Effects of crystallinity and dispersity on the self-assembly behavior of block co-oligomers in water†

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Self-assembly of block copolymers in solution is a topic of great interest in polymer science due to the potential for applications as a drug carrier system. In bulk, fully discrete polymers have been shown to self-assemble in extremely well-defined structures, but the effect of full discreteness on self-assembly in solution is less known. Furthermore, little is known about the effect of molar mass dispersity on crystallization driven self-assembly. Here, we investigate both the effects of dispersity and crystallinity on the self-assembly behavior of low molecular weight poly(lactic acid)-poly(ethylene glycol) block co-oligomers (BCOs) in solution. The results show that the introduction of dispersity and/or crystallinity does not significantly affect spherical and cylindrical morphologies, but vesicular structures are affected. The introduction of dispersity in amorphous vesicle forming BCOs lowers the reproducibility of preparations in solution. For crystalline BCOs, the introduction of dispersity leads to a clear decrease of ordering in bulk and it prevents crystallization of the LLA block in solution. This all arises already at a low dispersity (Đ ≤ 1.06), highlighting the effect of dispersity on assemblies of low MW BCOs. It also underlines the need to take dispersity into account when aiming for homogeneous well-defined structures in solution.

Introduction

Block copolymers (BCPs) are an exciting class of macromolecules that have seized the attention of many polymer chemists. Applications range from the development of drug delivery vehicles in aqueous media to nanolithography in bulk.1–5 Although controlled polymerizations have optimized BCPs, the presence of dispersity is an intrinsic property, both at the level of their molar mass (molar mass dispersity, D) and at the level of compositional dispersity. Natural polymers such as DNA and peptides are discrete with a D of 1.00 and are sequence-defined. However, achieving such controlled uniformity in synthetic polymers has been a great challenge in polymer science.6,7 Fully discrete low molecular weight oligomers with a D of 1.00 have been obtained via iterative synthesis methods,8 using solid-phase9–12 or solution based approaches.13–18 or by combining controlled polymerization techniques with flash chromatography.19–21 In addition, discrete block co-oligomers (BCOs) have been prepared by these methods, affording systems that at low molecular weights form highly-ordered, phase-segregated structures in bulk.22–27 The introduction of dispersity in these low molecular weight BCOs has a dramatic effect on the nanophase separation and long range ordering.28–32

In solution the assembly of block copolymers is driven by the difference of lyophobicity between the blocks. Upon solvation in a selective solvent, the lyophobic blocks will rearrange to minimize their contact with the solvent, while the lyophilic blocks will remain solvated. This leads to a segregated structure, in which both a dense lyophobic core, consisting of collapsed chains, and a swollen lyophilic corona are present.33,34 The final morphology in thermodynamic equilibrium is determined by the volume ratio of the lyophobic and the lyophilic blocks according to the theory of Israelachvili.35 In case of disperse lyophobic and lyophilic blocks, the volume ratios throughout the whole population of polymer chains might vary, causing a mixture of morphologies in the final assembly.36 Especially on smaller aggregates of less than 1000 molecules this is a serious drawback, as all aggregates
will have different compositions and thus different morphologies. This effect of dispersity and resulting mixture of morphologies has been shown for ABA and AB type block copolymers. Increasing dispersity in either the core or corona forming blocks shifted the morphology from vesicles towards a mixture of vesicles, spheres and worm-like assemblies, or from vesicles to spheres.37–39

In addition to dispersity, crystallinity has shown to be a driving force for morphology change. Crystallization-driven self-assembly of amphiphilic block copolymers has been used to obtain non-spherical morphologies in solution, by using a crystallizable core-forming block. The crystallinity of the core-forming block can be used to switch the morphology from the predicted spherical morphology into cylinders, as the crystalline core of these micelles can act as nuclei after which epitaxial growth can occur.40–44

In previous work of our group, the effect of dispersity on the crystallinity in self-assembled low molecular weight ABA type BCOs was studied in water.45 Similar to the observations in bulk, the introduction of dispersity showed a significant effect on the homogeneity of the obtained particle morphologies in water. To explore the effect of lack of dispersity in these low molecular BCOs, a library of AB type BCOs was synthesized with full control over sequence and molar mass distributions. To this end, previously used,46 but compare discrete and disperse, as well as the same blocks (oligo(lactic acid) and oligo(ethylene glycol)) we have selected a ratio of the two blocks that ensures the formation of vesicles, spheres and amorphous lactic acid blocks. To this end, we investigated AB type BCOs consisting out of the hydrophobic block, and discrete oligo(ethylene glycol) 11-, 17- and 48-mers were used as hydrophilic blocks, to obtain a variety of morphologies upon self-assembly of the BCOs in water. For bilayer morphologies it is known that they can adopt flat, curved, or closed vesicular structures in solution, depending on the ability of the hydrophobic block to bend into the closed vesicular structure. An excellent agreement between theoretically predicted size and morphology was found for all these discrete crystalline BCOs, but it remained unclear to what extent crystallinity and dispersity played a role in the formation of stable and reproducible vesicular structures.

To continue our previous work, we here like to answer the following question: what has a greater influence on the morphology formed by a low MW BCO, crystallinity or dispersity? Therefore, we investigate AB type BCOs consisting out of the same blocks (oligo(lactic acid) and oligo(ethylene glycol)) we previously used,46 but compare discrete and disperse, as well as crystalline and amorphous lactic acid blocks. To this end, we have selected a ratio of the two blocks that ensures the formation of vesicles (LA16EG11). These vesicle forming BCOs were characterized in bulk and in aqueous solution using a combination of scattering, differential scanning calorimetry (DSC) and total internal reflection fluorescence microscopy (TIRF). To shed light on the effect of dispersity and crystallinity on other morphologies, sets of LA16EG17 (cylinders) and LA16EG48 (spherical micelles) were also studied. The packing of the LA16 block in these assemblies in water was analyzed using small angle neutron scattering (SANS) and by assessing the solubilization region of the hydrophobic dye Nile Red (NR).

Results and discussion
Design and synthesis of BCOs
We prepared oligo(lactic acid) of 16 repeat units from L-lactide (LLA16), and selected commercially available oligo(ethylene glycol) of 11 repeat units (EG11). Dispersity is introduced into the LA block (discrete LLA16 versus disperse LLA16) where the dispersity is indicated by using the tilde symbol to study the effect of dispersity on crystalline BCOs (LLA16EG11 and LLA16EG11). To understand the effect of dispersity without crystallinity present, we synthesized a set of amorphous BCOs starting from the racemic mixture of L- and L-lactide nLLA16 and introduced dispersity in the LA block (nLLA16). The synthesis of discrete TBDMS-LLA16-COOH and TBDMS-dLLA16-COOH is based on the previously reported synthetic strategy developed by Hawker’s group,13 and later slightly modified by our group.22 Subsequent ligation with commercially available discrete MeO-EG11-OH resulted in discrete BCOs nLLA16EG11 and dLLA16EG11 (Fig. 1a). For nLLA16EG11 ring-opening polymerization (ROP) of L-lactide was performed using MeO-EG11-OH as initiator. For dLLA16EG11 first a TBDMS-nLLA16-COOH precursor was synthesized via ROP of nLLA16 using benzyl alcohol as initiator and subsequently coupled to MeO-EG11-OH. The TBDMS group was not removed to increase the stability of the BCOs.47,48

All compounds were purified by automated column chromatography and fully analyzed by 1H NMR, 13C NMR and matrix-assisted laser desorption/ionization mass spectrometry (MALDI-ToF) mass spectrometry (Fig. S1–S28†). Full synthetic details on the preparation of the BCOs can be found in the ESI.†

While 1H-NMR spectra of all four compounds look identical and give similar degrees of polymerization, MALDI-ToF spectra reveal large differences between the discrete and disperse compounds (Fig. 1b). nLLA16EG11 has a narrower distribution than dLLA16EG11 due to a column purification in the synthetic procedure, but the range of D is in both cases low enough to

Fig. 1 (a) Chemical structures of studied BCOs: crystalline discrete LLA16EG11, crystalline disperse LLA16EG11, amorphous discrete nLLA16EG11 and amorphous disperse dLLA16EG11. (b) MALDI-ToF spectra of all BCOs.
not expect pronounced effects of the difference in dispersity. \( \text{lLA}_{16}\text{EG}_{11} \) and \( \text{dLLA}_{16}\text{EG}_{11} \) show only a single peak in MALDI-ToF-MS, underlining the discreteness of these BCOs, while a distribution of peaks is observed for disperse BCOs \( \text{lLA}_{16}\text{EG}_{11} \) and \( \text{dLLA}_{16}\text{EG}_{11} \). We note that the values for the molar mass dispersity, \( D \), are narrow, and range between 1.01–1.06 for the disperse compounds. These disperse oligomers are thus a good reference to gain insight into what extent dispersity matters.

In addition to the series LA\(_{16}\text{EG}_{11} \) anticipated to form lamellar structures in water, we also synthesized the series LA\(_{16}\text{EG}_{17} \) (cylindrical morphologies) and LA\(_{16}\text{EG}_{48} \) (micellar morphologies) with discrete/disperse and/or amorphous/crystalline lactic acid blocks. The synthetic details and characterization of these BCOs are given in Schemes S1–S3 and Fig. S1–S29.

### Bulk properties of BCOs

Crystalline oligomers \( \text{lLA}_{16}\text{EG}_{11} \) and \( \text{lLA}_{16}\text{EG}_{17} \) were obtained as waxy solids at room temperature, while amorphous \( \text{dLLA}_{16}\text{EG}_{11} \) and \( \text{dLLA}_{16}\text{EG}_{17} \) were obtained as viscous oils. Their thermal behavior was investigated using DSC (Table 1, Fig. S29†). For the amorphous BCOs \( \text{dLLA}_{16}\text{EG}_{11} \) and \( \text{dLLA}_{16}\text{EG}_{17} \) the only visible transition is a glass transition temperature, \( T_g \). For both crystalline BCOs \( \text{lLA}_{16}\text{EG}_{11} \) and \( \text{lLA}_{16}\text{EG}_{17} \), a clear crystallization transition was observed upon cooling. The enthalpy of crystallization of \( \text{lLA}_{16}\text{EG}_{11} \) is lower than for \( \text{dLLA}_{16}\text{EG}_{11} \), suggesting that the crystalline packing is less defined for the disperse variant. However, disperse \( \text{lLA}_{16}\text{EG}_{17} \) crystallizes at a higher temperature than its discrete counterpart \( \text{lLA}_{16}\text{EG}_{11} \) (38 °C vs. 22 °C), indicating that the LA chains with DP > 16 in the disperse BCO nucleate crystallization at a higher temperature. Furthermore, by introducing dispersity into the \( \text{lLA}_{16} \) block, melting occurs over a broad temperature range. This is in contrast to the sharp melting transition that was observed for the discrete variant (Fig. S29†). These observations are in line with those previously observed in BCOs comprising oligodimethylsiloxanes and \( \text{LLA} \).30

The packing and long-range ordering of the oligomers in bulk at room temperature was further investigated with X-ray scattering experiments. The discrete crystalline oligomer \( \text{lLA}_{16}\text{EG}_{11} \) shows a sharp principal scattering peak and higher order Bragg reflections, indicating a highly structured long-range packing of the chains (Table 1, Fig. 2). The ratios of the Bragg reflections (\( \sqrt{4}, \sqrt{9}, \sqrt{16}, \sqrt{25} \)) indicate a lamellar phase (LAM) with an interlayer distance of 8 nm. Furthermore, multiple scattering peaks can be observed in the WAXS region, corresponding to the inter-chain packing of the LA block, typical for the crystalline packing of lactic acid chains.49 Introducing dispersity in the crystalline oligomer \( \text{lLA}_{16}\text{EG}_{11} \) gives rise to broadening of the primary scattering peak indicating a less ordered structure and smaller crystalline domains. In addition, the maximum of the first-order reflection which arise from the lamellar packing, \( q^* \), shifts to smaller \( q \)-values, corresponding to longer interlayer distances on the order of 10 nm. This is likely caused by the increasing amount of \( \text{LLA} \) blocks with DP > 16. At higher \( q \) values we can still see the peaks typical for crystalline packing of the lactic acid chain, at the same values as for the discrete block variant. These crystalline features probably arise from the longer lactic acid chains with DP > 16, as the shorter chains are not able to crystallize on short timescales.30 For the amorphous BCOs, \( \text{dLLA}_{16}\text{EG}_{11} \) and \( \text{dLLA}_{16}\text{EG}_{17} \), the absence of scattering peaks corroborates the thermal data, as no crystallization for the \( \text{dLLA}_{16} \) chains is visible.

### Table 1: Appearance, thermal properties and phase behavior of the BCOs

| Oligomer          | \( D \) | Appearance\(^a\) | \( T_g \) [°C] | \( T_c \) [°C] | \( \Delta H \) [kJ mol\(^{-1}\)] | Phase\(^b\) | \( d^* \) \(^c\) [nm] |
|------------------|--------|------------------|---------------|--------------|----------------------------|-----------|------------------|
| \( \text{lLA}_{16}\text{EG}_{11} \) | 1.00   | Wax              | —             | 22           | 32                         | LAM       | 8.0              |
| \( \text{lLA}_{16}\text{EG}_{17} \) | 1.06   | Wax              | —             | 38           | 22                         | LAM       | 10.1             |
| \( \text{dLLA}_{16}\text{EG}_{11} \) | 1.00   | Viscous oil      | —33           | —            | —                          | DIS       | —                |
| \( \text{dLLA}_{16}\text{EG}_{17} \) | 1.01   | Viscous oil      | —19           | —            | —                          | DIS       | —                |

\(^a\) Physical appearance at room temperature, directly after drying in vacuo. \(^b\) Bulk morphology determined with SAXS at room temperature. \(^c\) Domain spacing, calculated as \( d^* = 2\pi/q^* \). LAM = lamellar, DIS = disordered.
Self-assembly of BCOs in water

To study the effect of dispersity without any crystallinity present, we first self-assembled the amorphous DLLA$_{16}$EG$_{11}$ and DLLA$_{\sim}$$_{16}$EG$_{11}$ in water. A slow solvent switch was used, as our previous work showed that dropwise addition of water to a THF stock solution leads to the predicted thermodynamically stable vesicular structures. After self-assembly in water, multi-angle light scattering was used to measure the sizes of the morphologies formed and to obtain information on the morphology of the self-assembled structures. The results show that the hydrodynamic radius ($R_H$) is comparable for both DLLA$_{16}$EG$_{11}$ and DLLA$_{\sim}$$_{16}$EG$_{11}$ (Fig. 3a), and a log–log representation of the corresponding static light scattering data indicate that for both BCOs a vesicular structure is adapted, as $I \propto q^{-2}$ (Fig. 3b). These results indicate that introducing dispersity does not lead to differences in the nature of the particles formed by the amorphous BCOs. To visualize the structures in solution formed by DLLA$_{16}$EG$_{11}$ and DLLA$_{\sim}$$_{16}$EG$_{11}$, the particles were analyzed with TIRF microscopy after addition of Nile Red (Fig. S30†). Nile Red (NR) is a hydrophobic solvatochromic dye, which accumulates into hydrophobic domains where it fluoresces. Spherical, non-interacting particles were observed for both discrete and disperse BCOs, corroborating the results obtained from scattering techniques. Overall, the introduction of dispersity into amorphous BCOs does not lead to morphology differences upon particle formation in water. However, the reproducibility of sample preparation is affected by the presence of dispersity (Fig. 3c). After 1 week, the sizes obtained from discrete DLLA$_{16}$EG$_{11}$ are comparable, while for several preparations of DLLA$_{\sim}$$_{16}$EG$_{11}$ the sizes vary. Even though the difference in dispersity (1.00 vs. 1.01) is small, the consequences for a reproducible sample preparation are significant.

Our previous work showed that it was possible to obtain spherical vesicular structures from crystalline tLA$_{16}$EG$_{11}$ by using a slow solvent switch. Important to note here, is that sample preparation and history play an important role in obtaining these vesicular structures. Over time, the BCO crystallizes in bulk, which reduces the solubility of this compound and leads to irreproducible sample preparations in water. Rigorous dissolution in the appropriate solvent and carefully following the dissolution overtime prior to sample preparation in water are required for the sample preparation procedure to be reproducible.

Upon self-assembly of disperse tLA$_{\sim}$$_{16}$EG$_{11}$ in water using the slow solvent switch, particles with a spherical shape were obtained (Fig. 4a). To assess the capability of disperse tLA chains to crystallize in solution, tLA$_{\sim}$$_{16}$EG$_{11}$ was assembled at higher concentrations and subjected to micro-DSC. In contrast to tLA$_{16}$EG$_{11}$, no transitions were visible during the heating run (Fig. 4b). Thus, in solution, the disperse nature of the tLA chains prevents crystallization.

To further investigate the packing of the LA chains in the assembled structures, SANS experiments were performed for the four BCOs in water (Fig. 5a, Table 2). For tLA$_{16}$EG$_{11}$, we previously showed that the small angle X-ray scattering data could be fitted as a flat homogenous lamellar structure with a layer thickness of 5.9 nm, and thus 11.8 nm for the total bilayer. This indicates that the tLA$_{16}$ chains in the lamellar structures may be packed in a sort of intercalating or collapsed fashion (Fig. 5b), as the lamellar domain spacing of one fully extended tLA$_{16}$ chain is 6.0 nm.

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**Fig. 3** Light scattering results of the amorphous BCOs upon self-assembly in water. (a) $I$ vs. $q^2$ plot to determine the $R_H$ of the spherical bilayer structures. (b) Scattering intensity $I$ vs. $q$ to probe the shape of the self-assembled structures. (c) Obtained $R_H$ of multiple sample preparations.

**Fig. 4** (a) TIRF image of disperse tLA$_{\sim}$$_{16}$EG$_{11}$ upon self-assembly in water. (b) Micro-DSC traces of tLA$_{16}$EG$_{11}$ and tLA$_{\sim}$$_{16}$EG$_{11}$ at 5 mg mL$^{-1}$ after self-assembly in water.
In contrast, the patterns obtained with SANS here were shifted to longer distances. The scattering data could only be described as lamellae if polydispersity in the distribution of the hydrophobic bilayer was considered. Fitting the neutron scattering data with a head/tail lamellar structure (see ESI for details†), gives a bilayer thickness of 14.8 nm for particles formed by tLA16EG11, corresponding with a hydrophobic tail length of the tLA block of 4.3 nm. This larger tail length compared to previously published results likely arises from a different extent of crystallinity present in these structures. Crystalline regions are likely to pack tail-to-tail (Fig. 5b) rather than the intercalating arrangement of more amorphous samples. This would lead to an overall larger hydrophobic tail length (Fig. 5b). As mentioned previously, over time part of the bulk material crystallizes, and it is possible that sample history of the bulk BCOs was not fully removed before making the samples in water. A competing process between self-assembly and crystallization of the aged BCOs upon self-assembly in water can occur, resulting in a mixture of aggregates with a different extent of crystallinity and thus different sizes than observed before. For the scattering profiles in solution, the effect of chain length dispersity was not so pronounced: the scattering data with a head/tail lamellar structure (see ESI for details†), gives a bilayer thickness of 14.8 nm for particles formed by tLA16EG11, corresponding with a hydrophobic tail length of the tLA block of 4.3 nm. This larger tail length compared to previously published results likely arises from a different extent of crystallinity present in these structures. Crystalline regions are likely to pack tail-to-tail (Fig. 5b) rather than the intercalating arrangement of more amorphous samples. This would lead to an overall larger hydrophobic tail length (Fig. 5b). As mentioned previously, over time part of the bulk material crystallizes, and it is possible that sample history of the bulk BCOs was not fully removed before making the samples in water. A competing process between self-assembly and crystallization of the aged BCOs upon self-assembly in water can occur, resulting in a mixture of aggregates with a different extent of crystallinity and thus different sizes than observed before. For the scattering profiles in solution, the effect of chain length dispersity was not so pronounced: the scattering pattern and fit results were similar for tLA16EG11 and tLA16EG11, with a smaller bilayer thickness of 12.2 nm for disperse tLA16EG11.

The effect of (lack of) crystallinity, however, was clearly visible. For both amorphous tLA16EG11 and tLA16EG11 the fit results gave a smaller bilayer thickness of 10.6 nm. The hydrophobic tail length of the LA block was 2.3 nm, roughly half the value of the hydrophobic tail length of the crystalline bilayer in tLA16EG11, indicating a more intercalating type of packing of the tLA tails. This, together with the crystallization of tLA16EG11 in solution, implies that the ability of the hydrophobic block in tLA16EG11 to crystallize gives rise to a different type of packing in the obtained vesicular structures.

To further assess the nature of the packing of the LA block, we applied the dye NR as a probe. It was recently reported that the emission wavelength, $\lambda_{\text{max,em}}$, of this dye can be used to assess whether the dye is located in the hydrophobic part of the bilayer or in the corona region after solubilization in self-assembled structures. When comparing the $\lambda_{\text{max,em}}$ of NR mixed into tLA16EG11 or tLA16EG11 (Fig. 5c), a clear difference is observed. The $\lambda_{\text{max,em}}$ is at higher wavelengths for the particles prepared from crystalline tLA16EG11 than for the particles prepared from amorphous tLA16EG11 (~620 nm versus ~605 nm). This indicates that due to the crystallinity of the tLA block, the dye is not able to enter the hydrophobic bilayer and is more located towards the bilayer-corona interface.

Influence of dispersity and crystallinity on other morphologies

The combined effects of dispersity and crystallinity in the BCO series tLA16EG11 do not have a significant effect on the type of morphologies formed, as in all cases vesicular structures are formed in water. However, we found pronounced differences in the reliability of sample preparation and on the bilayer thickness of the vesicular structures in water when dispersity and crystallinity were introduced. The question then is, in how far is this effect dependent on morphology? Therefore, we investigated BCOs based on tLA16EG17 and tLA16EG48, which previously were shown to form cylindrical and spherical micelles, respectively. The dissolution process was carefully followed over time to obtain a reproducible sample preparation procedure as these samples are also sensitive to crystallinity and sample history.

For the tLA16EG17 series, a slow solvent switch by addition of water to the organic stock solution using a syringe pump was used to obtain the particles in solution. SANS measurements and theoretical calculations have predicted that this BCO com-

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**Table 2** SANS scattering results of tLA16EG11 vesicle forming BCOs in water

| Oligomer      | $D$  | Bilayer thickness $^a$ [nm] | Hydrophobic tail length $^a$ [nm] |
|---------------|------|---------------------------|----------------------------------|
| tLA16EG11     | 1.000| 14.8                      | 4.3                              |
| tLA16EG11     | 1.06 | 12.2                      | 3.0                              |
| tLA16EG11     | 1.000| 10.6                      | 2.3                              |
| tLA16EG11     | 1.01 | 10.6                      | 2.3                              |

$^a$ See Fig. 5b. $^b$ Previous SAXS results showed that tLA16EG11 forms vesicular structures with a bilayer thickness of 11.8 nm.}

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**Fig. 5** (a) SANS scattering profiles and corresponding fits in D$_2$O, 1 mg mL$^{-1}$. Data is shifted vertically for clarity. (b) Schematic representation of possible bilayer structures in solution, tLA16 in red, EG11 in blue. (c) Normalized emission spectra of the dye NR in self-assembled tLA16EG11 and tLA16EG11 in water.
position has the tendency to aggregate into cylindrical micelles. The theoretical models, however, do not account for sample crystallinity, which may be detrimental to self-assembly. SANS profiles for the BCOs of the LA₁₆EG₁₇ series (crystalline, non-crystalline, discrete and disperse) were all characterized by a decay at low q with a slope value of −1, suggesting the presence of rod-like aggregates. Therefore, we speculate that the crystallinity does not significantly influence the morphology of the cylindrical micelles (Fig. 6a). Following this assumption, the data was fit using a core–shell cylinder model. Similar to the LLA₁₆EG₁₁ system, the crystalline variants of LLA₁₆EG₁₇ have a slightly larger cross-sectional radius compared to the non-crystalline ones (Table 3). The packing of the molecules in the cylindrical aggregate may restrict the crystallization of the LA blocks. It is noteworthy that the maximum concentration that could be achieved with this sample preparation was too low to perform reliable micro-DSC experiments. Therefore, it is unclear whether the LA block can, in fact, crystallize in the core of the cylindrical micelles.

Finally, we measured the emission spectra of dye NR in the presence of all four BCOs (Fig. 6b), which are all very similar. Interestingly, the $\lambda_{\text{max,em}}$ is close to that observed for the lamellar morphologies formed by LLA₁₆EG₁₁. This indicates that the dye is not able to enter the core and is more located towards the core–corona interface. The fact that all emission spectra overlay, suggests that the LA block forms a densely packed core, regardless of dispersity and/or crystallinity.

In the case of the series LA₁₆EG₄₈, a fast solvent switch by quick injection of organic stock solution into water was used to obtain self-assembled structures, similar to previously published procedures. The SANS profile overlay (Fig. 7a), which indicates that neither the presence/absence of crystallinity, nor the presence/absence of dispersity significantly affects the size of the structures formed. In addition, the emission spectra of the dye NR are near identical for all BCOs, pointing to a highly similar solubilization region of dye NR (Fig. 7b). The $\lambda_{\text{max,em}}$ of approximately 620 nm indicates a tightly packed core, as this lies close to the value obtained for assemblies of LLA₁₆EG₁₁. Taken together, these results reveal that for BCOs predicted to form spherical micelles, introduction of dispersity or crystallinity does not lead to noteworthy differences in terms of packing of the core LA block.

**Conclusion**

We successfully synthesized a set of amphiphilic low molecular weight block co-oligomers. By preparing $\alpha$-lactic and $\delta$-lactic acid oligomers in an iterative manner and subsequently ligating to discrete ethylene glycol blocks, fully discrete diblock co-oligomers were obtained. Additionally, by introducing dispersity in crystalline and amorphous lactic acid oligomers, the effect of dispersity with and without crystallinity present was studied. By ligating a hydrophobic LA₁₆ block with EG₁₁, the formation of vesicular structures in solution was ensured. In bulk, discrete crystalline LLA₁₆EG₁₁ formed highly ordered phase separated structures, but upon introduction of dispersity in the LA block, this ordering was clearly diminished even though the disperse counterpart has a $D$ value of only 1.06. In solution, both amorphous BCOs, δLLA₁₆EG₁₁ and δLLA₁₆EG₁₁, formed the expected spherical vesicular structure. Particles made from disperse δLLA₁₆EG₁₁ showed a lower reproducibility in terms of size, which is quite remarkable as the $D$ value is only 1.01, highlighting the effect that dispersity can have on these low molecular weight oligomers in solution. Introducing crystallinity in the BCOs gave rise to a different type of core packing in the obtained lamellar structures. For crystalline BCOs the introduction of dispersity affected the packing of the LA block, as the LA chains in LLA₁₆EG₁₁ do not show a melting transition upon self-assembly in water and SANS results indicated a smaller bilayer thickness for LLA₁₆EG₁₁.

**Table 3** SANS scattering results of LA₁₆EG₁₇ cylindrical micelle forming BCOs using a core–shell cylinder model

| Oligomer        | $D$ | Cross-sectional radius [nm] | Core radius [nm] |
|-----------------|-----|-------------------------------|-----------------|
| LLA₁₆EG₁₇       | 1.00| 6.5                           | 2.5             |
| LLA₁₆EG₁₇       | 1.04| 6.5                           | 2.5             |
| $\alpha$LLA₁₆EG₁₇ | 1.00| 5.0                           | 1.0             |
| $\delta$LLA₁₆EG₁₇ | 1.01| 5.0                           | 1.0             |

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combined effects of dispersity and crystallinity were not evident. Only for vesicle forming low MW BCOs, introducing dispersity in crystalline compounds has a noticeable effect on packing of the hydrophobic block, both in bulk and in solution. As the difference in dispersity for this set of BCOs is so small, this highlights the importance of taking dispersity into account in low molecular weight systems.

Conflicts of interest

There are no conflicts to declare.

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