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Enhanced Self-Assembly and Mechanical Properties of Cellulose-Based Triblock Copolymers: Comparisons with Amylose-Based Triblock Copolymers

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ABSTRACT

Herein, we compared the microphase-separation behavior and mechanical properties of cellulose- and amylose-based block copolymers (BCPs). Various celooligosaccharide triacetate-*b*-poly(δ -decanolactone)-*b*-celooligosaccharide triacetates (AcCel_{*n*}-*b*-PDL-*b*-AcCel_{*n*}), which are cellulose-based ABA-type BCPs, with PDL molecular weights of approximately 5, 10, and 20 kg mol⁻¹ and PDL volume fractions of 0.65, 0.77, and 0.87, were synthesized from α,ω -diazido-end-functionalized PDLs and propargyl-end-functionalized celooligosaccharide triacetates via click chemistry. We adopted the cellodextrin-phosphorylase-mediated oligomerization of α -D-glucose-1-phosphate in the presence of a propargyl-end-functionalized cellobiose primer to synthesize the functional celooligosaccharide segment. The maltooligosaccharide triacetate-*b*-poly(δ -decanolactone)-*b*-maltooligosaccharide triacetate (AcMal_{*n*}-*b*-PDL-*b*-AcMal_{*n*}) amylose counterparts were also synthesized in a similar manner. Small-angle X-ray scattering experiments and atomic force microscopy revealed that the AcCel_{*n*}-*b*-PDL-*b*-AcCel_{*n*} are more likely to microphase-separate into ordered nanostructures compared to the AcMal_{*n*}-*b*-PDL-*b*-AcMal_{*n*}, despite their comparable chemical compositions and molecular weights. Furthermore, the AcCel_{*n*}-*b*-PDL-*b*-AcCel_{*n*} exhibited significantly superior mechanical performance compared to their amylose counterparts under tensile testing, with the Young's modulus and stress at break of AcCel_{*n*}-*b*-PDL_{10k}-*b*-AcCel_{*n*} being 2.3- and 1.8-times higher, respectively, than those of AcMal_{*n*}-*b*-PDL_{10k}-*b*-AcMal_{*n*}. The enhanced microphase-separation and mechanical properties of the AcCel_{*n*}-*b*-PDL-*b*-AcCel_{*n*} were found to be attributable to the stiffness and crystalline nature of the AcCel_{*n*} segments. These results demonstrate the advantages of using cellulose derivatives to synthesize novel bio-functional materials.

Introduction

The development of polymeric materials from natural and renewable resources has recently gained international momentum in order to mitigate the increasing environmental issues.¹⁻³ Carbohydrates are particularly interesting as renewable raw materials because they are readily available from a wide range of plant biomass resources, such as sugar corn, sugarcane, seaweed, waste wood, and agricultural waste.¹ Furthermore, the unique characteristics of carbohydrates, including their biocompatibilities, biodegradabilities, strong hydrophilicities, optical activities, and low toxicities, make them significantly attractive raw materials for creating novel sustainable polymers. Currently, the polymer industry employs carbohydrates mostly as carbon resources for producing bioplastics, such as polylactide, poly(3-hydroxybutyric acid), poly(ethylene furanoate), and poly(2,5-furandicarboxylic acid), through chemical or fermentation processes, despite the fact that cellulose and its derivatives have been used as precursors in their native states.¹⁻³ Using carbohydrates in their native or marginally modified forms may save the energy and time required to transform them to green and sustainable materials, while preserving their unique characteristics.

Directly combining carbohydrate molecules with synthetic polymers is an effective method for achieving the abovementioned goals; this combination can be achieved mainly through grafting synthetic polymer chains onto polysaccharide main chains,⁴⁻⁷ grafting poly/oligosaccharide chains onto synthetic polymer main chains,^{8,9} and tethering poly/oligosaccharides to synthetic polymers to form block copolymers (BCPs).¹⁰⁻¹³ Among these, the approach that creates carbohydrate-based BCPs is particularly attractive because of the microphase-separation and micelle-forming abilities of these polymers, which are useful when designing elastomers,¹⁴⁻¹⁶ lithographic materials,¹⁷⁻¹⁹ electrical materials,²⁰⁻²³ and drug-delivery nanocarriers.²⁴⁻²⁶ Carbohydrate-based BCPs remain neglected compared to the other two carbohydrate-containing polymer types because of the

difficulties associated with their syntheses. Recent progress in polymer chemistry has witnessed the evolution of strategies for the production of carbohydrate-based BCPs since the pioneering syntheses of carbohydrate-based BCPs were published in 1961.^{13,27} For example, cellulose-*b*-polystyrene, dextran-*b*-polystyrene, and maltoheptaose-*b*-poly(ϵ -caprolactone) have been successfully synthesized by radical polymerization and ring-opening polymerization from the reducing end of the carbohydrate segment.^{13,28–31} However, such a synthetic strategy is incompatible with ionic polymerization methods. An alternative pathway for producing carbohydrate-BCPs involves the end-to-end coupling of end-functionalized carbohydrates and synthetic polymer segments, to which various synthetic polymer segments can be joined.^{11,12,17–19,26} In particular, the copper-catalyzed azido-alkyne cycloaddition (CuAAC) reaction has been widely employed for end-to-end coupling because of its high efficiency, simplicity, and functional group tolerance.^{32–34}

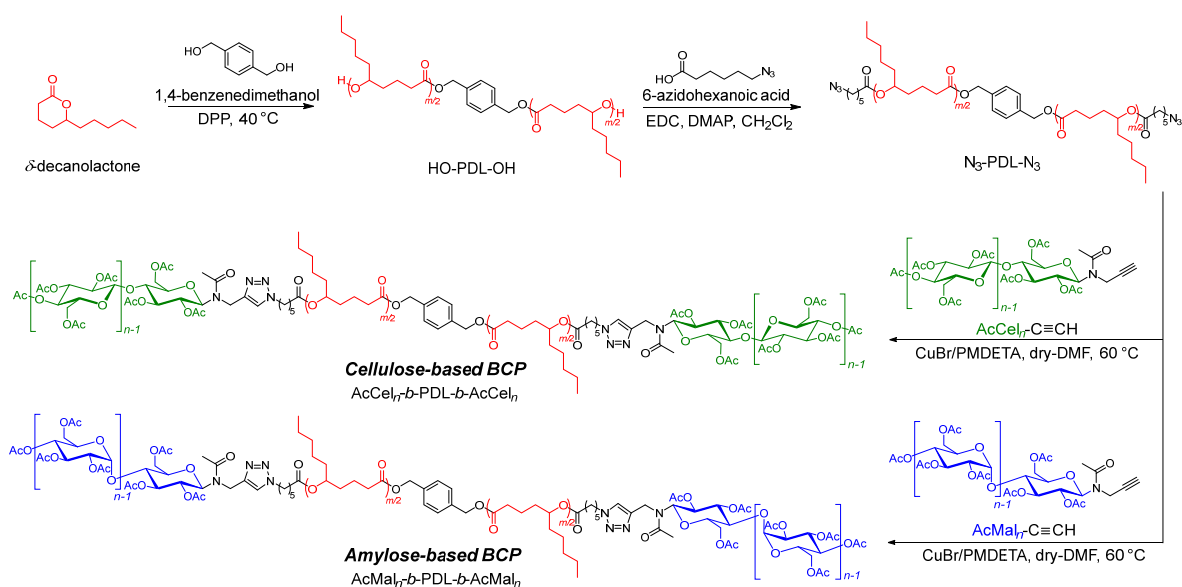
Most carbohydrate-based BCPs comprise dextran and amylose. Surprisingly, there are a few reports on cellulose-containing BCPs, although cellulose and amylose comprise glucose and similar primary structures, with the only structural difference being the presence of α -1,4 linkages in amylose and β -1,4 linkages in cellulose. Consequently, little is known about the self-assembly behavior of cellulose-based BCPs in their solid and solution states, and their fundamental physical properties have not been studied in detail to date. Because cellulose is inedible and is the most abundant polymer on Earth, it is highly attractive as a block partner for constructing carbohydrate-based BCPs. Moreover, it is probable that such cellulose-based BCPs will exhibit enhanced self-assembly properties as well as good mechanical performance because cellulose is highly rigid and crystalline.

However, directly using native cellulose to construct BCPs is significantly challenging because cellulose does not dissolve in common solvents or melt even at high temperatures because of strong inter- and intramolecular hydrogen bonding. This fundamental drawback has historically been overcome by chemically modifying the hydroxyl groups with other functional groups, such as cellulose ethers or esters. Cellulose acetate, one of the most common cellulose derivatives, is readily soluble in common organic solvents and melt-processable. Hence, it has been widely commercialized at different degrees of substitution for use as membranes, textiles, protecting films for polarizers, and cigarette filters. In addition, cellulose acetate has recently gained considerable attention because of its potential biodegradability.^{35,36} Therefore, carbohydrate-based BCPs containing cellulose acetate segments are of significant interest in terms of their synthesis, self-assembly, and physical properties. Recently, the groups of Matson, Edgar, and Kamitakahara reported BCPs containing cellulose derivatives such as cellulose acetate-*b*-polybutadiene-*b*-cellulose acetate,³⁷ cellulose acetate-*b*-oligoamide,³⁸ and trimethyl cellulose-*b*-poly(ethylene glycol).³⁹ However, the microphase-separation behavior and mechanical properties of these BCPs have not been investigated. Furthermore, the advantages of using cellulose derivatives over other carbohydrates remain unclear.

Herein, we report a pioneering comparative study of cellulose-based BCPs and their amylose-based counterparts with the aim of unveiling the unique characteristics imparted by cellulose segments. To this end, we designed and synthesized ABA-type cellooligosaccharide- and maltooligosaccharide-based BCPs, namely, cellooligosaccharide triacetate-*b*-poly(δ -decanolactone)-*b*-cellooligosaccharide triacetates (AcCel_{*n*}-*b*-PDL-*b*-AcCel_{*n*}s) and maltooligosaccharide triacetate-*b*-poly(δ -decanolactone)-*b*-maltooligosaccharide triacetates (AcMal_{*n*}-*b*-PDL-*b*-AcMal_{*n*}s), as shown in **Scheme 1**. Poly(δ -decanolactone) (PDL) is an

amorphous, rubbery, aliphatic polyester that can be synthesized from plant-derived lactone monomers. The “hard moiety-*b*-soft moiety-*b*-hard moiety” ABA-type triblock architecture is important because it endows the BCPs with elastomer-like properties characteristic of the well-known styrene-based thermoplastic elastomers. In fact, our group recently reported completely bio-based elastomers consisting of a maltooligosaccharide as the hard segment and PDL as the soft segment.¹⁴ Importantly, X-ray scattering experiments and atomic force microscopy revealed that the AcCel_{*n*}-*b*-PDL-*b*-AcCel_{*n*}s are more likely to microphase-separate compared to their amylose counterparts. Furthermore, we found that the AcCel_{*n*}-*b*-PDL-*b*-AcCel_{*n*}s have significantly superior mechanical properties compared to the AcMal_{*n*}-*b*-PDL-*b*-AcMal_{*n*}s, in terms of Young’s moduli, elongations at break, and toughnesses.

Scheme 1. Syntheses of AcCel_{*n*}-*b*-PDL-*b*-AcCel_{*n*} and AcMal_{*n*}-*b*-PDL-*b*-AcMal_{*n*}



Results and Discussion

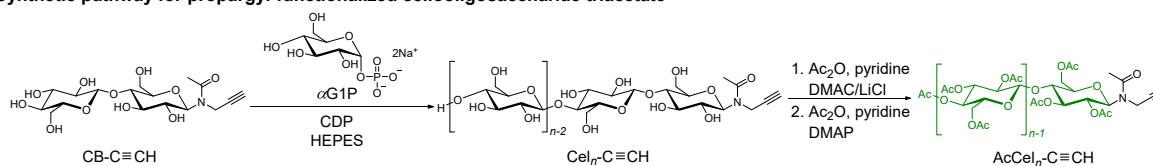
Preparation of propargyl-functionalized celooligosaccharide triacetate and maltooligosaccharide triacetate

We initially synthesized propargyl-functionalized celooligosaccharide triacetate ($\text{AcCel}_n\text{-C}\equiv\text{CH}$) and its maltooligosaccharide counterpart ($\text{AcMal}_n\text{-C}\equiv\text{CH}$). Some researchers reported the syntheses of celooligosaccharides with different functional groups at the reducing end by the cellodextrin-phosphorylase-mediated (CDP-mediated) oligomerization of α -D-glucose-1-phosphate (α G1P) (glycosyl donor) in the presence of anomeric *O*-functionalized glucose (primer).^{40–44} Inspired by this approach, we envisaged the efficient CDP-mediated synthesis of propargyl-functionalized celooligosaccharides ($\text{Cel}_n\text{-C}\equiv\text{CH}$) using propargyl-functionalized primers (**Scheme 2**, upper). In this work, we employed a CDP obtained from *Clostridium thermocellum*. Because His-tagged CDP expressed in *Escherichia coli* BL21(DE3) harbors the pET28a-CDP vector,⁴⁵ it was purified by nickel-nitrilotriacetic acid affinity chromatography.^{40,45} CDP was obtained at a concentration of 7.7 mg mL^{-1} using the Bradford method. According to previous reports on CDP-mediated celooligosaccharide syntheses, glucose and cellobiose are known to act as primers.^{40,42} Therefore, to identify the optimal primer structure for the preparation of propargyl-functionalized celooligosaccharides, we examined celooligosaccharide synthesis using glucose and cellobiose as primers (50 mM) in the presence of CDP (10 mg L^{-1}) and α G1P (200 mM) in 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid (HEPES) buffer solutions (pH = 7.5) at 60°C for 7 d. Both reactions produced a celooligosaccharide as a water-insoluble product; however, the yield was significantly higher when cellobiose ($\sim 23\%$) was used as the primer rather than glucose ($\sim 4\%$). Based on these results, we employed *N*-acetyl-propargyl D-(+)-cellobiose ($\text{CB-C}\equiv\text{CH}$; **Figure S1** for ^1H NMR and FT-IR spectra) as the primer for the synthesis of

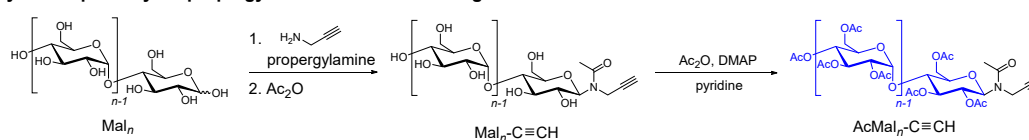
$\text{Cel}_n\text{-C}\equiv\text{CH}$. The desired product was successfully synthesized in 23.0% yield under the aforementioned conditions using $\text{CB-C}\equiv\text{CH}$ as the primer.

Scheme 2. Syntheses of propargyl-functionalized celooligosaccharide triacetate and maltooligosaccharide triacetate

Synthetic pathway for propargyl-functionalized celooligosaccharide triacetate



Synthetic pathway for propargyl-functionalized maltooligosaccharide triacetate



The ^1H NMR spectrum of the prepared $\text{Cel}_n\text{-C}\equiv\text{CH}$ exhibited characteristic signals for the proton on the reducing terminal (I) at 5.39 and 4.88 ppm, the methine group (a) at 2.54 ppm, and the methyl groups (b) at 2.30 and 2.23 ppm; these signals were justifiably assigned to the expected structure (**Figure 1** (a), upper). The number-average degree of polymerization of the prepared $\text{Cel}_n\text{-C}\equiv\text{CH}$ was calculated to be 7.19 by integrating and comparing the signals corresponding to the anomeric protons of the reducing terminal (I) and repeating units (I' and I''), which corresponds to a number-averaged molecular weight ($M_{n,\text{NMR}}$) of 1,260. The FT-IR spectrum of the product shows a characteristic absorption band at $1,645\text{ cm}^{-1}$ that corresponds to the stretching of the amido $\text{C}=\text{O}$ group on the anomeric functional group (**Figure S3** (a)). To investigate the structure in detail, the product was subjected to matrix-assisted laser desorption/ionization time-of-flight mass spectrometry (MALDI-TOF MS). The MALDI-TOF mass spectrum shows two series of repeating peaks separated by approximately 162 Da in the 900–1,600 Da range, among

which one series is consistent with the Na^+ adducts, while the other with the K^+ adducts of $\text{Cel}_n\text{-C}\equiv\text{CH}$ (**Figure S2**). For example, the peak at $m/z = 1254.4$ matches the theoretical molecular mass for the Na^+ adduct of the 7-mer in $\text{Cel}_n\text{-C}\equiv\text{CH}$ ($[\text{M} + \text{Na}]^+ = 1254.4$ Da). The synthesized $\text{Cel}_n\text{-C}\equiv\text{CH}$ mainly comprises 6- to 8-mers according to the MALDI-TOF mass spectrum. The number-average molecular weight was determined by MALDI-TOF MS ($M_{n,\text{MALDI}}$) to be 1,190, which is in good agreement with the $M_{n,\text{NMR}}$ value (1,230).

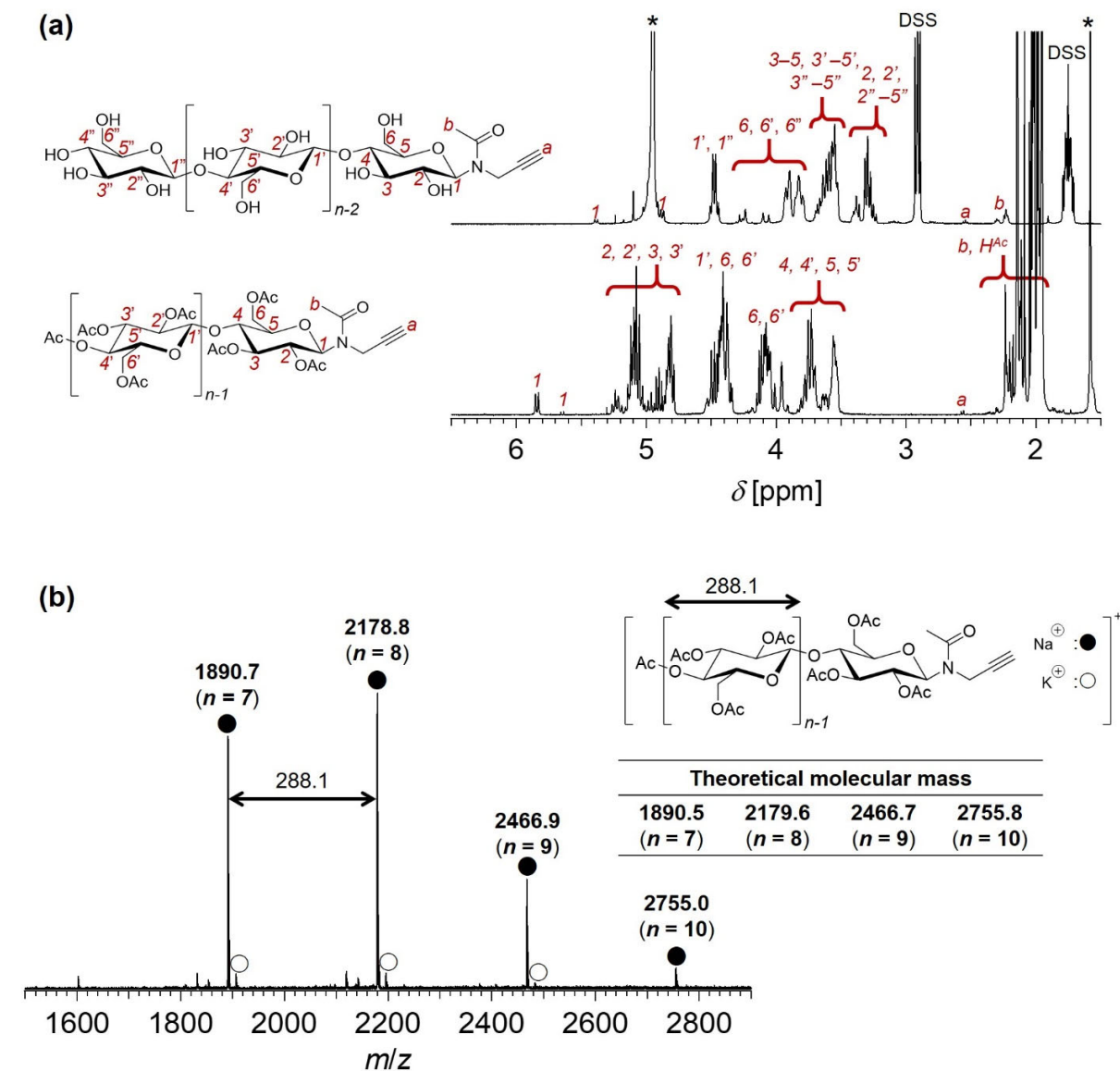


Figure 1. Characterization of $\text{Cel}_n\text{-C}\equiv\text{CH}$ and $\text{AcCel}_n\text{-C}\equiv\text{CH}$. (a) ^1H NMR spectra of $\text{Cel}_n\text{-C}\equiv\text{CH}$ (upper) in D_2O containing sodium trimethylsilylpropanesulfonate (DSS) and 10% (w/w) NaOD and $\text{AcCel}_n\text{-C}\equiv\text{CH}$ in CDCl_3 (lower) (400 MHz). (b) MALDI-TOF mass spectrum and theoretical molecular masses of $\text{AcCel}_n\text{-C}\equiv\text{CH}$.

The prepared $\text{Cel}_n\text{-C}\equiv\text{CH}$ was subsequently subjected to acetylation following solvent-exchange with H_2O , acetone, and *N,N*-dimethylacetamide (DMAc). Acetylation was carried out using Ac_2O and pyridine in LiCl/DMAc for 5 d.⁴⁶ However, ^1H NMR and FT-IR spectroscopy of the product revealed incomplete acetylation. Therefore, the partially acetylated product was further treated with Ac_2O and pyridine in the presence of 4-(dimethylamino)pyridine (DMAP) to ensure complete acetylation. The ^1H NMR spectrum of the final product exhibited the characteristic signals of the acetyl groups (**Figure 1** (a), lower). The number-average degree of polymerization of the prepared $\text{AcCel}_n\text{-C}\equiv\text{CH}$ was calculated to be 6.76 by integrating and comparing the signals corresponding to the anomeric protons of the reducing terminal (*I*) and repeating units (*2*, *2'*, *3*, and *3'*), which corresponds to a $M_{n,\text{NMR}}$ of 2,090. The FT-IR spectrum of the product lacks the absorption band attributable to the hydroxyl groups and exhibits new absorption bands attributable to ester bonds (**Figure S3** (a)). The MALDI-TOF mass spectrum exhibited two series of repeated peaks in the 1,800–3,000 Da range separated by approximately 288 Da, which correspond to the repeating unit of $\text{AcCel}_n\text{-C}\equiv\text{CH}$ (**Figure 1** (b)). These spectroscopic results quantitatively confirmed that all hydroxyl groups of $\text{Cel}_n\text{-C}\equiv\text{CH}$ had been acetylated. The size exclusion chromatography (SEC) trace of $\text{AcCel}_n\text{-C}\equiv\text{CH}$ exhibited a monomodal peak and a narrow dispersity ($D = 1.06$), and the SEC-based number-average molecular weight ($M_{n,\text{SEC}}$) was calculated to be 2,080 when calibrated against a polystyrene standard (**Figure S3** (b)). In summary, we successfully prepared a well-defined and perfectly acetylated $\text{AcCel}_n\text{-C}\equiv\text{CH}$ with an average DP of 6.76.

$\text{AcMal}_n\text{-C}\equiv\text{CH}$, which is the amylose counterpart of $\text{AcCel}_n\text{-C}\equiv\text{CH}$, was successfully prepared in three steps (**Scheme 2**, lower). According to a previously reported procedure,⁴⁷ a commercially available maltooligosaccharide was treated with propargyl amine in the absence of

a solvent followed by Ac₂O in MeOH, to produce propargyl-functionalized maltooligosaccharide (Mal_n-C≡CH). Subsequent *O*-acetylation with Ac₂O in the presence of DMAP in pyridine produced AcMal_n-C≡CH, which mainly comprises 6- to 8-mers according to MALDI-TOF MS. The $M_{n,NMR}$ and D values of the prepared AcMal_n-C≡CH were determined to be 2,020 (DP = 6.51) and 1.09, respectively, according to ¹H NMR spectroscopy and SEC (see **Figures S4** and **S5** for more details).

Triblock copolymer synthesis

Ring-opening polymerization (ROP) of δ -decanolactone was carried at 40 °C using diphenyl phosphate (DPP) and 1,4-benzenedimethanol (BDM) as the catalyst and initiator, respectively, according the procedure reported previously by us, to produce poly(δ -decanolactone) diol (HO-PDL-OH) as the central soft segment of the target triblock copolymer.¹⁴ In this work, PDLs with three molecular weights were synthesized by varying the initial monomer-to-initiator ratio (HO-PDL_{6k}-OH, $M_{n,NMR} = 5,590$, $D = 1.07$; HO-PDL_{10k}-OH, $M_{n,NMR} = 10,000$, $D = 1.06$; HO-PDL_{22k}-OH, $M_{n,NMR} = 21,600$, $D = 1.06$). The HO-PDL-OH groups were thereafter treated with 6-azidohexanoic acid (N₃-COOH) to give the corresponding α,ω -diazido-functionalized PDLs (N₃-PDL_{6k}-N₃, $M_{n,NMR} = 5,900$, $D = 1.07$; N₃-PDL_{10k}-N₃, $M_{n,NMR} = 10,300$, $D = 1.06$; N₃-PDL_{22k}-N₃, $M_{n,NMR} = 21,900$, $D = 1.09$; **Figures S6–S8** show the ¹H NMR spectra of the HO-PDL-OHs and N₃-PDL-N₃s).

Finally, the copper-catalyzed azido-alkyne click reactions between the prepared N₃-PDL-N₃s and AcCel_n-C≡CH or AcMal_n-C≡CH were carried out in DMF using CuBr/PMDETA as the catalyst to produce the target ABA-type triblock BCPs, namely, AcCel_n-*b*-PDL_{6k}-*b*-AcCel_n,

AcCel_n-*b*-PDL_{10k}-*b*-AcCel_n, AcCel_n-*b*-PDL_{22k}-*b*-AcCel_n, AcMal_n-*b*-PDL_{6k}-*b*-AcMal_n, AcMal_n-*b*-PDL_{10k}-*b*-AcMal_n, and AcMal_n-*b*-PDL_{22k}-*b*-AcMal_n, in isolated yields of 26%–80%. The FT-IR spectra of the products confirmed the disappearance of the absorption band at approximately 2,100 cm⁻¹ attributable to azido groups (**Figures S9–14 (b)**), and each SEC elution peak appeared in a higher molecular weight region compared to the azido-functionalized PDL peak, which suggests that each click reaction proceeded quantitatively (**Figure 2**). Furthermore, the ¹H NMR spectra exhibit signals derived from the PDL and AcCel_n or AcMal_n segments that are assigned to the expected chemical structures, thereby confirming that the AcCel_n-*b*-PDL-*b*-AcCel_ns and AcMal_n-*b*-PDL-*b*-AcMal_ns had been successfully synthesized (**Figures S9–S14 (a)**). The molecular weights, *D* values, and volume fractions of the PDL segments (*f*_{PDL}) of the prepared AcCel_n-*b*-PDL-*b*-AcCel_ns and AcMal_n-*b*-PDL-*b*-AcMal_ns are summarized in **Table 1**.

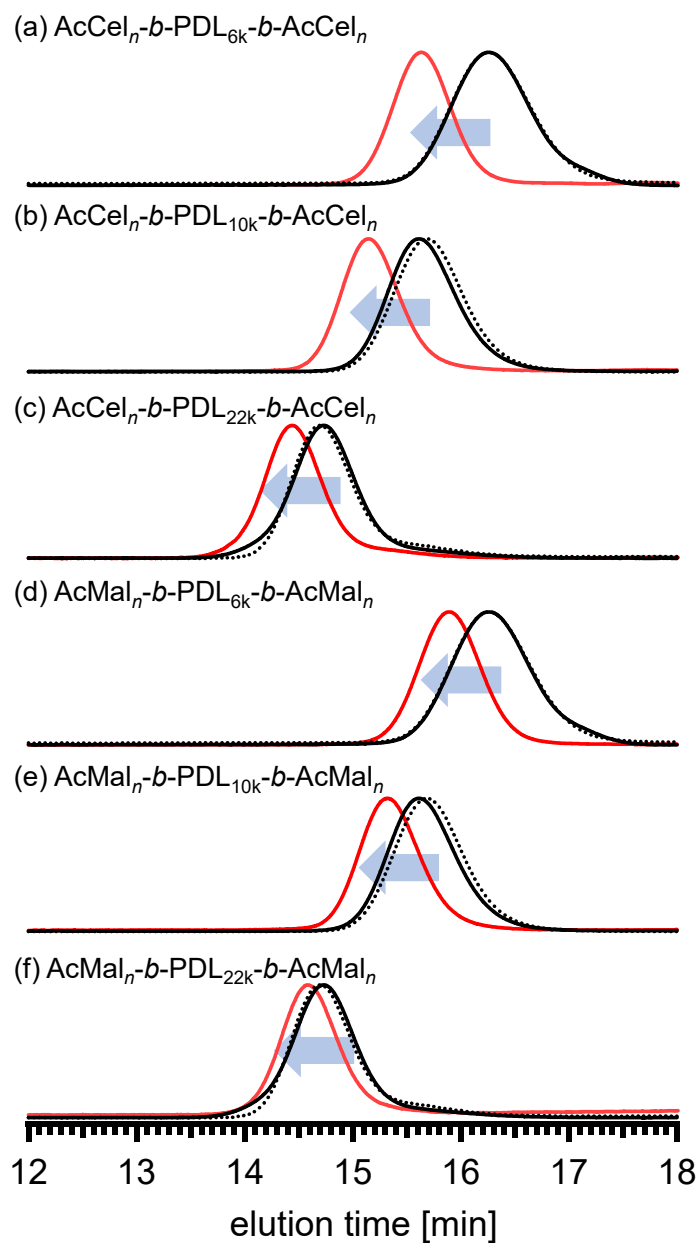


Figure 2. SEC traces of (a) $\text{AcCel}_n\text{-}b\text{-PDL}_{6k}\text{-}b\text{-AcCel}_n$, (b) $\text{AcCel}_n\text{-}b\text{-PDL}_{10k}\text{-}b\text{-AcCel}_n$, (c) $\text{AcCel}_n\text{-}b\text{-PDL}_{22k}\text{-}b\text{-AcCel}_n$, (d) $\text{AcMal}_n\text{-}b\text{-PDL}_{6k}\text{-}b\text{-AcMal}_n$, (e) $\text{AcMal}_n\text{-}b\text{-PDL}_{10k}\text{-}b\text{-AcMal}_n$, and (f) $\text{AcMal}_n\text{-}b\text{-PDL}_{22k}\text{-}b\text{-AcMal}_n$ (eluent, THF; flow rate, 1.0 mL min^{-1}). The solid red, solid black, and dotted black lines represent the BCPs and their corresponding $\text{N}_3\text{-PDL-N}_3$ and HO-PDL-OH units, respectively.

Table 1. Molecular characteristics of AcCel_n-*b*-PDL-*b*-AcCel_{ns} and AcMal_n-*b*-PDL-*b*-AcMal_{ns}

Polymer	<i>M</i> _n	<i>D</i> ^c	<i>T</i> _{g,PDL} ^d [°C]	<i>T</i> _{g,Cel} ^d [°C]	<i>T</i> _{m,Cel} ^d [°C]	<i>T</i> _{g,Mal} ^d [°C]	<i>f</i> _{PDL} ^e
AcCel _n -C≡CH	2,090 ^a	1.06	-	110	167	-	-
AcMal _n -C≡CH	2,020 ^a	1.09	-	-	-	101	-
AcCel _n - <i>b</i> -PDL _{6k} - <i>b</i> -AcCel _n	10,100 ^b	1.04	-52	86	-	-	0.65
AcCel _n - <i>b</i> -PDL _{10k} - <i>b</i> -AcCel _n	14,500 ^b	1.04	-52	97	162	-	0.77
AcCel _n - <i>b</i> -PDL _{22k} - <i>b</i> -AcCel _n	26,100 ^b	1.06	-53	106	-	-	0.87
AcMal _n - <i>b</i> -PDL _{6k} - <i>b</i> -AcMal _n	9,900 ^b	1.04	-52	-	-	60	0.64
AcMal _n - <i>b</i> -PDL _{10k} - <i>b</i> -AcMal _n	14,300 ^b	1.04	-51	-	-	60	0.76
AcMal _n - <i>b</i> -PDL _{22k} - <i>b</i> -AcMal _n	25,900 ^b	1.04	-54	-	-	55	0.87

^aDetermined by ¹H NMR spectroscopy in CDCl₃. ^bCalculated from the molecular weight of AcCel_n-C≡CH or AcMal_n-C≡CH determined by MALDI-TOF and the molecular weight of the corresponding N₃-PDL-N₃ determined by ¹H NMR spectroscopy in CDCl₃. ^cDetermined by SEC in THF using polystyrene standards. ^dDetermined by DSC at a heating rate of 10 °C min⁻¹. ^eCalculated using the density of each block: 1.29 g cm⁻³ for AcCel_n-C≡CH,⁴⁸ 1.20 g cm⁻³ for AcMal_n-C≡CH,⁴⁹ and 0.97 g cm⁻³ for PDL.⁵⁰

Thermal properties

To gain fundamental insight into the differences in the physical properties originating from the oligosaccharide blocks, we initially evaluated the thermal properties of the AcCel_n-*b*-PDL-*b*-AcCel_{ns} and AcMal_n-*b*-PDL-*b*-AcMal_{ns} by thermogravimetric analysis (TGA) and differential scanning calorimetry (DSC) in a nitrogen atmosphere. According to the TGA data, all of the studied BCP samples were thermally stable up to approximately 280 °C and exhibited 5% weight-loss temperatures (*T*_{5%}) in the 290–320 °C range (**Figures S15** and **S16**). Visual inspection during TGA revealed that all BCP samples changed their appearance; they transformed from a solid to a

melt state close to the glass transition temperature of the AcCel_n or AcMal_n segment, consistent with thermoplastic behavior. The DSC curves of the AcCel_n-*b*-PDL-*b*-AcCel_ns and AcMal_n-*b*-PDL-*b*-AcMal_ns acquired during the second heating process exhibited two baseline shifts derived from the glass transitions of the PDL and AcCel_n or AcMal_n segments (**Figure 3**). The glass transition temperatures of the PDL ($T_{g,PDL}$), AcCel_n ($T_{g,Cel}$), and AcMal_n segments ($T_{g,Mal}$) are approximately -50, 95, and 60 °C, respectively (**Figures 3 and S17**). The fact that $T_{g,PDL}$ and $T_{g,Cel}$ (or $T_{g,Mal}$) were observed separately indicates phase separation between the PDL and AcCel_n (or AcMal_n) segments. The $T_{g,Mal}$ values of the AcMal_n-*b*-PDL-*b*-AcMal_ns were considerably lower than that of AcMal_n-C≡CH, which indicates partial mixing of the AcMal_n and PDL domains. Similarly, partial mixing of the AcCel_n and PDL domains resulted in lower $T_{g,Cel}$ values for the AcCel_n-*b*-PDL-*b*-AcCel_ns than AcCel_n-C≡CH. Interestingly, $T_{g,Cel}$ was observed to increase with increasing f_{PDL} , which is attributable to increasing segregation strength associated with increasing AcCel_n-*b*-PDL-*b*-AcCel_n total molecular weight. Notably, the DSC curve of AcCel_n-*b*-PDL_{10k}-*b*-AcCel_n exhibited a small endothermic peak at 163 °C due to the melting of the AcCel_n segment ($T_{m,Cel}$), which indicates that the BCP is semi-crystalline in nature. The DSC trace of AcCel_n-C≡CH clearly exhibits a $T_{m,Cel}$ at 167 °C; however, the traces for AcCel_n-*b*-PDL_{6k}-*b*-AcCel_n and AcCel_n-*b*-PDL_{22k}-*b*-AcCel_n did not exhibit any melting peaks. Given the small area of the melting transition of the AcCel_n segment in AcCel_n-*b*-PDL_{10k}-*b*-AcCel_n, crystallization tends to occur slowly. In contrast, the AcMal_n-*b*-PDL-*b*-AcMal_ns and AcMal_n-C≡CH did not exhibit any melting transition, which confirms the amorphous nature of the AcMal_n segment.

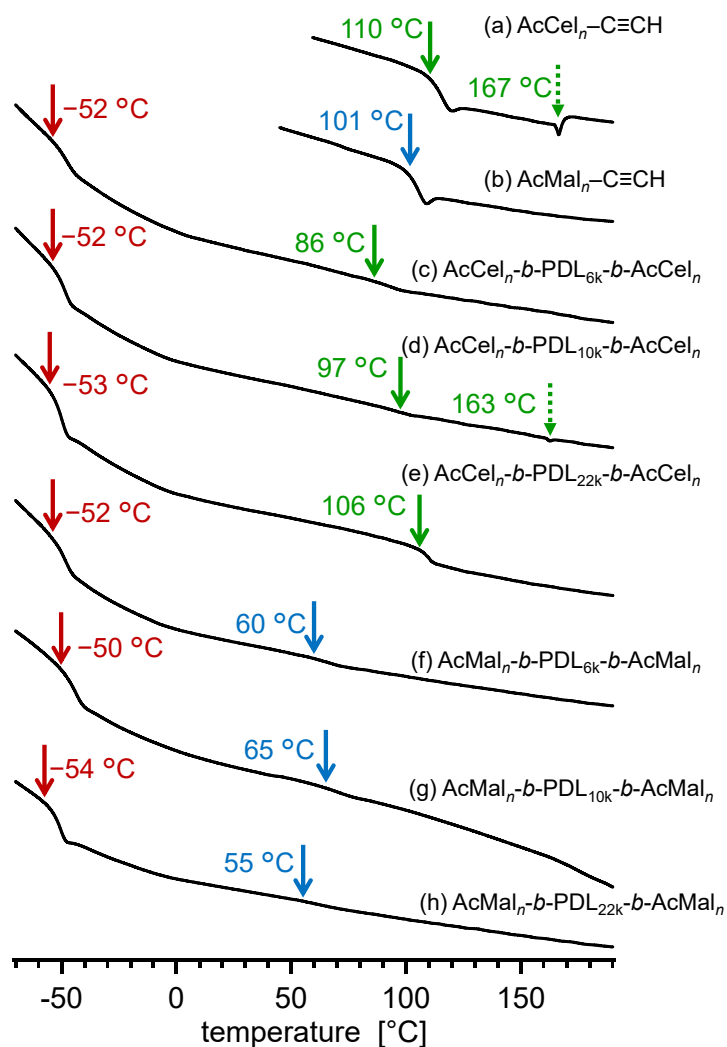


Figure 3. DSC curves of (a) $\text{AcCel}_n\text{-C}\equiv\text{CH}$, (b) $\text{AcMal}_n\text{-C}\equiv\text{CH}$, (c) $\text{AcCel}_n\text{-}b\text{-PDL}_{6k}\text{-}b\text{-AcCel}_n$, (d) $\text{AcCel}_n\text{-}b\text{-PDL}_{10k}\text{-}b\text{-AcCel}_n$, (e) $\text{AcCel}_n\text{-}b\text{-PDL}_{22k}\text{-}b\text{-AcCel}_n$, (f) $\text{AcMal}_n\text{-}b\text{-PDL}_{6k}\text{-}b\text{-AcMal}_n$, (g) $\text{AcMal}_n\text{-}b\text{-PDL}_{10k}\text{-}b\text{-AcMal}_n$, and (h) $\text{AcMal}_n\text{-}b\text{-PDL}_{22k}\text{-}b\text{-AcMal}_n$ during the second heating run (heating rate: $10\text{ }^\circ\text{C min}^{-1}$).

Microphase-separated structures

To investigate the impact of the oligosaccharide segments on the BCP self-assembly behavior, the microphase-separated structures of the $\text{AcCel}_n\text{-}b\text{-PDL-}b\text{-AcCel}_n$ s and $\text{AcMal}_n\text{-}b\text{-PDL-}b\text{-AcMal}_n$ s were investigated through small-angle X-ray scattering (SAXS) experiments on bulk samples, which were prepared by casting toluene solutions of the BCPs, followed by drying for 15 h at atmospheric pressure and for 1 d under vacuum at room temperature. The prepared films were subsequently thermally annealed under vacuum at 130 °C for 6 or 24 h.

The SAXS profiles of the non-annealed $\text{AcCel}_n\text{-}b\text{-PDL-}b\text{-AcCel}_n$ s and $\text{AcMal}_n\text{-}b\text{-PDL-}b\text{-AcMal}_n$ s exhibited only primary peaks indicative of weak segregation between the PDL and acetyl oligosaccharide segments with no periodically ordered structures (**Figure 4**). The mean distance between the phase-separated microdomains was calculated to be approximately 10 nm based on the Bragg equation using the position of the primary peak (q^*). In contrast, dramatic differences were observed in the microphase-separated structures of the $\text{AcCel}_n\text{-}b\text{-PDL-}b\text{-AcCel}_n$ s and $\text{AcMal}_n\text{-}b\text{-PDL-}b\text{-AcMal}_n$ s upon thermal annealing. The SAXS profiles of the $\text{AcMal}_n\text{-}b\text{-PDL-}b\text{-AcMal}_n$ s exhibited only primary peaks, regardless of the annealing conditions, which implies that these BCPs do not meet the requirements for ordered microphase-separation. In contrast, the SAXS profiles of the $\text{AcCel}_n\text{-}b\text{-PDL-}b\text{-AcCel}_n$ s exhibited primary peaks and additional higher-order scattering peaks upon thermal annealing at 130 °C for appropriate durations. The SAXS profile of $\text{AcCel}_n\text{-}b\text{-PDL}_{6k}\text{-}b\text{-AcCel}_n$ exhibited a higher-order scattering peak at $3q^*$ in addition to q^* , indicative of a lamellar (LAM) structure (**Figure 4 (a)**); no even-ordered peak appeared in this SAXS profile, which indicates that the volumes of the AcCel_n and PDL microdomains are nearly equal.⁵¹ The SAXS profiles of $\text{AcCel}_n\text{-}b\text{-PDL}_{10k}\text{-}b\text{-AcCel}_n$ and $\text{AcCel}_n\text{-}b\text{-PDL}_{22k}\text{-}b\text{-AcCel}_n$ also exhibited higher-order scattering peaks at $\sqrt{3}q^*$, $2q^*$, $\sqrt{7}q^*$, and $3q^*$, and $\sqrt{2}q^*$, $\sqrt{3}q^*$, $\sqrt{5}q^*$, and $\sqrt{7}q^*$,

indicative of hexagonal cylindrical (HEX) and body-centered cubic (BCC) spherical structures, respectively (**Figure 4** (c) and (e), respectively). The domain-spacing (d) of the thermally annealed $\text{AcCel}_n\text{-}b\text{-PDL}_{6k}\text{-}b\text{-AcCel}_n$, $\text{AcCel}_n\text{-}b\text{-PDL}_{10k}\text{-}b\text{-AcCel}_n$, and $\text{AcCel}_n\text{-}b\text{-PDL}_{22k}\text{-}b\text{-AcCel}_n$ were determined to be 9.6, 10.3, and 12.1 nm, respectively; the d value increases with increasing molecular weight. Overall, the $\text{AcCel}_n\text{-}b\text{-PDL-}b\text{-AcCel}_n$ s are more likely to undergo ordered microphase-separation than the $\text{AcMal}_n\text{-}b\text{-PDL-}b\text{-AcMal}_n$ s, despite being composed of the same monomeric units and possessing comparable chemical compositions. This is partly attributable to the rigidity granted to the AcCel_n segments by their β -1,4-linked cellulose backbone. The longer Kuhn and persistence lengths of cellulose compared to those of amylose^{13,52,53} induce microphase-separation in the $\text{AcCel}_n\text{-}b\text{-PDL-}b\text{-AcCel}_n$ s. The fact that $\text{AcCel}_n\text{-C}\equiv\text{CH}$ has a larger $M_{n,\text{SEC}}$ than $\text{AcMal}_n\text{-C}\equiv\text{CH}$, despite their comparable $M_{n,\text{NMR}}$ values, implies that the AcCel_n segment is more rigid than AcMal_n . In addition, the AcCel_n segment exhibited the potential to crystallize, as revealed by DSC, which indicates its stronger tendency to self-aggregate. The wide-angle X-ray diffraction (WAXD) data also confirmed that the AcCel_n segment is crystalline in nature (**Figures S18** and **S19**). While the WAXD profile of $\text{AcMal}_n\text{-C}\equiv\text{CH}$ and the $\text{AcMal}_n\text{-}b\text{-PDL-}b\text{-AcMal}_n$ s showed only amorphous halos, $\text{AcCel}_n\text{-C}\equiv\text{CH}$ and the $\text{AcCel}_n\text{-}b\text{-PDL-}b\text{-AcCel}_n$ s showed diffraction peaks assignable to cellulose triacetate I crystals.^{54,55} This may further enhance microphase-separation in the $\text{AcCel}_n\text{-}b\text{-PDL-}b\text{-AcCel}_n$ s and significantly alter the self-assembly properties of the $\text{AcCel}_n\text{-}b\text{-PDL-}b\text{-AcCel}_n$ s and $\text{AcMal}_n\text{-}b\text{-PDL-}b\text{-AcMal}_n$ s.

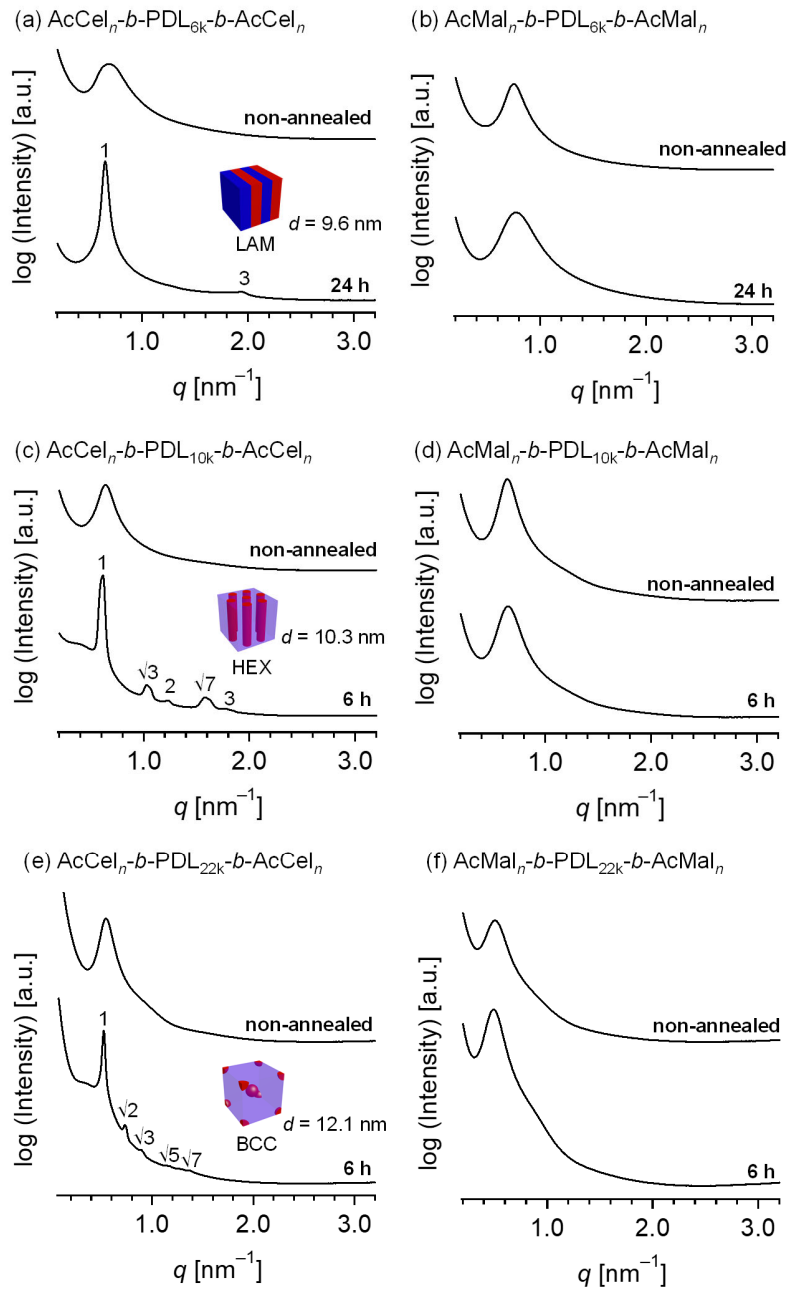


Figure 4. SAXS profiles of (a) $\text{AcCel}_n\text{-}b\text{-PDL}_{6k}\text{-}b\text{-AcCel}_n$, (b) $\text{AcMal}_n\text{-}b\text{-PDL}_{6k}\text{-}b\text{-AcMal}_n$, (c) $\text{AcCel}_n\text{-}b\text{-PDL}_{10k}\text{-}b\text{-AcCel}_n$, (d) $\text{AcMal}_n\text{-}b\text{-PDL}_{10k}\text{-}b\text{-AcMal}_n$, (e) $\text{AcCel}_n\text{-}b\text{-PDL}_{22k}\text{-}b\text{-AcCel}_n$, and (f) $\text{AcMal}_n\text{-}b\text{-PDL}_{22k}\text{-}b\text{-AcMal}_n$ without thermal annealing and after annealing at 130 °C for 6 or 24 h.

To further investigate the microphase-separation behavior, we subjected BCP thin films to atomic force microscopy (AFM) (**Figure S20**); the thin-film samples were prepared by spin-coating 1% (w/w) solutions of $\text{AcCel}_n\text{-}b\text{-PDL-}b\text{-AcCel}_n$ s and $\text{AcMal}_n\text{-}b\text{-PDL-}b\text{-AcMal}_n$ s in toluene onto bare Si substrates. The AFM phase images of the non-annealed $\text{AcCel}_n\text{-}b\text{-PDL}_{6k}\text{-}b\text{-AcCel}_n$, $\text{AcCel}_n\text{-}b\text{-PDL}_{10k}\text{-}b\text{-AcCel}_n$, $\text{AcMal}_n\text{-}b\text{-PDL}_{6k}\text{-}b\text{-AcMal}_n$, and $\text{AcMal}_n\text{-}b\text{-PDL}_{10k}\text{-}b\text{-AcMal}_n$ thin films show less-ordered fingerprint patterns. In contrast, the non-annealed $\text{AcCel}_n\text{-}b\text{-PDL}_{22k}\text{-}b\text{-AcCel}_n$ and $\text{AcMal}_n\text{-}b\text{-PDL}_{22k}\text{-}b\text{-AcMal}_n$ thin films show less-ordered spheres. These results are consistent with the fact that featureless SAXS profiles were observed for the non-annealed samples in the bulk. The morphologies of the $\text{AcCel}_n\text{-}b\text{-PDL-}b\text{-AcCel}_n$ s were significantly altered by thermal annealing in the thin-film and bulk states. The AFM phase image of the $\text{AcCel}_n\text{-}b\text{-PDL}_{6k}\text{-}b\text{-AcCel}_n$ thin film showed a peculiar morphology consisting of rectangular units, which are related to the crystallization of the AcCel_n segment.⁵⁶ A highly ordered fingerprint-like pattern was observed in the thermally annealed $\text{AcCel}_n\text{-}b\text{-PDL}_{10k}\text{-}b\text{-AcCel}_n$ thin film, which is attributable to the horizontally orientated HEX structure, considering the bulk SAXS result. The spherical structures observed in the thermally annealed $\text{AcCel}_n\text{-}b\text{-PDL}_{22k}\text{-}b\text{-AcCel}_n$ are attributable to the BCC spherical structure. The AFM images of $\text{AcCel}_n\text{-}b\text{-PDL}_{10k}\text{-}b\text{-AcCel}_n$ and $\text{AcCel}_n\text{-}b\text{-PDL}_{22k}\text{-}b\text{-AcCel}_n$ revealed the d values of 10.8 and 12.0 nm, respectively, which were in good agreement with the values derived from the bulk SAXS results. As expected from the bulk SAXS results, the $\text{AcMal}_n\text{-}b\text{-PDL-}b\text{-AcMal}_n$ thin films did not exhibit any ordered patterns following thermal annealing, which verifies that the cellulose-based BCPs are more likely to microphase-separate than their amylose counterparts.

Mechanical properties

Finally, to investigate the impact of the hard oligosaccharide segment on mechanical properties, dog-bone-shaped specimens of solvent-cast BCPs were subjected to tensile testing. Note that it was difficult to subject $\text{AcMal}_n\text{-}b\text{-PDL}_{22k}\text{-}b\text{-AcMal}_n$ to tensile testing because it is a sticky material. The other BCPs exhibited elastic properties, which confirms that the hard microphase-separated oligosaccharide domains act as physical crosslinks between the rubbery PDL chains. The stress–strain curves and key mechanical properties (Young’s modulus (E), strain at break (ϵ_b), stress at break (σ_b), and toughness, and their average values and standard deviations) of the BCP specimens are shown in **Figure 5** and **Table 2**, respectively.

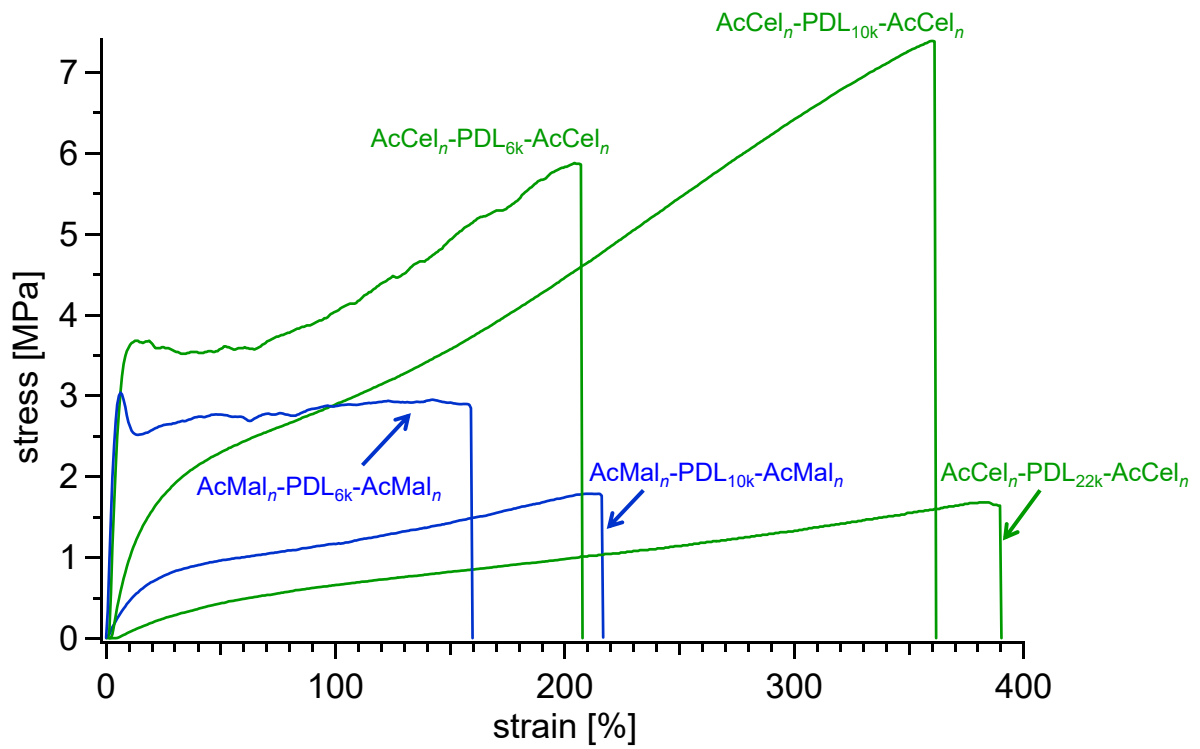


Figure 5. Typical stress–strain curves of the $\text{AcCel}_n\text{-}b\text{-PDL}_{6k}\text{-}b\text{-AcCel}_n$, $\text{AcCel}_n\text{-}b\text{-PDL}_{10k}\text{-}b\text{-AcCel}_n$, $\text{AcCel}_n\text{-}b\text{-PDL}_{22k}\text{-}b\text{-AcCel}_n$, $\text{AcMal}_n\text{-}b\text{-PDL}_{6k}\text{-}b\text{-AcMal}_n$, and $\text{AcMal}_n\text{-}b\text{-PDL}_{10k}\text{-}b\text{-AcMal}_n$ specimens (crosshead speed: 10 mm min^{-1}).

Table 2. Tensile properties of the BCP specimens^a

BCP	E^a [MPa]	ε_b^a [%]	σ_b^a [MPa]	Toughness ^a [MJ m ⁻³]
AcCel _n - <i>b</i> -PDL _{6k} - <i>b</i> -AcCel _n	69±5.1	209±17.6	5.8±0.082	8.9±0.90
AcCel _n - <i>b</i> -PDL _{10k} - <i>b</i> -AcCel _n	13±1.7	356±8.99	7.1±0.47	15±1.2
AcCel _n - <i>b</i> -PDL _{22k} - <i>b</i> -AcCel _n	1.3±0.062	385±16.0	1.7±0.047	3.8±0.17
AcMal _n - <i>b</i> -PDL _{6k} - <i>b</i> -AcMal _n	61±6.8	150±14.6	2.8±0.12	4.2±0.48
AcMal _n - <i>b</i> -PDL _{10k} - <i>b</i> -AcMal _n	5.6±0.68	198±13.6	1.6±0.14	2.2±0.29

^aTensile properties are shown as average values (with standard deviations) for three specimens.

The stress–strain curves show that AcCel_n-*b*-PDL_{6k}-*b*-AcCel_n and AcMal_n-*b*-PDL_{6k}-*b*-AcMal_n, which possess the lowest f_{PDL} values, exhibit yield points, suggestive of plastic-like behavior irrespective of the type of hard segment. In contrast, AcCel_n-*b*-PDL_{10k}-*b*-AcCel_n, AcCel_n-*b*-PDL_{22k}-*b*-AcCel_n, and AcMal_n-*b*-PDL_{10k}-*b*-AcMal_n did not exhibit clear yield points, suggesting that BCPs with longer PDL segments exhibit elastomer-like behavior, which is supported by the results of cyclic tensile testing (**Figure S21**), which suggest that the areas enclosed by hysteresis loops, which correlate with the level of plastic deformation, diminish with increasing f_{PDL} . No obvious residual strain was observed following five loading/unloading cycles for AcCel_n-*b*-PDL_{22k}-*b*-AcCel_n, confirming its high level of elastic recovery. A general trend in which ε_b increases and E decreases with increasing f_{PDL} was observed; for example, the ε_b value of AcCel_n-*b*-PDL_{22k}-*b*-AcCel_n was 1.8-times higher than that of AcCel_n-*b*-PDL_{6k}-*b*-AcCel_n, and the E value of AcCel_n-*b*-PDL_{6k}-*b*-AcCel_n was 53-times higher than that of AcCel_n-*b*-PDL_{22k}-*b*-AcCel_n. This trend was observed when the ε_b and E values for AcMal_n-*b*-PDL_{6k}-*b*-AcMal_n and AcMal_n-*b*-PDL_{10k}-*b*-AcMal_n were compared.

The BCPs containing AcCel_n segments possess significantly superior mechanical properties to those containing AcMal_n segments, despite their comparable molecular weights and chemical compositions. Specifically, the E , ϵ_b , σ_b , and toughness values of AcCel_n-*b*-PDL_{10k}-*b*-AcCel_n were 2.3-, 1.8-, 4.4-, and 6.8-times greater than those of AcMal_n-*b*-PDL_{10k}-*b*-AcMal_n, respectively. Furthermore, stress on the AcCel_n-*b*-PDL_{6k}-*b*-AcCel_n specimen gradually increased beyond the yield point, while AcMal_n-*b*-PDL_{6k}-*b*-AcMal_n maintained constant stress beyond the yield point until rupture. AcCel_n-*b*-PDL-*b*-AcCel_n strain hardening reveals that cellulosic hard-segment pull-out was more effectively hindered in these polymers compared to that in their amylose counterparts.⁵⁷ The stronger crosslinking ability of the AcCel_n segment is partly attributable to its crystalline nature, in contrast to the amorphous nature of AcMal_n. X-ray crystallography previously revealed that cellulose triacetate and amylose triacetate adopt sheet and helix conformations, respectively.^{48,49,58,59} Evidently, the sheet-like conformation is preferred for creating larger interchain attractive forces, which may be responsible for the stronger crosslinking of the AcCel_n segment. In addition, the higher potential to form ordered microphase-separated structures seems to be another important factor that contributes to the superior mechanical performance of the AcCel_n-*b*-PDL-*b*-AcCel_ns.

Figure S22 compares σ_b and E of AcCel_n-*b*-PDL_{6k}-*b*-AcCel_n, AcCel_n-*b*-PDL_{10k}-*b*-AcCel_n, AcCel_n-*b*-PDL_{22k}-*b*-AcCel_n, AcMal_n-*b*-PDL_{6k}-*b*-AcMal_n, and AcMal_n-*b*-PDL_{10k}-*b*-AcMal_n with those of commercial thermoplastic elastomers (TPEs). Notably, the E value of AcCel_n-*b*-PDL_{10k}-*b*-AcCel_n was comparable to or even higher than those of commercial TPEs, such as melt-processable rubber and TPEs containing polyamide hard blocks. The σ_b of AcCel_n-PDL_{10k}-*b*-AcCel_n was also comparable to those of commercial TPEs, such as melt-processable rubber and

silicone TPEs. This comparison highlights the potential of the AcCel_n-*b*-PDL-*b*-AcCel_ns as fully bio-based high-performance elastomers.

Conclusions

In this work, novel cellulose-based BCPs (AcCel_n-*b*-PDL-*b*-AcCel_ns) and their amylose counterparts (AcMal_n-*b*-PDL-*b*-AcMal_ns) were precisely synthesized. The AcCel_n-*b*-PDL-*b*-AcCel_ns were found to be more likely to self-assemble than the AcMal_n-*b*-PDL-*b*-AcMal_ns in their bulk and thin-film states. Because these BCPs have comparable chemical compositions and molecular weights, the dramatic difference in their self-assembly behavior stems from the differences in the types of bonding (i.e., β -1,4 or α -1,4 linkages) in their oligosaccharide segments. More interestingly, the AcCel_n-*b*-PDL-*b*-AcCel_ns exhibit superior mechanical properties than the AcMal_n-*b*-PDL-*b*-AcMal_ns, which indicates that stronger physical crosslinking was achieved in the AcCel_n segments. In addition, the Young's modulus (~69 MPa) and elongation at break (~5.8 MPa) of the AcCel_n-*b*-PDL-*b*-AcCel_ns were comparable to or higher than those of commercially available TPEs. To the best of our knowledge, this is the first report that investigates in detail how small differences in the primary structures of oligosaccharide segments (i.e., α -1,4 or β -1,4 linkages) affect the microphase-separation and mechanical properties of carbohydrate-based BCPs. Overall, we verified the applicability of cellulose-based BCPs exhibit as bio-based and sustainable polymeric materials. Encouraged by these promising results, we are currently working toward the syntheses of bio-based plastics, elastomers, and fibers that possess sufficiently high strengths and toughnesses for practical use by either optimizing the degree of substitution on the cellulose segment or by introducing other soft middle segments.

ASSOCIATED CONTENT

Supporting Information. Experimental section, comparative NMR and FT-IR spectra of precursors, MALDI-TOF mass spectra, comparative FT-IR spectra of precursors and cellulose-based products, SEC traces of cellulosic products, TGA, DSC, AFM, and cyclic tensile testing results.

Author Contributions

K.T., T.I., and T.S. designed the experiments. S.K. wrote the manuscript. S.K. and Y.T. synthesized and characterized the BCPs and analyzed the X-ray data. N.S. and K.I. constructed the plasmid to express CDP and established the process for preparing CDP. T.Y. contributed to the discussion. The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript.

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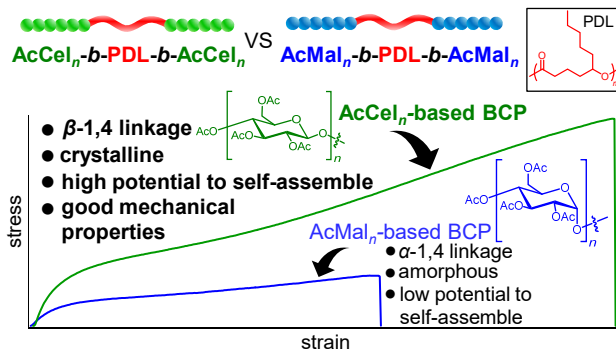
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SYNOPSIS: Environmentally friendly cellulose-based block copolymers were self-assembled and exhibited higher mechanical strength than commercially available thermoplastic elastomers.