft potential as described in the references above:

$$\alpha_{ij} = 25 + 3.497 \chi_{ij}$$  \hspace{1cm} (2)

We get the following values for the interaction parameters: $\alpha_{DI} = 35$, $\alpha_{IM} = 55$ and $\alpha_{DM} = 91$. We will discuss the validity of these values based on this simple solubility parameter approach below when comparing simulation results with the experimental structure determination.

**Self-Consistent Field Theory (SCFT).** The SCFT of ABC miktoarm star terpolymer melts has been described previously. Briefly, the Hamiltonian of the ABC miktoarm melt was defined followed by standard field theoretical transformations leading to a field theoretical representation. Standard saddle-point approximation leads to a set of non-linear SCFT equations, which were then solved iteratively to obtain the spatial distribution of the volume fraction of the polymer segments. The modified diffusion equations appearing at the saddle-point were solved using pseudo-spectral method. Polyswift ++ provides an efficient implementation of the SCFT of ABC miktoarm star terpolymer melts and was used for all the SCFT calculations in this work. Other details for evolving the SCFT equations are presented in Ref. The Hamiltonian of the ABC miktoarm polymer is briefly described here and the details are described in Ref. Three Gaussian chains, each of length $N_i l_i$, are defined, where $i$ is the index of the block ($i = 1, 2, 3$ represents D, I and M, respectively), $N_i$ and $l_i$ stand for the number and length of a Kuhn segment in $i$th block, respectively. The polymer segments interact with dissimilar polymer segments by an effective short-range mean field term defined as $\chi_{ij}$ where $i$ varies between 1 and 2 and $j$ varies between 2 and 3 ($j >
All the $l_i$'s are assumed to be the same ($l = 0.51$ nm), where $l = \nu^{1/3}$ and $\nu$ is the geometric mean of the molar volume of all three monomers. $N_i$ of each polymer is obtained by equating the coarse grain model contour length against atomistic contour length.

$$N_i = (2DP - 1)b/l$$

Where $DP$ is the degree of polymerization, defined as polymer molecular weight/monomer molecular weight, $b$ is the atomistic backbone bond length ($b = C-C$ bond length = 0.154 nm for PI and PMMA, and $b = Si-O$ bond length = 0.163 nm for PDMS. All the length scales are scaled with respect to $R_g = l(N/6)^{1/2}$, where $N = \sum_{i=1}^{3} N_i$. For all calculations, chain fraction $f_i = N_i/N$ is assumed to be equal to the volume fraction and $N$ is used to estimate the interaction parameter, $\chi_{ij}N$. ABC miktoarm polymer melts are reported to change morphologies over a narrow range of volume fractions, especially around symmetric composition, which justifies the choice of volume fraction for modeling the system.\textsuperscript{29,31} The pairwise interaction parameters were obtained from the studies reported in literature: $\chi_{DM}$ is reported to be $\sim 0.19$ at 150 °C, the annealing temperature used in our work,\textsuperscript{12} $\chi_{IM}$ it is reported to be $\sim 0.08$ at 50 °C\textsuperscript{69} (assumed to be the same at 150 °C here) and $\chi_{ID}$ is reported to be $\sim 0.08$ based on solubility parameter estimation,\textsuperscript{70} but was found to better fit our observations if a value of 0.0554 was used. If a $\chi_{ID}$ value of 0.08 is chosen instead of 0.0554 then [4.8.8] tiling pattern tends to have lower free energy than the experimentally observed [6.6.6] tiling. Zone annealing was used to equilibrate the system.\textsuperscript{42,71} The length and interaction parameters are given in Table 1.

Simulation box parameters for the SCFT calculations are as follows: $\Delta$ is the uniform grid size and $N_xN_yN_z$ are the number of grids in the x, y, z directions, respectively. For DIM-0.8, DIM-2.0 and DIM-4.9, calculations were performed in two-dimension (2D) whereas three-dimensional (3D) calculation was performed for DIM-13. For the 2D calculations, $\Delta = 0.15 R_g$, $N_x = N_y = 64$, and $N_z = 1$ and for the 3D calculations, $\Delta = 0.30 R_g$ and $N_x = N_y = N_z = 32$ were used.

<table>
<thead>
<tr>
<th></th>
<th>$f_D$</th>
<th>$f_I$</th>
<th>$f_M$</th>
<th>$N$</th>
<th>$\chi_{ID}N$</th>
<th>$\chi_{IM}N$</th>
<th>$\chi_{DM}N$</th>
</tr>
</thead>
<tbody>
<tr>
<td>DIM-0.8</td>
<td>0.36</td>
<td>0.38</td>
<td>0.26</td>
<td>368</td>
<td>20.4</td>
<td>29.4</td>
<td>69.9</td>
</tr>
<tr>
<td>DIM-2.0</td>
<td>0.24</td>
<td>0.26</td>
<td>0.49</td>
<td>520</td>
<td>28.8</td>
<td>41.6</td>
<td>98.8</td>
</tr>
<tr>
<td>DIM-4.9</td>
<td>0.14</td>
<td>0.15</td>
<td>0.71</td>
<td>878</td>
<td>48.6</td>
<td>70.2</td>
<td>166.7</td>
</tr>
<tr>
<td>DIM-13</td>
<td>0.07</td>
<td>0.07</td>
<td>0.86</td>
<td>1819</td>
<td>100.8</td>
<td>145.5</td>
<td>345.5</td>
</tr>
</tbody>
</table>
Static Structure Factor from SCFT. Discrete Fourier transform was used to obtain the static structure factor from the volumetric mean of the electron scattering length density (SLD), \( \rho(x, y, z) \), to qualitatively model the SAXS experiments, where \( \rho(x, y, z) = \sum_{i=1}^{3} SLD_i \phi_i(x, y, z) \) and \( SLD_i \) and \( \phi_i \) represent the SLD and volume fraction of the component \( i \) at equilibrium, respectively (\( z = 0 \) for 2D calculations). The SLD (in \( 10^6 \text{Å}^{-2} \)) for D, I and M components are 8.8, 8.6 and 10.8, respectively.\(^{58}\) The FFTW library\(^{72}\) was used to perform Fourier transformation of \( \rho(x, y, z) \):

\[
\Omega(q_x, q_y, q_z) = \text{FT}\{ \rho(x, y, z) \},
\]

where \( q_x = \frac{2 \pi l}{L_x}, q_y = \frac{2 \pi m}{L_y} \) and \( q_z = \frac{2 \pi n}{L_z} \) are the scattering wave vector components in the \( x, y, z \) directions, respectively and \( l, m, \) and \( n \) are integers ranging from 0 to \( N_x/2, N_y/2 \) and \( N_z/2 \), respectively. The static structure factor \( S(q^*) \) is obtained as a function of \( q^* = (q_x^2 + q_y^2 + q_z^2)^{1/2} \) using \( S(q^*) = \frac{\langle \Omega(q^*) \Omega(-q^*) \rangle}{(N_xN_yN_z)} \) where \( \Omega(q^*) \) and \( \Omega(-q^*) \) represents the Fourier transform of \( \rho(x, y, z) \) and its complex conjugate, respectively (\( q^* \) is a dimensionless vector). The angular brackets \( <> \) represent the averaging over the Fourier space.

Results and Discussion

Synthesis. The synthesis of PDMS containing miktoarm stars was realized by developing a new protocol relying on a novel double functional capping agent, 4-allyl-1,1-diphenylethylene (DPE-All). The capping agent was successfully synthesized and immobilized onto hydride terminated PDMS through a hydrosilylation reaction facilitated by Karstedt's catalyst. In general, the hydrosilylation reaction is sensitive to the substitution of an alkene: terminal monosubstituted alkenes were found to be the most reactive, while internal and 1,1-gem double bonds are less prone to react.\(^{73,74}\) Even selective hydrosilylation of monosubstituted alkenes in the presence of internal alkenes was reported.\(^{75}\) Capitalizing on the high regio-selectivity of the hydrosilylation process we reacted the allyl group of DPE-All with PDMS-H while retaining 1,1-diphenylethylene functionality. The progress of the reaction could be conveniently monitored by the disappearance of a heptet at 4.70 ppm (RMe_2Si-H) and appearance of a doublet of doublets at 5.42 ppm originating from the olefinic
DPE protons in the modified PDMS. The PDMS-DPE macromonomer produced was further reacted with living poly(isoprenyl)-Li at -45 °C. Attempts to conduct the reaction at lower temperatures were not successful, due to the reduced reactivity of poly(isoprenyl)-Li. In the final step, a 5x molar excess of LiCl relative to the initiator was used to reduce the nucleophilicity of the growing PMMA-Li centers, which would otherwise be active enough towards the carbonyl groups of the monomer/polymer. As a result, well-defined miktoarm star block terpolymers were obtained (Table 2 and Figure 2).

Table 2. Characteristics of the synthesized star block terpolymers.

<table>
<thead>
<tr>
<th>Name</th>
<th>MW, kDa (NMR)</th>
<th>PDI (GPC)</th>
<th>Dx : Dy : Mz (NMR)</th>
<th>Morphology (SAXS)</th>
<th>$d_0$ (SAXS), nm</th>
</tr>
</thead>
<tbody>
<tr>
<td>D</td>
<td>16.1</td>
<td>1.07</td>
<td>1 : 0 : 0</td>
<td>n/a</td>
<td>n/a</td>
</tr>
<tr>
<td>I</td>
<td>16.4</td>
<td>1.03</td>
<td>0 : 1.1 : 0</td>
<td>n/a</td>
<td>n/a</td>
</tr>
<tr>
<td>DI</td>
<td>32.5</td>
<td>1.05</td>
<td>1 : 1.1 : 0</td>
<td>n/a</td>
<td>n/a</td>
</tr>
<tr>
<td>DIM-0.8</td>
<td>46.5</td>
<td>1.08</td>
<td>1 : 1.1 : 0.8</td>
<td>HEX</td>
<td>28.9</td>
</tr>
<tr>
<td>DIM-2.0</td>
<td>71.7</td>
<td>1.06</td>
<td>1 : 1.1 : 2.0</td>
<td>LAM</td>
<td>40.8</td>
</tr>
<tr>
<td>DIM-4.9</td>
<td>131</td>
<td>1.14</td>
<td>1 : 1.1 : 4.9</td>
<td>LAM</td>
<td>54.6</td>
</tr>
<tr>
<td>DIM-13</td>
<td>287</td>
<td>1.07</td>
<td>1 : 1.1 : 13.0</td>
<td>HEX</td>
<td>56.0</td>
</tr>
</tbody>
</table>

$^a$Volume ratios (v:v:v) of the blocks were estimated by $^1$H NMR (CDCl$_3$, 400 MHz) using densities of PDMS (0.965 g/ml), poly(1,4-isoprene) (0.91 g/ml), PMMA (1.18 g/ml) and the following NMR peaks (in ppm): PDMS (-CH$_3$) 0.07, PMMA (-O-CH$_3$) 3.60 and PI (olefinic H) 4.68, 4.76, 5.12. $^b$Hexagonally packed cylinders (HEX) and lamellar (LAM) morphologies were determined from SAXS relative peak positions. The equilibrium domain spacing ($d_0 = 2\pi/q^*$) was calculated from SAXS.

Figure 2. GPC curves of the synthesized DIM stars and their precursors.

**TEM.** The analysis of the bulk morphologies by TEM (Figure 3) indicate that sample DIM-0.8 forms well-ordered [6.6.6] tiling patterns in agreement with the volume fractions of the corresponding DIM arms (1:1.1:0.8, v:v:v). Since all samples were exposed to OsO$_4$ vapor, the PI component was selectively stained and appears black, while PMMA and PDMS, even though unstained, still have sufficient natural contrast and appear grey and white respectively. A slightly higher PI content relative
to PDMS and PMMA in DIM-0.8 results in visually larger black domains in the cross-section compared to grey and white domains (Figure 3A). A similar [6.6.6] symmetry was observed for ISV stars comprised of polyisoprene (I), polystyrene (S) and poly(2-vinylpyridine) (V) having almost identical volume ratios (1:1:1:0.7, v:v:v) but a three times higher total molecular weight (46.5 vs. 134 kDa). The morphology of DIM-2.0 and DIM-4.9 is represented by alternating lamellar [ALT.

LAM] structure, which consists of PMMA lamellae separated by combination of PI and PDMS lamellae. Since PI and PDMS domains resemble cylinders, the structure is sometimes referred to as Lamellar+Cylinder. Such a morphology was observed for ISV (1:1:3, v:v:v) and PS-PI-PFS (1:1.4:4.7, v:v:v) miktoarm star terpolymers, and was predicted theoretically for ideal ABC star terpolymers over a broad composition range (1:1:x, 2.5 ≤ x ≤ 6.5). The equilibrium bulk morphology of DIM-13 is best described by the cylinders made up of stacked disks of alternating PI and PDMS in a matrix of PMMA. This morphology is referred to as a columnar piled disk or lamellar-within-cylindrical structure in the literature and designated as alternating cylinders [ALT.

CYL] in the present work.

Figure 3. (Top 2 rows) TEM images of OsO4 stained DIM-0.8 (A), DIM-2.0 (B), DIM-4.9 (C) and DIM-13 (D) with (bottom row) the corresponding computer generated 3D bulk morphologies. Color code: D – blue, I – green, M - red
**SAXS.** The morphologies of DIM miktoarm star terpolymers in the bulk were also investigated using SAXS after thermal annealing (Figure 4). The calculated electron densities ($10^{23} \text{electrons/cm}^3$) for PDMS (3.13), PI (3.06) and PMMA (3.83) indicate that the contrast between PDMS and PI components is low and these microdomains will be difficult to distinguish by SAXS. Therefore, the main contribution to scattering profiles in Figure 4, arises from the size and shape of the PMMA microdomains and the spatial correlations of this microdomains. For DIM-0.8, the six reflections at scattering vector ($q$) ratios of $1:\sqrt{3} :4 :\sqrt{7} :\sqrt{13} :\sqrt{16}$ relative to the first order peak ($q^*$) were observed, characterized by $d_0 = 2\pi/q^* = 28.9 \text{ nm (}\quad q^* = 0.217 \text{ nm}^{-1}\quad)$, indicating that the PMMA arrange in hexagonally packed cylindrical [HEX] microdomains with a unit cell size $a = 2d_0/\sqrt{3} = 33.4 \text{ nm}$. This is consistent with the [6.6.6] tiling seen in the TEM images and with our DPD modeling as well as SCFT SAXS predictions.\textsuperscript{26}

With an increase in PMMA molecular weight, the scattering profile indicates that the PMMA microdomains are lamellar [LAM], in keeping with the TEM and DPD results. For DIM-2.0, the lamellae have a $d_0$ of 40.8 nm ($q^* = 0.154 \text{ nm}^{-1}$). For DIM-4.9 the $d_0$ of the [LAM] microdomains increases to 54.6 nm. Again, in keeping with the DPD predictions and the TEM results, the [LAM] transforms into [HEX] cylindrical microdomains for the largest molar mass of PMMA, DIM-13, although the peaks are less pronounced than for the low molecular weight sample. The two low $q$ peaks can be indexed as the $1:\sqrt{3}$ peaks of the hexagonal peak sequence leading to a $d_0 = 56.0 \text{ nm (}\quad q^* = 0.112 \text{ nm}^{-1}\quad)$ giving a unit cell size of 64.7 nm. From these results, it follows that the morphology of the DIM miktoarm terpolymers changes from [HEX] to [LAM] and then back to a [HEX] structure due to an increase in the PMMA molecular weight which causes a progressively higher degree of compositional asymmetry. These results are in agreement with the modelling data depicted on Figure 1, despite the inherent simplifications in the simulations.
Figure 4. SAXS intensity profiles as a function of scattering vector ($q$) for DIM miktoarm terpolymers. The samples were prepared using solvent casting and thermally annealed at 150 °C under vacuum for 5 days. The intensity profiles were vertically shifted according to the molecular weight of PMMA. The black arrows are $q/q^*$ values while the red arrows show DI correlation peak.

**SCFT Modeling.** The SCFT morphologies of the DIM miktoarm polymer melts are shown in Figure 5. Overall, the morphologies agree with the experimental micrographs shown in Figure 3 with the only exception being the case of DIM-13 where concentric cylinders, instead of stacked alternative cylinders of PI and PDMS, are predicted. As per the parameters listed in Table 1, the repulsion between PDMS and PMMA are far stronger in comparison to other pairs, which results in the preference, in the simulations, for an inner cylindrical microdomain of PDMS surrounded by PI which shields it from nonfavorable interactions with PMMA (i.e., a core-shell morphology). One possible origin for this discrepancy is that the interactions between the PDMS and PMMA are mediated by the presence of the solvent during the preparation of the TEM samples, decreasing the non-favorable interactions between these segments. Evidence for this is seen in the DPD simulations where the [ALT. CYL] morphology for DIM-13 was observed using symmetrical interaction parameters only. When the interaction parameters were estimated from solubility parameters, the DPD simulations returned core-shell morphology for DIM-13, in agreement with SCFT predictions.
The static structure factors obtained for different compositions of DIM are shown in Figure 6. The DIM-0.8 shows relative peaks at $1 : \sqrt{3} : \sqrt{4} : \sqrt{7}$, which qualitatively agrees with the peaks observed in the experimental SAXS data (Figure 4) and with the SCFT predictions by others.\textsuperscript{26} The first peak is split into two sub-peaks in Figure 6a, which is attributed to the imperfect match between the computational lattice parameters and the period of the ordered phases.\textsuperscript{26} The DIM-2.0 and DIM-4.9 exhibit peaks at $1 : 2 : 3$ positions corresponding to the lamellar spacing of PMMA as well as smaller peak designated as $1^*$ on Figure 6b-c. Since the electron contrast between PMMA and the other two components is much higher compared to the contrast between PDMS and PI, relatively small peak is expected for the lamella formed by PDMS and PI domains. The peaks spaced as $1 : 2 : 3$ are observed experimentally by SAXS for DIM-2.0 and DIM-4.9 (Figure 4) but the predicted $1^*$ peak was not detected. This is attributed to the low intensity of the $1^*$ peak which makes it difficult to resolve such feature experimentally. Finally, for DIM-13, the modelled peaks, $1 : \sqrt{3} : \sqrt{4} : \sqrt{9} : \sqrt{13}$, agree with $1 : \sqrt{3}$ experimental ones.
We should point out that the results presented in Figure 5 and in turn, Figure 6, were obtained by solving the SCFT equations after starting from the field configurations initialized with random numbers and by ramping up the Flory’s chi parameters gradually to the values presented in Table 1. We have compared free energies of the morphologies obtained with initial configurations generated by different seeds for the random number generator and in some cases, with the inclusion of conformational asymmetry between the blocks in an *ad hoc* manner. These comparisons of the free energies have revealed that the sample DIM-2.0 can have multiple metastable structures with their free energies quite close to each other, in agreement with previous reports in the literature. All of the other samples exhibited clear free energy minima. The pseudo-spectral method used to solve the SCFT equations is quite useful for exploring different morphologies without any *a priori* knowledge of their symmetry but cannot be used to resolve infinitesimal differences in the free energies in a reliable manner. Figure 5 reveals that indeed a minimal model constructed here while ignoring the effects of conformational asymmetries and for fixed values of the Flory’s chi parameters taken from the literature can reproduce the experimental results. We plan to present many details of the SCFT based modeling of ABC star terpolymers along with the effects of the conformational asymmetries in a forthcoming publication.

**Thin film morphologies.** In the thin film regime the morphologies are influenced by the interactions between the blocks and the two interfaces: substrate (silica) and free surface (air). The arms with low surface tension – PDMS ($\gamma_{PDMS} = 21$ mN/m) and PI ($\gamma_{PI} = 31$ mN/m) – would tend to preferentially segregate to the polymer/air interface, while PMMA ($\gamma_{PMMA} = 41$ mN/m), the most polar of the blocks, would preferentially segregate to the substrate (the thin silicon oxide layer coating the silicon substrate). As a consequence, the equilibrium morphology for DIM-2.0 transforms from [ALT. LAM] in the bulk to a [4.8.8] tiling pattern in thin films (compare Figures 3B and 7A). Since the thickness of the films (40 nm) was close to the equilibrium domain spacing ($d_0 = 40.8$ nm) and the wetting is antisymmetric, a monolayer of DIM stars was achieved with all components present at
the air interface, resulting in a relatively non-frustrated morphology oriented orthogonal to the film surface.\textsuperscript{79}

Another important effect originates from the confined movement of the ABC junction points which are allowed to migrate only in one dimension compared to the two dimensional movement of connecting points in AB block copolymers.\textsuperscript{49,80} This, coupled with the connectivity of the three blocks precludes the formation of the PDMS wetting layer on the top of the microphase separated thin film, which is well-recognized in PDMS containing block copolymer films.\textsuperscript{12,81} As a consequence, the surface morphology could be directly imaged by SEM without the need of short CF\textsubscript{4} plasma etching to remove the PDMS top brush layer (Figure 7A). Moreover, all three components could be distinguished by direct observation with SEM (Figure 7A inset): white (PDMS), grey (PMMA) and PI (dark).

In another example, in contrast to [ALT. LAM] in the bulk, core shell cylinders appear on silica substrate for DIM-4.9 (Figures 3C and 7B). In this scenario, rather than forming lamellae, as in the bulk, PMMA forms a matrix (Figure 7G) thereby maximizing its interactions with the substrate. The existence of all patterns was confirmed by selective removal of PMMA and PI by isotropic oxygen plasma etching (50W, 0.3 mbar, 5 min). Upon such treatment PDMS is converted into silicon oxycarbide forming characteristic patterns resistive to further oxygen plasma etching (Figure 7 C, D, F, H). Both patterns identified in this work, namely, the square array of equally spaced silicon oxycarbide domains (pitch = 25 nm) and the free standing cylindrical shells of oxidized PDMS (pitch = 53 nm), represent unique surface morphologies unattainable by means of diblock copolymer lithography. Needless to say, the abovementioned patterns could further be transferred to underlying silica substrate by CF\textsubscript{4} plasma exposure or used as prepared for a range of applications where nanopatterned substrates are required.\textsuperscript{82}

As for the samples DIM-0.8 and DIM-13, our attempts to observe their surface patterns were not successful for two possible reasons. In the case of DIM-0.8 the film de-wets the substrate immediately after spin coating from toluene solution. Presumably, the low PMMA content in this sample precludes effective adsorption of the DIM-0.8 onto the silicon oxide. For DIM-13, despite the fact that uniform
films were always produced via spin coating, reduced chain mobility at such high molecular weight (287 kDa) may account for the inability of the copolymer to form well-defined surface morphologies.

Figure 7. SEM images of the thin films of DIM-2.0 (A, C, E, F) and DIM-4.9 (B, D, G, H) after SVA in acetone (A, B) followed by oxygen plasma etching (C, D, F, H). Bottom row represents computer generated images of the corresponding morphologies before (E, G) and after oxygen plasma etching (F, H). Color code: D – blue, I – green, M - red

Conclusions

A series of DIM miktoarm stars consisting of polydimethylsiloxane [D], poly(1,4-isoprene) [I] and poly(methyl methacrylate) [M] arms were synthesized by anionic polymerization using a novel linking agent, 4-allyl-1,1-diphenylethylene. The bulk morphology of DIM was investigated as a function of the volume ratio of M/D = M/I = 0.8-13 while the ratio of D/I was fixed at ~ 1. A systematic increase in the molecular weight of the M arm resulted in the morphological transition from [6.6.6] cylindrical tiling to [ALT. LAM] and back to [ALT. CYL] in the bulk. The real space morphologies were determined using TEM, DPD and SCFT, while SAXS and Fourier transform of
SCFT results were used to access the scattering profiles of the polymers. The presence of the PDMS component resistant to oxygen plasma etching allowed us to produce regular square array of oxidized PDMS dots (pitch = 25 nm) and empty core cylinders (pitch = 53 nm) on silica – the patterns which were drastically different from the observed bulk morphologies. The ability to form regular etch resistant patterns on silica wafers makes synthesized DIM interesting candidates for nanoscopic lithography applications.  

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Bulk and surface morphologies of ABC miktoarm star terpolymers comprised of PDMS, PI and PMMA arms

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