Shell Architecture Strongly Influences the Glass Transition, Surface Mobility, and Elasticity of Polymer Core-Shell Nanoparticles

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INTRODUCTION

Nanoscaled polymeric materials are increasingly used in advanced technologies, for example, as separation membranes,1,2 drug-delivery vehicles,3 reinforcing modifiers,4 and photonic5 and phononic crystals.6 These technologies utilize one-dimensional (e.g., thin films) and three-dimensional (e.g., nanoparticles (NPs)) confinement to modify polymer properties. When the length scale of a polymeric system is reduced to the nanoscale, polymeric materials show deviations in physical properties,9 such as viscosity,10–12 elasticity,13,14 and glass-transition temperature (Tg) from the bulk.15–23 Although the mechanism behind these deviations is of much debate, interfacial and surface effects dominate confined polymer properties due to the large ratio of surface area to volume.24 The influence of polymer–polymer interfaces on the Tg of confined polymers remains an open question.25,26 Polymer Tg near the polymer–polymer interface has been observed in stacked multilayer films,27–29 and phase-separated block copolymers.30–32 The interfacial width, contrast between bulk Tg values, chain connectivity, and relative Debye–Waller factors may all play a role in determining the change in Tg across a polymer–polymer interface.30,31 Less well-studied is the influence of polymer–polymer interfaces on the mechanical properties of confined polymers.13,32–36 Various techniques including buckling,13 nanobubble inflation,34 and nanoindentation34,37 have been used to measure the Young modulus or compliance of polymer thin films both above and below the Tg. Below the Tg both a substantial reduction13,33 and an increase37 in the Young modulus, depending on the nature of the underlying substrate, have been measured for films with a thickness less than ~40 nm. In rubbery films, above the Tg, stiffening has been observed under nanoscale confinement.32 Although these experiments have been conducted in a variety of geometries including freestanding films, films supported atop elastomers, and films with and without a topcoat polymer layer supported atop silicon, a thorough investigation of the mechanical properties near polymer–polymer interfaces of confined systems has not yet been conducted.

ABSTRACT: Despite the growing application of nanostructured polymeric materials, there still remains a large gap in our understanding of polymer mechanics and thermal stability under confinement and near polymer–polymer interfaces. In particular, the knowledge of polymer nanoparticle thermal stability and mechanics is of great importance for their application in drug delivery, phononics, and photonics. Here, we quantified the effects of a polymer shell layer on the modulus and glass-transition temperature (Tg) of polymer core–shell nanoparticles via Brillouin light spectroscopy and modulated differential scanning calorimetry, respectively. Nanoparticles consisting of a polystyrene (PS) core and shell layers of poly(n-butyl methacrylate) (PBMA) were characterized as model systems. We found that the high Tg of the PS core was largely unaffected by the presence of an outer polymer shell, whereas the lower Tg of the PBMA shell layer decreased with increasing PBMA thickness. The surface mobility was revealed at a temperature about 15 K lower than the Tg. The in

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Supporting Information

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been undertaken. To understand the effect of polymer interfaces on the modulus and \( T_g \) we recently introduced polystyrene (PS) core–shell NPs with varying ultrathin shells via seeded surfactant-free polymerization and layer-by-layer adsorption.\(^{43} \) We found a significant change in the elastic modulus and thermal behavior of NPs by the introduction of ultrathin shell layers with different chemical compositions via Brillouin light-scattering spectroscopy (BLS).\(^{38} \)

Recently, focus has been placed on studying deviations in the glass transition of nanoparticles.\(^{38-44} \) Spherical geometries have received less attention partially due to the limited number of experimental techniques available to measure the material properties of nanoparticles. Zhang et al.\(^{40} \) first measured a decrease in the \( T_g \) (∼50 K) for ∼90 nm PS NPs prepared by surfactant-free emulsion polymerization. When capped with a silica shell, the \( T_g \) confinement effect was absent due to the removal of the free surface.\(^{40} \) Feng et al.\(^{41} \) modified PS NPs with different types of surfactants and discovered a strong surfactant effect on \( T_g \) deviations for NPs less than 200 nm in diameter, thus emphasizing the importance of the NP surface in determining confinement effects. Furthermore, recent measurements of molecular mobility in NPs\(^{42,43} \) have shown a decoupling between the decrease in the NP \( T_g \) with decreasing diameter and no change in the segmental mobility as measured by dielectric relaxation spectroscopy\(^{45} \) and fast-scanning calorimetry.\(^{46} \)

Here, we examine PS-core NPs with a low-\( T_g \) poly(butyl methacrylate) (PBMA) shell layer of varying thickness to understand the effects of the outermost surface and core–shell interface on the thermomechanical properties of the shell. The PS core was synthesized by surfactant-free polymerization. Seeded surfactant-free polymerization was adapted to produce the PS–PBMA and PS–poly(methyl methacrylate) (PMMA) core–shell NPs. The synthesis of core–shell NPs is elaborated in the Supporting Information. BLS with a high-resolution tandem Fabry–Perot interferometer was used to measure the elastic properties, particle surface softening temperature (\( T_s \)), and \( T_g \). Modulated differential scanning calorimetry (mDSC) was used to measure the \( T_g \) of the NPs. To understand the influence of the shell layer on the thermomechanical properties via BLS and mDSC, we have also investigated the thermal behavior of contiguous films prepared by annealing NPs.

**RESULTS AND DISCUSSION**

**Nanoparticle Shear Elastic Modulus.** Figure 1 shows the anti-Stokes side of the BLS spectra of four different close-packed clusters of core–shell NPs, PS–PS, PS–PMMA\(_{30}\), PS–PBMA\(_{30}\), and PS–PBMA\(_{45}\) at 294 K with the particle size information listed in Table 1. For all core–shell NPs, the diameter was obtained from the SEM images, averaging the size of 100 randomly chosen particles. The identification number within the nomenclature represents the thickness of the shell in nanometers and a change in the shell volume fraction. In the quasi-core–shell PS–PS NP, the thin PS shell layer, which has the same chemistry as the PS core, shows negligible effects on the particle elasticity\(^{38} \) and hence is used as a reference. To compare the spectral features of systems differing both in structure and size, we account for the latter by scaling the particle vibration frequencies by the NP diameter (\( d \)). The BLS spectra were then plotted against the particle diameter normalized frequency \( (\omega d) \) as shown in Figure 1a. In principle, BLS probes the spheroidal mechanical eigenmodes of NPs. The eigenfrequencies of these modes are labeled \( j(n_l) \), where \( n \) and \( l \) are integers defining the radial and angular dependence of the displacement, respectively.\(^{37} \) To determine the thermomechanical properties of NPs, we focused on the spectral position and line shape of the \( (1,2) \) mode. Here, 1 and 2 stand for the radial and angular vibrations of spheres, respectively.\(^{48} \) As depicted in Figure 1, in each case, the \( (1,2) \) modes correspond to Lorentzian doublets (solid gray lines) peaked at \( f_1 \) and \( f_2 \). The split and blue shift of the \( f(1,2) \) with respect to the isolated NPs is the result of attractive

**Table 1. Dimensions of Shell Architecture of PS-Based Core–Shell Nanoparticles**

| samples       | PS–PS | PS–PBMA\(_30\) | PS–PBMA\(_{45}\) | PS–PMMA\(_{30}\) | PS–PMMA\(_{45}\) |
|---------------|-------|----------------|----------------|----------------|----------------|
| core diameter (nm) | 222 ± 7 | 220 ± 7 | 222 ± 7 | 201 ± 9 | 220 ± 7 |
| NP diameter (d) (nm) | 256 ± 6 | 234 ± 9 | 290 ± 11 | 317 ±12 | 236 ± 7 | 254 ± 6 |
| shell thickness (nm) | 18 ± 3 | 7 ± 5 | 34 ± 5 | 58 ± 6 | 8 ± 3 | 16 ± 3 |
| shell vol. fraction | 0.35 ± 0.1 | 0.17 ± 0.07 | 0.55 ± 0.1 | 0.75 ± 0.08 | 0.19 ± 0.1 | 0.33 ± 0.1 |

*Particle size was measured by SEM images.*
interactions between NPs in a colloidal cluster.\textsuperscript{47–53} The (1,2) peak position clearly shifts depending on the chemical composition and thickness of the shell as indicated by the two dashed vertical lines in Figure 1a. For example, the \( f(1,2) \) of PS–PBMA undergo a red shift compared to the quasi-core–shell PS–PS NPs. The blue (red) shift in the spectra implies a stiffening (softening) of the NP modulus. Interestingly, the enhanced modulus of the PS–PMMA core–shell NPs is independent of the PMMA shell layer thickness (see Figure S1). For the PS–PBMA NPs, the red shift of (1,2) increases with the thickness of the PBMA shell layer from PS–PMMA to PS–PBMA\(_{38} \) (Figure S1). This trend reflects a reduction in the NP modulus as a function of the PBMA shell layer thickness, as discussed in detail below.

To account for the thermal phonon population and better reveal the low-frequency side of the BLS spectra near the elastic Rayleigh peak, Figure 1b displays the power spectra \( I f^2 \). A clear low-frequency peak (arrows) appears in the power spectrum. This (1,1) mode\textsuperscript{\textsuperscript{54}} is related to longitudinal phonons in the cluster of the NPs induced by their mutual adhesion, and its frequency \( f(1,1) \) can be described by the Johnson–Kendall–Roberts model.\textsuperscript{54} The precise \( f(1,1) \) position is defined as the cutoff frequency in the right tail where the intensity is 20% of the \( f(1,1) \) maximum intensity in the power spectrum. When this peak position is plotted against diameter, it follows the scaling \( f(1,1) \sim d^{-7/6} \) as shown in Figure S2. Although all core–shell NPs follow the scaling because of a similar adhesion energy,\textsuperscript{\textsuperscript{54}} PS–PBMA\(_{38} \) data deviated from the solid line, suggesting a higher adhesion energy.

We have recently reported that \( f(1,2) = 2f_1 - f_2 \) can account for the effect of interparticle interactions, providing a good approximation of the frequency \( f(1,2) \) of the single (1,2) mode of individual NPs \( (f_1 = f_2) \).\textsuperscript{52} To estimate the shear modulus \( G \) of NPs, the equation

\[
C_t = f_1 f(1,2) \cdot d/A = \sqrt{G/\rho}
\]

was applied.\textsuperscript{55} Here, \( C_t \) is the effective transverse sound velocity of NPs, \( d \) is the diameter, and \( \rho \) is the density of particles. The value of the Lamb constant, \( A_t \), depends on the Poisson ratio (Figure S3), and for the present case, \( A_t = 0.84 \) for PS spheres was used. Figure 2 depicts the \( G \) vs shell thickness plot for the studied NPs (the plot for \( G \) vs shell volume fraction is shown Figure S4). \( C_t \) and \( G \) values for the NPs are listed in Table S1. Note that given the high frequency of the NP vibrations, \( G \) refers to the elastic (frequency-independent) modulus. As the thickness of the soft PBMA shell increases, \( C_t \) and \( G \) decrease. In Figure 2, the estimated shear modulus of the NPs is plotted against shell thickness. Based on previous studies,\textsuperscript{38,52} the particle size does not affect the shear modulus, but the chemical composition plays a more important role in determining the NP shear modulus in homogeneous NPs. PS and the quasi-PS–PS core–shell NPs have a similar shear modulus in spite of their different particle sizes and polymerization processes. The shell-forming process has no discernible effect on the shear modulus, but the chemical composition of the shell can impact the shear modulus of NPs.\textsuperscript{38} PS NPs with a high-modulus PMMA shell manifest about a 10% higher modulus than their precursor PS core even with the addition of a thin (8 nm), harder shell layer. The hardening trend is also observed for the PS–PMMA\(_{16} \) NPs, but it is mainly due to the increase of density as the NPs’ \( C_t \) remains virtually constant; there is no significant blue shift for PS–PMMA\(_{16} \) (see Figure S1). Compared to PS–PS NPs, a lower shear modulus was revealed for PS–PBMA NPs. The NP modulus decreased with increasing PBMA thickness, reaching a value \( \sim 12\% \) lower than that of PS–PS NPs for a 58 nm PBMA shell (see Figure 2). The enhancement of the modulus in the case of the PS–PMMA NP corroborates the notion that a thin shell layer of an

Figure 1. (a) Anti-Stokes side of the Brillouin light-scattering (BLS) vibrational spectra of the shell architecture of PS–PS, PS–PMMA\(_{34} \), PS–PBMA\(_{34} \), and PS–PBMA\(_{38} \) vs reduced frequency, \( f_d \), with \( d \) being the particle diameter. The frequencies \( f_1, f_2 \), denote the spectral split of the (1,2) mode and the vertical dotted lines indicate \( f_1, f_3, \) and \( f(1,3) \) of the PS–PS NPs. (b) BLS power spectra, \( I f^2 \), vs frequency, \( f_1 \) at 294 K. The arrows indicate the interaction-induced (1,1) branch.

Figure 2. Shear modulus, \( G \), of PS and core–shell architecture, PS–PS, the PS–PMMA group (PS–PMMA\(_{16} \), PS–PMMA\(_{18} \), and PS–PMMA\(_{34} \)), and the PS–PMMA group (PS–PMMA\(_{34} \), and PS–PMMA\(_{38} \)), NPs plotted against the shell thickness, \( k \). The dashed straight line indicates the shear modulus for the bulk PBMA, whereas the open square and solid circles refer to the shear modulus of the precursor PS and PS–PMMA core–shell nanoparticles.\textsuperscript{38} The red triangles refer to the shear modulus of PS–PBMA core-shell NPs. The shaded area indicates the maximum error for the modulus of PS. The red and green solid lines indicate the calculated shear modulus for PS–PBMA and PS–PMMA core–shell NPs at a constant PS core diameter \( (d = 220 \text{ nm}) \) using bulk moduli values for the PS core and two different values for the PMMA \( (C_t = 1500 \text{ and } 1400 \text{ m/s}) \) and PBMA \( (C_t = 900 \text{ and } 1050 \text{ m/s}) \) shells. A similar plot for \( G \) but as function of the shell volume fraction is shown in Figure S4 in the Supporting Information.
elastically different material, atop the PS core, can significantly modify the core–shell NP elasticity.

The finite element method (FEM) was used to calculate the shear modulus of the PS–PMMA and PS–PBMA NPs assuming no interactions among NPs and perfect bonding between the core and shell interface and bulk elastic properties of each component (see Figure S5). In this case, the (1,2) peak is single in contrast to the interaction-induced double-spectral shape (see Figures 1 and S1, S5). Using \( C_t = 1400 \text{ m/s} \) for PMMA and \( C_t = 1050 \text{ m/s} \) for PBMA, \( G \) is presented by the lower green and upper red lines in Figure 2. For PBMA, \( G \) was obtained from the experimental longitudinal value \( C_l = 2200 \text{ m/s} \) and using the value 0.35 for the Poisson ratio. The agreement is only qualitative, since the calculated \( G \) overestimates the experimental value of \( G \) for PS–PBMA (upper red line, \( C_l = 1050 \text{ m/s} \)) and underestimates the \( G \) value for the two PS–PMMA NPs (lower green line, \( C_l = 1400 \text{ m/s} \)). To obtain quantitative agreement between FEM calculations and the experiment, the transverse sound velocity \( C_t \) for the PBMA shell should deviate from its bulk value. The lower red line in Figure 2 corresponds to \( C_t = 900 \text{ m/s} \), being about 15% lower than that in the bulk PBMA. The deviation between the assumed PBMA shell and PBMA bulk \( C_t \) values decreases with increasing PBMA shell thickness as indicated by the deviation of the experimental \( G \) values from the two red lines. For a thinner PBMA shell, PS–PBMA, the \( C_t \) value (shell) is about 15% lower than that in the bulk PBMA (1050 m/s), whereas for PS–PBMA, its elastic \( G \) can be captured with the bulk PBMA \( C_t \). In analogy, for the PS–PMMA NPs, a higher \( C_t \) (>1500 m/s) than for the bulk PMMA is required as indicated by the upper green line.

Although for colloids there are no other experiments on the elastic modulus, the impact of confinement on the Young modulus of polymer thin films has been investigated by buckling instability techniques, \(^{13,33-35} \) atomic force microscopy (AFM), \(^{23,37,58-59} \) and a nanobubble inflation method. \(^{32} \) A reduction in the glassy modulus has been reported for thin (<40 nm) PS films atop elastic poly(dimethylsiloxane) substrates and has been assigned to a free surface layer with a reduced modulus. \(^{33,57} \) An enhancement of the glassy modulus was reported near interfaces when thin polymer films were supported on hard substrates (e.g., silicon, glass). \(^{23,37,58-59} \) Quasi-static finite-element simulations \(^{59} \) of the AFM experiment assuming elastic properties of the substrate and the glassy polymer (PMMA) film indicate stiffening near the rigid substrate. Very recent molecular dynamics simulations \(^{60} \) have further suggested that the stiffness gradient length scale increases with temperature from below to above the polymer glass transition. Both simulations are based on several elastic and energetic parameters for the polymer interactions with the substrate and the indenter. Hence, the determining factor for either reduction or enhancement of the modulus for film–substrate systems is still unclear. Another system of polymer nanofibers such as PS and Nylon-11 shows a dramatic elastic modulus increase as the diameter drops below 200 nm. \(^{61,62} \) Very recently, the enhanced elasticity in films of polymer-tethered nanoparticles at a low grafting density was attributed to strong polymer–polymer interactions compared to densely tethered NPs with short chains. \(^{63} \) For the spherical confinement of Figure 2, the NP shear modulus can be rationalized assuming hardening for the PMMA and softening for the PBMA shell atop the same PS core.

**Nanoparticle Surface Mobility and Glass Transition.** The sensitive dependence of the (1,2) branch on the NP adhesion was utilized to directly probe the NP surface mobility and identify \( T_s \) as the temperature at which a crossover from the red to blue frequency shift was observed. \(^{52} \) It originates from the activation of the outermost surface mobility of NPs. \(^{56,57} \) At \( T_s \), the surface of the polymer particles undergoes a glass-to-rubber crossover. At temperatures above \( T_s \), interparticle interactions become activated and thereby increase the contact area between NPs. \(^{52} \) The resulting reinforced adhesion force is at the origin of the \( f(1,2) \) blue shift above the \( T_s \). The presence of a mobile surface layer was first reported by the whispering gallery mode light localization, which revealed a negative optical birefringence. \(^{54} \) The frequencies \( f_1 \) and \( f_2 \) of the (1,2) branch obtained by the representation of the (1,2) peak in BLS spectra by a double Lorentzian line shape are shown in Figure 3a for the PS–PS,
spectrum displayed only the longitudinal acoustic peak. In the case of PS–PBMA NPs, the incompatibility of the two components preserves the PS cores from merging within the annealed PBMA matrix, as indicated by the SEM images (see Figure S9). For PS–PS NPs, both \( T_g (\sim 352 \, \text{K}) \) and \( T_g (\sim 371 \, \text{K}) \) assume the same values for the PS NPs in our previous work.\(^{38} \) This indicates that the performed polymerization process, necessary for the formation of the shell, does not influence \( T_g \) or \( T_g \).

A thin (8 nm) PMMA shell atop PS NPs causes an increase of both \( T_g (\sim 8 \, \text{K}) \) and \( T_g (\sim 5 \, \text{K}) \) compared to those of PS–PS NPs. As in the case of \( G \), both transition temperatures are robust upon further increase of the PMMA thickness (16 nm) as the \( T_g \) has already reached the value for bulk PMMA. From the two characteristic temperatures of the NPs, only a single \( T_g \) can be measured by DSC as shown in Figure 3b. This \( T_g \) assumes the same value in both PS–PMMA\(_{8} \) NPs in agreement with the BLS, and it is indistinguishable, within experimental error, from the glass-transition temperature of PS–PS. For PS on PMMA bilayer films\(^{37} \) experiencing hard confinement next to PMMA (higher \( T_g \) than PS), the \( T_g \) in the PS domain deviated from the bulk \( T_g \) of PS below about 100 nm away from the PS/PMMA interface. A local \( T_g \) elevation of about 7 K was reported 25 nm from the PMMA layer, but no data exist for PMMA on the PS film geometry. The relatively small difference (about 15 K) in the bulk \( T_g \) between PS and PMMA limits the resolution of possible dynamic asymmetries in the PS core vs the PMMA shell. For a comparison, \( \Delta T_g = T_g (\text{PS}) - T_g (\text{PBMA}) \approx 80 \, \text{K} \) between the PS and PBMA components.

For PS–PBMA NPs, the large dynamic asymmetry between the core and shell revealed (i) a strong dependence of both \( T_g \) and \( T_g \) (see Figures 3a and S6a) and (ii) low and high glass-transition temperatures (\( T_g \), \( T_g \)) (see Figure 3b) on the PMMA thickness as in the case of \( G \) (see Figure 2), but (iii) a robust value for \( T_g \) = \( T_g (\text{PS}) \) BLS can access only \( T_g \) and \( T_g \) due to the disappearance of the NP vibration at \( T = T_g < T_g \) (see Figures 3a and S6), whereas DSC probes both glass transitions (see Figure 3b). These characteristic temperatures for all systems are compiled in Table S2. The glass-transition enhancement and the relation between \( T_g \) and \( T_g \) as a function of the proximity to the PS surface is illustrated in the lower and upper panels of Figure 4. To compare data of systems with different geometries, we plot normalized enhancements, \( \Delta T_g = (T_g (\text{PS}) - T_{g,\text{PBMA}})/(T_g (\text{PS}) - T_{g,\text{PBMA}}) \), where \( T_g (\text{PS}) \) is either \( T_g \) (filled red circles) or high \( T_g \) (blue solid circles). To appreciate the actual \( T_g \) changes, we have replotted Figure 4 as \( T_g \) vs. \( h \) in Figure S10.

The robustness of the PS \( T_g \) (solid blue circles in Figure 4b) is an unexpected result in view of the flat PBMA/PS bilayer films. For the latter, the glass-transition temperature was determined by fluorescence intensity measurements of pyrene-labeled layers (PS or PBMA) inserted at different distances, \( h \), (dashed line in the insets in Figure 4b) from the PS/PBMA interface (solid line in the upper bilayer model).\(^{26} \) The \( T_g \) in the PS layer softens with proximity to the PBMA interface as indicated in Figure 4b (blue open triangles) and reaches the bulk PS value at about 200 nm from the interface. Alternatively, the \( T_g \) in the PBMA layer is enhanced with increasing distance from the interface, but the dependence is weaker than that for the PS layer (open red squares). For the spherical geometry, the enhancement of the \( T_g \) in the PBMA shell is stronger than that in the corresponding bilayer film (red circles vs red squares) but shows a similar trend. The representation of the \( T_g (h) \) dependence by an exponential decay (red solid line) cannot describe the initial \( T_g \) softening (\( h < 8 \, \text{nm} \)) as indicated by the intercept (\( h = 0 \)) value (\( <1 \)). Hence, the glass dynamics appears to reflect a topology effect since in bilayer films the layers are less confined along the direction parallel to the film. The higher \( T_g \) of PBMA even at about 60 nm thickness is apparent from the robust surface of PS–PBMA NPs compared to that of PBMA NPs with \( T_g = 293 \, \text{K} \).

The huge \( T_g \) enhancement effect in the PBMA shell suggests that free surface effects are significantly counter-balanced by the influence of the polymer–polymer interface. This is supported by the observation of very similar DSC \( T_g \)’s in the PS–PBMA NPs and the corresponding contiguous films of PS (cores) in the PBMA matrix (no free interface) in Figure S11. This effect mimics that observed in polymers tightly adsorbed on solid interfaces. Here, the effect of the free surface is erased by the presence of such an adsorbed layer.\(^{26} \) On the same line of reasoning, we can consider the (soft)polymer/(hard) polymer interface as analogous to that of polymer/hard substrates, where the analogy is marked by the presence of extensive van der Waals interaction involving several pinpoints on each polymer chain, whose effects propagate to the free interface.\(^{60,65−68} \) However, it is worth noting that polymers in contact with an inorganic interface at best exhibit the bulk \( T_g \) or a \( T_g \) only a few degrees higher than that of the bulk. Therefore, the large \( T_g \) enhancement observed here may be at least partly explained by the presence of an interfacial mixed

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**Figure 4.** (a) Difference between the low glass-transition and the softening transition temperatures for the PS–PS and three PS–PBMA NPs depending on the shell thickness, \( h \). (b) Normalized glass-transition temperature contrast, \( \Delta T_g = (T_g (h) - T_{g,\text{PBMA}})/(T_g (\text{PS}) - T_{g,\text{PBMA}}) \), for the low \( T_g \) (blue solid circles) and high \( T_g \) (red solid line) glass transitions in PS–PBMA as a function of the shell thickness. Note that \( T_g \) refers to the PS core bearing a PBMA shell of thickness, \( h \). The red solid line denotes exponential decay excluding the data point for PS–PS (\( \Delta T_g = 1 \)). To display the actual \( T_g \) change, \( T_g (h) \) is shown in Figure S10 in the Supporting Information. The comparison with a flat PS/PBMA bilayer film is indicated by the red square symbols for the \( T_g (\text{PBMA}) \) in PBMA on the PS layer, whereas the blue triangle symbols are for \( T_g (\text{PS}) \) on the PBMA layer.\(^{26} \) \( h \) refers to the distance from the PS/PBMA interface.
layer. The presence of such a layer has been demonstrated\textsuperscript{69} and was found to be about 8 nm. This thickness may explain the very large $T_g$ deviation found for PS–PBMA$_h$. The effect of the free (PBMA) surface is, however, evident in the NP’s thermal behavior as indicated by the presence of the softening $T_g (<T_g)$. It does relate to $T_g$ and the difference $T_g(T_h - T_g)$ ~ 15 K seems to be virtually independent of the PBMA thickness as shown in Figure 4a, suggesting a constant free surface contribution to $T_g$.

## CONCLUSIONS

We synthesized core–shell nanoparticles with a PS core and low-$T_g$ PBMA or high-$T_g$ PMMA shell layer of varying thicknesses. We used Brillouin light scattering to measure the vibrational frequency spectrum of each nanoparticle type. We found that PS–PMMA core–shell particles exhibited a small, shell-thickness-independent increase in the elastic modulus. The elastic modulus of PS–PBMA core–shell particles decreased as the PBMA shell thickness increased. The change in the elastic modulus of the core–shell particles is not well-described by a finite-element analysis assuming bulk moduli for each component. Our results are more consistent with modulus hardening in the PMMA shell layer and softening in the PBMA shell layer under the spherical confinement induced by the NP core–shell geometry.

Temperature-dependent Brillouin light spectroscopy reports the softening temperature as an increase in the vibrational frequency. Additionally, modulated DSC was used to measure the $T_g$ of the nanoparticles. Both the softening temperature and $T_g$ for PS–PMMA core–shell particles increased slightly, consistent with the ~5 K higher PS–PMMA $T_g$ compared with the PS $T_g$. The softening temperature dramatically decreased for PS–PBMA particles with thick shells. Modulated DSC measured two $T_g$’s for PS–PBMA NPs: a low-shell-thickness-dependent $T_g$ for the PBMA shell and a high, constant $T_g$ for the PS core. The softening temperature measured via BLS was consistently ~15 K below the low $T_g$ measured by mDSC, suggesting that BLS is less sensitive to the shell of the nanoparticle and that the PBMA shell layer has enhanced mobility at the free surface. However, the PS–PBMA interaction dominates the mDSC $T_g$ measured for the PBMA shell layer. Finally, according to a recently published article,\textsuperscript{28} on supported flat bilayer films, the $T_g$ of PS with a rubbery tocastop is affected only for PS thicknesses below about 60 nm. This is consistent with the unaffected $T_g$ of the PS NPs ($d ~ 220$ nm) with a PBMA shell. In contrast, the $T_g$ of PS on the top of an unsupported free-PBMA film\textsuperscript{26} is unaffected only for PS thicknesses above 200 nm.

## ASSOCIATED CONTENT

* Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.macromol.9b00766.

BLS spectra of all core–shell NPs (PS–PS, PS–PBMA$_{n}$, PS–PMMA$_{h}$, PS–PBMA$_{m}^h$, PS–PMMA$_{n}^h$, and PS–PMMA$_{h}$), relation of the interaction-induced mode, shear modulus data from characteristic particle vibration frequencies, $T_g$ and $T_f$ from BLS and mDSC, comparison of experimental data with FEM, temperature dependency of eigenfrequencies of all core–shell NPs, heat maps of PS–PS and PS–PBMA$_{n}$ groups, and mDSC thermograms (PDF)

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