Architecting polymersomes from spherical to tubules by molecular alterations

Abstract

Polymersomes formed by the self-assembling of block copolymers gained huge attraction recently due to it numerous potential applications especially as nano-reactors, tunable controlled delivery carriers, templates for bio-mineralization etc. The self-assembly of these block copolymers amphiphiles into monodisperse structures in aqueous solutions is influenced by their hydrophobic and hydrophilic molecular regions. However the commonly investigated structures are spherical vesicles, micelles and worm like micelles. This study focuses on generating tubular vesicles or “Tubularosomes” by altering the amphiphilic block length of the polymer chain. Polyoxazoline (POx) is covalently conjugated with polydimethyl siloxane (PDMS) to form block copolymers and explored the self-assembling behavior with special focus on the influence of chain length forming various structures. Additionally, the current limitations in the reproducibility and biocompatibility during the synthesis and utilization of biologically important polymer vesicles are explored along with enabling the development of these nanostructures towards real-world applications.

Keywords: Polymersomes, Tubularosomes, Vesicles, Packing parameter

Introduction

There is a constant demand for well-defined block-copolymers with reproducible molecular characteristics and polydispersity, which are a prerequisite for various applications including controlled biological delivery systems, hydrogels, catalysis etc. “Polymersomes” or polymer vesicles are self-assembled structures formed from block copolymers. Compared to traditional liposomes, polymersomes provide robustness and enhanced stability both in terms of chemical and physical attributes and hence have gained popularity for numerous potential.6-14 The hydrophilic and hydrophobic properties of the diblock-copolymers makes polymersomes a potential competitor for various nano-medical systems.15-17 The hydrophilic corona can be utilized to encapsulate water soluble agents while the hydrophobic part can have various proteins embedded. The size and structure of polymersomes plays a crucial role in implementing functions.9 The amphiphilic can self-assemble into simple morphologies like spherical micelles, simple vesicles, onion ring shaped vesicles, rods or more complex structures like toroid, compound micelles with inverted core shell structures.18 Morphology of the block-copolymers is controlled by different forces such as interactive forces between hydrophobic and hydrophilic blocks, degree of stretching of core blocks and repulsive interaction among corona chains.19,20 One of the major parameter affecting the self-assembly of block-copolymers can be explained in terms of classical dimensionless “packing parameter” or “Pc”.13,19,21 Pc is mathematically represented as: P = v/a1, where v is the volume of hydrophobic chain, a1 is area occupied by hydrophilic head group and l1 is the length of molecule. With P values below 0.5, the self-assemblies are in form of spheres, cylinders or worm like micelles, whereas with increasing P values until 1, bilayers are formed. Factors such as solution pH, temperature, type of organic solvent, initial copolymer concentration, chemical structure of copolymer, presence of additives and hydrophilic/hydrophobic ratio determine the packing factor.14-17 When the hydrophilic to hydrophobic ratio is greater than 1:1 the self-assembly is usually micelles. Ratios less than 1:2 usually favor vesicles while that less than 1:3 may form complex structures like inverted micro-structures along with vesicles.19,22 Changing the block length of hydrophilic or hydrophobic chain in the copolymers may lead to switching between morphologies or make the copolymer sensitive to stimuli like pH, temperature and solvent polarity.20 This helps in controlled release phenomena as well as in sensing applications. In earlier works by Choe et al.23 the amphiphilic block co-polypeptide was tuned to achieve different hydrophobic lengths. Varying hydrophobic chain length led to monodisperse vesicles with low toxicity and better stability. Yingchao Chen et al.24 blended amphiphilic block copolymers poly(acrylic acid)-block-poly-isoprene and poly(acrylic acid)-block-polystyrene to yield various self-assembly from spherical to rod like micelles. Extreme slow or medium rate of water addition to the co-polymer blends resulted in mixture of vesicle and rod like conformations. In similar study by Yoshida25 spherical vesicles produced only by poly(methacrylic acid)-block-poly(methyl methacrylate-random-methacrylic acid) copolymer were changed into worm-like vesicles with an uneven surface upon long poly(methacrylic acid) copolymer incorporation. In the last 20 years polymer vesicles with spherical structure morphology has been the focus of extensive research. However, other interesting structures like cylindrical or rod like shape have been gaining interest particularly due to their potential application in the field of medicine and nanotechnology.26 Due its typical structure and flexibility, cylindrical vesicles or micelles can stretch and orient themselves for flow-intensive systems such as phage-mimetic or micro-pore drug delivery systems.27-29 With spherical structures being thermodynamically stable, the cylindrical conformations were produced by blending degradable polyactic acid with inert block copolymer.26 This morphological...
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The triblock copolymers formed worm-like tubular to micro-phase separated regimes of hydrophobic POx domains and chemical structure were studied via transmission electron microscopy (TEM) and dynamic light scattering (DLS) (Figure 1).

**Materials and methods**

POX-b-PDMS-b-POX was synthesized via cationic ring opening polymerization of 2-ethyl oxazoline. The preliminary step had monomer 1,3-bis(4-hydroxybutyl)tetramethyldisiloxane initiated with one of the Brönsted acid, trifluoromethanesulfonic acid anhydride to form a living polymer chain. The length of POX block was adjusted by addition of varying the amount of fresh distilled 2-ethyl-2-oxazoline. Finally, the polymerization was terminated by addition of 0.5 M KOH in ethanol. For this study the hydrophobic PDMS block was maintained at 5600 g mol$^{-1}$ while the molecular weight of the hydrophilic end group POX was varied from 1088 to 1396 g mol$^{-1}$. For synthesis of POX-b-PDMS-b-POX, the PDMS block was maintained at 5600 g mol$^{-1}$ while the hydrophilic POX end group was kept at 1500 g mol$^{-1}$. Synthesis of PMOXA was accomplished using a propargyl tosylate initiator. PDMS was converted into tosylate end groups TsO-PDMS-OTs. This tosylation made it easier to be transformed into azides N$^3$-PDMS-N via nucleophilic substitution. Finally, copper-catalyzed alkyn-azide cycloaddition (CuAAC) click reaction was utilized to achieve high yield PMOXA-b-PDMS-b-PMOXA.

Both polymersomes and tubularsomes were prepared by popular “film rehydration method”. Film rehydration technique involved dissolving the block copolymers in organic solvent like chloroform with subsequent film formation by evaporation of the solvent using a rotavaporator. The film was further placed under high vacuum overnight to remove any traces of organic solvent. Consequently Milli-Q water was added to the dry film followed by continuous stirring for 4 hours to rehydrate the film. The suspension obtained was then extruded through two polycarbonate membrane filters (Whatman nucleopore membrane) with nominal diameter of 400 nm held in an extruder using 200 psi nitrogen gas. The number of times the sample was extruded was maintained at 10 passes to reach a uniform size distribution. The scanning electron micrograph in TEM mode was acquired on Hitachi S4800 field emission microscope operating at 30 kV and 20 µA. The diluted suspension of vesicles were dropped on carbon-coated TEM grids followed by removal of excess using Kim wipes. One drop of staining solution 2% uranyl acetate was placed on the grid for 30 seconds which was also removed by Kim wipes. The TEM grids were placed under vacuum for 1 hour to completely remove the solvent after which they were analyzed under microscope. Dynamic light scattering (DLS) study was carried out on Malvern Nano ZS zetasizer at room temperature with back scattering angle 0$^\circ$ = 173$^\circ$.

**Results and discussion**

Figures 2&3 shows the TEM images of tubularsomes of various POX-b-PDMS-b-POX chain length block copolymers and confirm that the major population of tubular morphology with dark regions corresponding to uranyl acetate staining of PDMS domains and lighter areas corresponding to hydrophilic POX segments. The individual tubular vesicle formed from POX-b-PDMS-b-POX had a diameter of 36 nm and length of 50–450 nm. The hydrodynamic diameters obtained from DLS, Figure 5, for POX-b-PDMS-b-POX were 100–500 nm and 110–600 nm for POX-b-PDMS-b-POX. As illustrated in Figure 4&5, the vesicle formed from PMOXA-b-PDMS-b-PMOXA triblock was spherical in structure with an average diameter of 250 nm and hydrodynamic diameter of 150–250 nm. The triblock copolymers formed worm like tubular to spherical structures upon varying the experimental parameters. The morphology change depended on the hydrophilic weight fraction, suggesting a strong effect of the POX block length as well as change of POX into PMOXA on self-assembly of the vesicles. Due to the amphiphilic nature of POX-b-PDMS-b-POX having both hydrophobic and hydrophilic domains, the volume fractions in aqueous solution formed a micro-phase separated regime of the polymer domains which self-assembled into lamellae structures. The triblock copolymer in the films showed only limited micro-phase separation and on contact with water or buffer solution resulted in micro-phase separation into hexagonally packed rods, with further hydration leading to the formation of lamellae. The spontaneous curvature formation of the triblock copolymer occurs when the effective hydrophilic area increases. This change in morphology is critical while designing vesicles for applications such as long tubular vesicles provide much needed surface area for encapsulation of drugs and easier transport through fine blood vessels (Figure 2-5).

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In summary, changing the hydrophilic chain length of the block copolymer constitutes an efficient and elegant way of controlling the stability, morphology and size of vesicles formation. The spontaneous formation of vesicles was simply achieved by changing the block length of same appropriately chosen hydrophilic material. Longer the hydrophilic chain, more pronounced spherical is structure of vesicles. This important research work helps us to tailor make the vesicles based on the applications in pharmacy, drug-delivery and cosmetics industry.

Conclusion

In summary, changing the hydrophilic chain length of the block copolymer constitutes an efficient and elegant way of controlling the stability, morphology and size of vesicles formation. The spontaneous formation of vesicles was simply achieved by changing the block length of same appropriately chosen hydrophilic material. Longer the hydrophilic chain, more pronounced spherical is structure of vesicles. This important research work helps us to tailor make the vesicles based on the applications in pharmacy, drug-delivery and cosmetics industry.

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Conflict of interest

There is no conflict of interest.

References


