pH-Induced Association and Dissociation of Intermolecular Complexes Formed by Hydrogen Bonding between Diblock Copolymers

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Abstract: Poly(sodium styrenesulfonate)–block–poly(acrylic acid) (PNaSS–b–PAA) and poly(sodium styrenesulfonate)–block–poly(N-isopropylacrylamide) (PNaSS–b–PNIPAM) were prepared via reversible addition–fragmentation chain transfer (RAFT) radical polymerization using a PNaSS-based macro-chain transfer agent. The molecular weight distributions ($M_w/M_n$) of PNaSS–b–PAA and PNaSS–b–PNIPAM were 1.18 and 1.39, respectively, suggesting that these polymers have controlled structures. When aqueous solutions of PNaSS–b–PAA and PNaSS–b–PNIPAM were mixed under acidic conditions, water-soluble PNaSS–b–PAA/PNaSS–b–PNIPAM complexes were formed as a result of hydrogen bonding interactions between the pendant carboxylic acids in the PAA block and the pendant amide groups in the PNIPAM block. The complex was characterized by $^1$H NMR, dynamic light scattering, static light scattering, and transmission electron microscope measurements. The light scattering intensity of the complex depended on the mixing ratio of PNaSS–b–PAA and PNaSS–b–PNIPAM. When the molar ratio of the N-isopropylacrylamide (NIPAM) and acrylic acid (AA) units was near unity, the light scattering intensity reached a maximum, indicating stoichiometric complex formation. The complex dissociated at a pH higher than 4.0 because the hydrogen bonding interactions disappeared due to deprotonation of the pendant carboxylic acids in the PAA block.

Keywords: block copolymers; RAFT polymerization; complex; hydrogen bonding interactions; pH-responsive

1. Introduction

Non-covalent interactions, such as hydrophobic [1], electrostatic [2,3], van der Waals [4], and hydrogen bonding interactions [5,6], can be a driving force for the complex formation of polymers. In particular, hydrogen bonding interactions are an important driving force for the self-organization of natural polymers such as polysaccharides, proteins, and deoxyribonucleic acid (DNA).

It is known that hydrogen bonding interactions between amide groups or poly(ethylene glycol) (PEG) and carboxylic acid groups promote self-association or complex formation in water [7–9]. Shieh et al. [10] prepared a series of poly(N-isopropylacrylamide-random-acrylic acid) (PNIPAM–r–AA) copolymers and examined their glass transition behavior. They found that the incorporation of acrylic acid units into the PNIPAM polymer enhances the glass transition temperature ($T_g$) due to intermolecular hydrogen bonding between the pendant isopropyl amide groups and the carboxylic acid groups using $^1$H NMR and Fourier-transform infrared analyses. As an example of such complex formation,
Bian et al. [11] investigated the interaction between poly(N,N-diethylacrylamide) (PDEA), which is an analog of PNIPAM, and poly(acrylic acid) (PAA). The complex is formed between the two polymers through hydrogen bonding interactions with a stoichiometry of \( r = 0.6 \) (\( r \) is the unit molar ratio of PAA/PDEA), and the complex formation depends on pH values. We lately reported [12] the complex formation behavior of poly(sodium stylenesulfonate)-block-PAA/PNaSS–block–poly(sodium stylenesulfonate) (PNaSS–b–PEG–b–PNaSS) with poly(methacrylic acid) (PMA). Both PNaSS–b–PEG–b–PNaSS and PMA were synthesized via reversible addition–fragmentation chain transfer (RAFT) radical polymerization. Below pH 5, water-soluble complexes were formed owing to the hydrogen bonding interactions between the PEG block in PNaSS–b–PEG–b–PNaSS and the carboxylic acids in PMA. The experimental data indicated that the PNaSS–b–PEG–b–PNaSS/PMA complex was spherical in shape. At pH greater than 5, the complex dissociated because the hydrogen bonding interaction disappeared due to deprotonation of the pendant carboxylic acids in PMA.

In the present study, we focused on the complex formation behavior owing to the hydrogen bonding interactions as a function of the solution temperature instead of the solution pH. Two species of diblock copolymers (Figure 1a), PNaSS–block–poly(acrylic acid) (PNaSS–b–PAA) and PNaSS–block–PNIPAM (PNaSS–b–PNIPAM), were synthesized via RAFT radical polymerization [13]. The PNIPAM blocks of PNaSS–b–PNIPAM are also assumed to associate above the lower critical solution temperature (LCST) to form core–corona-type multi-polymer micelles [14]. When PNaSS–b–PAA and PNaSS–b–PNIPAM were mixed in a 0.1 M NaCl aqueous solution at 20 °C, water-soluble PNaSS–b–PAA/PNaSS–b–PNIPAM complexes were formed through hydrogen bonding interactions between the PAA and PNIPAM blocks below pH 3.9 (Figure 1b). The complexes were maintained above the LCST of PNaSS–b–PNIPAM. The complexes dissociated under basic conditions when the hydrogen bonding interactions disappeared due to deprotonation of the pendant carboxylic acids in the PAA block.

![Figure 1](image-url)

**Figure 1.** (a) Chemical structures of diblock copolymers used in this study: Poly(sodium styrenesulfonate)\(_{58}\)-block-poly(acrylic acid)\(_{125}\) (PNaSS\(_{58}\)-b-PAA\(_{125}\)) and poly(sodium styrenesulfonate)\(_{58}\)-block-poly(N-isopropylacrylamide)\(_{115}\) (PNaSS\(_{58}\)-b-PNIPAM\(_{115}\)), and (b) schematic representation of polymer chain mixing in the core and corona micelles comprising PNaSS\(_{58}\)-b-PAA\(_{125}\) and PNaSS\(_{58}\)-b-PNIPAM\(_{115}\).

2. Experimental Section

2.1. Materials

Acrylic acid (AA) from Kanto Chemical (Tokyo, Japan) was dried over 4 Å molecular sieves and distilled under reduced pressure. \( N \)-isopropylacrylamide (NIPAM) from Aldrich (St. Louis, MO, USA) was purified by recrystallization from a mixed solvent of benzene and \( n \)-hexane. \( \alpha \)-Methyltrithiocarbonate-S-phenylaceticacid (MTPA) was synthesized as previously
reported [15]. Sodium styrenesulfonate (NaSS) from Tokyo Chemical Industry (Tokyo, Japan) and 4,4′-azobis(4-cyanopentanoic acid) (V-501) from Aldrich were used as received. Methanol was dried using molecular sieves and distilled. Deionized water was used.

2.2. Synthesis of PNaSS Macro-Chain Transfer Agent (PNaSS Macro–CTA)

NaSS (20.6 g, 100 mmol) and MTPA (0.26 g, 1.0 mmol) were dissolved in 180 mL of water, and V-501 (5.6 mg, 0.2 mmol) was added to the aqueous solution. Polymerization was carried out at 70 °C for 3 h under Ar atmosphere. After the polymerization, the mixture was dialyzed against pure water for a week and recovered by a freeze-drying technique (yield 14.8 g, number-average molecular weight \(M_n = 1.22 \times 10^4\) (gel permeation chromatography (GPC)), molecular weight distribution \(M_w/M_n = 1.19\), and degree of polymerization (DP) = 58). The obtained PNaSS could be used as a macro-CTA (PNaSS\textsubscript{58} macro–CTA).

2.3. Preparation of PNaSS\textsubscript{58}–b–P AA\textsubscript{125}

AA (0.86 g, 12 mmol) was dissolved in 15 mL of water, and PNaSS\textsubscript{58} macro–CTA (1.05 g, 0.087 mmol) and V-501 (5.0 mg, 0.018 mmol) were added to this solution. The mixture was deoxygenated by purging with Ar gas for 30 min. Block copolymerization was carried out at 70 °C for 2 h. The diblock copolymer was purified by dialysis against pure water for a week and then recovered using a freeze-drying technique. The diblock copolymer, PNaSS\textsubscript{58}–b–PAA\textsubscript{125}, was obtained (yield 1.79 g, \(M_n = 2.12 \times 10^4\) (\(^1\)H NMR), \(M_w/M_n = 1.18\), and DP of the PAA block = 125).

2.4. Preparation of PNaSS\textsubscript{58}–b–PNIPAM\textsubscript{115}

NIPAM (2.26 g, 10.2 mmol) was dissolved in a mixed solvent of water and methanol (12.4 mL, 1/1, v/v), and PNaSS\textsubscript{58} macro–CTA (0.96 g, 0.079 mmol) and V-501 (4.6 mg, 0.016 mmol) were added to this solution. The mixture was deoxygenated by purging with Ar gas for 30 min. Block copolymerization was carried out at 60 °C for 5 h. The diblock copolymer was purified by reprecipitation from a methanol solution into excess ether twice and then recovered using a freeze-drying technique. The diblock copolymer PNaSS\textsubscript{58}–b–PNIPAM\textsubscript{115} was obtained (yield 1.79 g, \(M_n = 2.52 \times 10^4\) (\(^1\)H NMR), \(M_w/M_n = 1.39\), and DP of the PNIPAM block = 115).

2.5. Preparation of the Water-Soluble Complex

Stock solutions of PNaSS\textsubscript{58}–b–PAA\textsubscript{125} and PNaSS\textsubscript{58}–b–PNIPAM\textsubscript{115} were prepared by dissolving each polymer in 0.1 M NaCl aqueous solutions of pH 3 and 10, respectively. To prepare the complex, the PNaSS\textsubscript{58}–b–PNIPAM\textsubscript{115} aqueous solution was added to the PNaSS\textsubscript{58}–b–PAA\textsubscript{125} aqueous solution over a period of five min, and the mixture was allowed to stand still for one day. The mixing ratio of the two diblock copolymers was adjusted based on the molar fraction of NIPAM units \(f_{\text{NIPAM}} = (\text{NIPAM})/((\text{NIPAM}) + (\text{AA}))\), where (NIPAM) and (AA) are the molar concentrations of NIPAM and AA units, respectively). The complex was prepared at \(f_{\text{NIPAM}} = 0.5\) unless otherwise noted.

2.6. Measurements

GPC measurements were performed with a Shodex 7.0 µm bead size GF-7F HQ column. A phosphate buffer at pH 9, containing 10 vol % acetonitrile was used as an eluent at a flow rate of 0.6 mL/min at 40 °C. \(M_n\) and \(M_w/M_n\) were calibrated with standard PNaSS samples of 11 different molecular weights ranging from \(1.37 \times 10^3\) to \(2.61 \times 10^6\). \(^1\)H NMR spectra were obtained with a Bruker DRX-500 spectrometer (Buick Rica, MA, USA) operating at 500 MHz. Light scattering measurements were performed using an Otsuka Electronics Photal DLS-7000DL equipped with a digital time correlator (ALV-5000E, Osaka, Japan). Sample solutions were filtered with a 0.2-µm membrane filter. For dynamic light scattering (DLS) measurements, the data obtained were analyzed with ALV software version 3.0 [16,17]. For static light scattering (SLS) measurements, weight-average
molecular weight ($M_w$), and radius of gyration ($R_g$) were estimated from Zimm plots [18]. Values of $dn/dC_P$ were determined with an Otsuka Electronics Photal DRM-1020 differential refractometer. Transmission electron microscopy (TEM) measurements were carried out using a JEOL JEM-2100 (Tokyo, Japan) at an accelerating voltage of 200 kV. The TEM sample was prepared by placing one droplet of the solution on a copper grid coated with Formvar. The sample was stained by sodium phosphotungstate and dried under reduced pressure. Percent transmittance (%) measurements were performed using a JASCO V-630 BIO UV–Vis spectrometer (Tokyo, Japan) with a 10 mm path length quartz cell. The temperature was increased from 20 to 80 °C with a heating rate of 1.0 °C-min⁻¹ using a JASCO ETC-717 thermostat system.

3. Results and Discussion

We prepared the diblock copolymers, PNaSS₈₈–b–PAA₁₁₅ and PNaSS₈₈–b–PNIPAM₁₁₅, via a RAFT technique using PNaSS₈₈ macro–CTA. The DP of PNaSS₈₈ macro–CTA, which was determined by GPC, was 58 (PNaSS₈₈–macro–CTA). The DPs of the PAA and PNIPAM blocks were 125 and 115, respectively. These values were calculated from $^1$H NMR peak area intensities derived from the PAA or PNIPAM block and an area derived from the PNaSS block. $M_n$ and $M_w / M_n$ of the diblock copolymers were estimated from GPC. The results for the characteristics of the diblock copolymers are listed in Table 1. The $M_w / M_n$ values are relatively small ($M_w / M_n < 1.4$), indicating that the controlled/living polymerizations proceeded successfully [13].

| Samples            | DP of PNaSS a | DP of PAA b | DP of PNIPAM b | $M_n$(theo) c | $M_w$(NMR) b $\times 10^{-4}$ | $M_w$(GPC) a $\times 10^{-4}$ | $M_w / M_n$ a |
|--------------------|---------------|-------------|----------------|--------------|-------------------------------|-------------------------------|---------------|
| PNaSS₈₈–b–PAA₁₁₅   | 58            | 125         |                | 2.07         | 2.12                          | 3.51                          | 1.18          |
| PNaSS₈₈–b–PNIPAM₁₁₅| 58            | 115         |                | 2.27         | 2.52                          | 1.21                          | 1.39          |

a Estimated from gel permeation chromatography (GPC) eluted with a phosphate buffer solution at pH 9 containing 10 vol % acetonitrile; b Estimated from $^1$H NMR; c Calculated from Equation (4).

When the polymerization is assumed an ideally living process, then the theoretical number-average molecular weight ($M_n$(theo)) can be estimated as

$$M_n$(theo) = \frac{[M]_0}{[CTA]_0} \times 100 \times M_m + M_{CTA} \tag{1}$$

where $[M]_0$ is the initial monomer concentration, $[CTA]_0$ is the initial PNaSS₈₈ macro–CTA concentration, $x_m$ is the conversion of the monomer, $M_m$ is the molecular weight of the monomer, and $M_{CTA}$ is the molecular weight of PNaSS₈₈ macro–CTA. The $M_n$(NMR) values for PNaSS₈₈–b–PAA₁₁₅ and PNaSS₈₈–b–PNIPAM₁₁₅ were calculated from the $^1$H NMR data. As shown in Table 1, the $M_n$(NMR) values for PNaSS₈₈–b–PAA₁₁₅ and PNaSS₈₈–b–PNIPAM₁₁₅ were in reasonable agreement with the $M_n$(theo) values. However, the $M_n$(theo) and $M_n$(GPC) values for both diblock copolymers were found to be slightly different. This may be because the volume-to-mass ratio for PNaSS is different from those for SS₈₈–b–PAA₁₁₅ and SS₈₈–b–PNIPAM₁₁₅ [19,20].

We attempted to monitor complex formation between PNaSS₈₈–b–PAA₁₁₅ and PNaSS₈₈–b–PNIPAM₁₁₅ induced by the solution pH using $^1$H NMR spectra measured in D₂O containing 0.1 M NaCl at pH 10 and 3. Figure 2 compares the $^1$H NMR spectra of mixed solutions at pH 10 and 3. In Figure 2a, the resonance peaks at 1.16 (f) and 3.91 ppm (c) are attributed to the methyl and methine protons, respectively, in the pendant isopropyl group of the PNIPAM block are observed. Moreover, the resonance bands owing to the pendant phenyl protons in the PNaSS block are also detected at 6.1–7.9 ppm (a and b). The resonance bands in the 1.28–2.38 ppm region (d and e) are attributed to the sum of the main chain of the diblock copolymers. At pH 3, the intensities of the resonance peaks f and
c derived from the PNIPAM block remarkably decreased, whereas those of a and b remained intact, as shown in Figure 2b. Besides this result, the resonance bands in the 1.28–2.38 ppm region (d and e) became broad. From the significant reduction of the motional freedom for the PNIPAM block at pH 3, one can predict that the complexes are formed from PNaSS$_{58}$–PAA$_{125}$ and PNaSS$_{58}$–PNIPAM$_{115}$ owing to the hydrogen bonding interactions between the pendant carboxylic acids in the PAA block and the amide groups in the PNIPAM block. At pH 10, however, the complexes dissociated owing to the disappearance of the hydrogen bonding interactions as a result of deprotonation of the carboxylic acids in the PAA block.

**Figure 2.** $^1$H NMR spectra and peak assignment of the mixture of PNaSS$_{58}$–PAA$_{125}$ and PNaSS$_{58}$–PNIPAM$_{115}$ in D$_2$O containing 0.1 M NaCl at (a) pH 10 and (b) pH 3.

Figure 3 shows the light scattering intensities for a mixture of PNaSS$_{58}$–PAA$_{125}$ and PNaSS$_{58}$–PNIPAM$_{115}$ in 0.1 M NaCl at pH 3 as a function of $f_{\text{NIPAM}}$. The total polymer concentration was kept constant at 4.4 g/L. An increase in the scattered intensity suggests an increase in the size of the complex. The maximum scattering intensity was observed at $f_{\text{NIPAM}} = 0.5$. This result indicates that a stoichiometric interaction in the mixture of PNaSS$_{58}$–PAA$_{125}$ and PNaSS$_{58}$–PNIPAM$_{115}$ led to form a complex with largest aggregation number. The complex with $f_{\text{NIPAM}} = 0.5$ was studied unless otherwise stated.

**Figure 3.** Light scattering intensities of the PNaSS$_{58}$–PAA$_{125}$/PNaSS$_{58}$–PNIPAM$_{115}$ complexes as a function of $f_{\text{NIPAM}}$ (= (NIPAM)/((NIPAM) + (AA))) in 0.1 M NaCl at pH 3. The total polymer concentration was fixed at 4.4 g/L.

Figure 4 shows the hydrodynamic radius ($R_h$) distributions for each diblock copolymer and the PNaSS$_{58}$–PAA$_{125}$/PNaSS$_{58}$–PNIPAM$_{115}$ complex at pH 3 and 10. The values of $R_h$ were determined by DLS in 0.1 M NaCl and are indicated in the figure. The $R_h$ values for PNaSS$_{58}$–PAA$_{125}$ and PNaSS$_{58}$–PNIPAM$_{115}$ were 3.5 and 3.7 nm, respectively, which are reasonable...
for a unimer state. The $R_h$ values of the complex at pH 3 and 10 were 15.0 and 3.7 nm, respectively. When PNaSS$_58$–b–PAA$_{125}$ and PNaSS$_58$–b–PNIPAM$_{115}$ were mixed at pH 3, a water-soluble complex was formed due to hydrogen bonding interactions. On the other hand, at pH 10 the complex dissociated because the hydrogen bonding interactions disappeared as a result of deprotonation of the pendant carboxylic acids.

In Figure 6a, the scattering intensity increases sharply with increases in the concentration above a critical concentration of 0.9 g/L. Figure 6b indicates that the $R_h$ values for the complex are practically constant ($R_h$ is approximately 15 nm) in the concentration range from 0.9 to 4.4 g/L. The results suggest that the formation of the complex between PNaSS$_{58}$–b–PAA$_{125}$ and PNaSS$_{58}$–b–PNIPAM$_{115}$ starts to occur above a threshold of 0.9 g/L. Figure 6b indicates that the apparent critical aggregate concentration (CAC) value is approximately 15 nm) in the concentration range from 0.9 to 4.4 g/L. The results suggest that the relaxation modes are virtually diffusive [21]. Thus, the relaxation rates ($\Gamma$) measured at different scattering angles ($\theta$) are plotted as a function of the square of the magnitude of the scattering vector ($q^2$) for the mixture of PNaSS$_{58}$–b–PAA$_{125}$ and PNaSS$_{58}$–b–PNIPAM$_{115}$ in 0.1 M NaCl at pH 3 in Figure 5. A linear plot passing through the origin suggests that the relaxation modes are virtually diffusive [21]. Thus, the $R_h$ values can be estimated at a fixed $\theta$ of 90° as the angular dependence is negligible.

![Figure 4](image-url)  
**Figure 4.** Hydrodynamic radius ($R_h$) distributions for (a) PNaSS$_{58}$–b–PAA$_{125}$ and (b) PNaSS$_{58}$–b–PNIPAM$_{115}$ in 0.1 M NaCl at pH 3, and the PNaSS$_{58}$–b–PAA$_{125}$/PNaSS$_{58}$–b–PNIPAM$_{115}$ complex at (c) pH 3 and (d) pH 10.

![Figure 5](image-url)  
**Figure 5.** The relaxation rate ($\Gamma$) plotted as a function of the square of the magnitude of the scattering vector ($q^2$) for the PNaSS$_{58}$–b–PAA$_{125}$/PNaSS$_{58}$–b–PNIPAM$_{115}$ complex at $C_p = 4.4$ g/L in 0.1 M NaCl at pH 3.

Figure 6 shows the light scattering intensities and $R_h$ values for the mixture of PNaSS$_{58}$–b–PAA$_{125}$ and PNaSS$_{58}$–b–PNIPAM$_{115}$ in 0.1 M NaCl at pH 3 as a function of the concentration of the mixture. In Figure 6a, the scattering intensity increases sharply with increases in the concentration above a threshold of 0.9 g/L. Figure 6b indicates that the $R_h$ values for the complex are practically constant ($R_h$ is approximately 15 nm) in the concentration range from 0.9 to 4.4 g/L. The results suggest that the formation of the complex between PNaSS$_{58}$–b–PAA$_{125}$ and PNaSS$_{58}$–b–PNIPAM$_{115}$ starts to occur
above a critical concentration of 0.9 g/L. The apparent critical aggregate concentration (CAC) value for the complex, estimated from the plots, is 0.9 g/L.

![Graph showing light scattering intensity and hydrodynamic radius as a function of concentration](image)

**Figure 6.** (a) Light scattering intensity and (b) hydrodynamic radius ($R_h$) of the PNaSS$_{58}$–b–PAA$_{125}$/PNaSS$_{58}$–b–PNIPAM$_{115}$ complex as a function of the concentration of the mixture in 0.1 M NaCl at pH 3.

The apparent values of $M_w$ and $R_g$, determined by SLS measurements, are listed in Table 2. Figure 7 shows a Zimm plot for the PNaSS$_{58}$–b–PAA$_{125}$/PNaSS$_{58}$–b–PNIPAM$_{115}$ complex, which is formed from a mixture of PNaSS$_{58}$–b–PAA$_{125}$ and PNaSS$_{58}$–b–PNIPAM$_{115}$ in 0.1 M NaCl at pH 3. The aggregation number ($N_{agg}$) was defined as the number of polymer chains forming one complex, which can be estimated from the $M_w$ values of the complex and unimer. The result of this calculation gives an $N_{agg}$ of 46 for the complex. The chain numbers of PNaSS$_{58}$–b–PAA$_{125}$ and PNaSS$_{58}$–b–PNIPAM$_{115}$ for single complex are 22 and 24, respectively, as calculated from $f_{NIPAM} = 0.5$ and the DP values of PAA and PNIPAM.

**Table 2.** Light scattering data for the PNaSS$_{58}$–b–PAA$_{125}$/PNaSS$_{58}$–b–PNIPAM$_{115}$ complex.

| pH | $R_h$ a (nm) | $M_w$(SLS) b $\times 10^{-5}$ | $R_g$ b (nm) | $R_g/R_h$ | $N_{agg}$ c |
|----|-------------|-------------------------------|-------------|-----------|-------------|
| 3  | 15.0        | 9.23                          | 12.9        | 0.86      | 46          |
| 10 | 3.7         | 0.20                          | 13.6        | 3.7       | 1           |

* a Estimated from dynamic light scattering (DLS); b Estimated from static light scattering (SLS); c Aggregation number calculated from $M_w$ of the aggregate determined at pH 3 and $M_w$ of the corresponding unimer at pH 10 determined from SLS.
As the pH decreased,

The light scattering intensity was nearly constant between pH 3.5 and 2.5 suggests that not the aggregation number but the compactness of the complex was practically constant.

The average radius estimated from the TEM images for the complex was 13.4 nm, which is similar to the shells. The average radius estimated from the TEM images for the complex was 13.4 nm, which is similar to a homogeneous hard sphere is 0.778 but this value increases substantially for less dense structures and polydisperse mixtures; for example, 

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Figure 7. Zimm plot for the PNaSSb–PAA125/PNaSSb–PNIPAM115 complex with f_{NIPAM} = 0.5 in 0.1 M NaCl at pH 3. Scattering angles (θ) range from 30° to 130° in 20° increments.

The \( R_g/R_h \) value indicates the shape of the molecular assemblies. The theoretical \( R_g/R_h \) value of a homogeneous hard sphere is 0.778 but this value increases substantially for less dense structures and polydisperse mixtures; for example, \( R_g/R_h = 1.5–1.7 \) for flexible linear chains in good solvents, whereas \( R_g/R_h \geq 2 \) for a rigid rod [22–24]. The \( R_g/R_h \) ratio for the complex (Table 2) was 0.86, suggesting that the shape of the complex may be spherical. The \( R_g/R_h \) ratio for the unimer was 3.7, which indicates that the unimer was in a relatively expanded conformation with polydispersity.

To confirm the shape and size of the PNaSSb–PAA125/PNaSSb–PNIPAM115 complex at pH 3, TEM measurements were performed (Figure 8). The complex formed spherical objects with almost uniform contrast, suggesting that it comprises micelles with PAA/PNIPAM cores and PNaSS shells. The average radius estimated from the TEM images for the complex was 13.4 nm, which is similar to the \( R_h \) value estimated from DLS.

Figure 8. Transmission electron microscopy (TEM) image of the PNaSSb–PAA125/PNaSSb–PNIPAM115 complex at pH 3.

Figure 9a shows the light scattering intensities for the complex of PNaSSb–PAA125/PNaSSb–PNIPAM115 as a function of pH. The light scattering intensity increased rapidly as the pH decreased from 4.0 to 3.5, which suggests that the complex was formed below pH 3.5. The light scattering intensity was nearly constant between pH 3.5 and 2.5 suggests that \( N_{agg} \) was constant in this pH region because the light scattering intensity is proportional to the molecular mass. Figure 9b shows \( R_h \) values for the complex as a function of pH. Above pH 4.0, the \( R_h \) values were of the order of 3 nm, suggesting that PNaSSb–PAA125 and PNaSSb–PNIPAM115 were in a unimer state. As the pH decreased, \( R_h \) started to increase at around pH 4.0, reaching a maximum value of 15.0 nm at pH 3.5. As the pH continued to decrease, \( R_h \) was nearly constant between pH 3.5 and 2.5, suggesting that not the aggregation number but the compactness of the complex was practically constant.

When the pH value was increased from 3 to 10 and subsequently decreased back to 3, pH-induced \( R_h \) changes were found to be completely reversible. Figure 10 shows the pH-induced changes of the
$R_h$ value of the PNaSS$_{58}$–b–PAA$_{125}$/PNaSS$_{58}$–b–PNIPAM$_{115}$ complex in 0.1 M NaCl cycled between pH 3 and 10 with 20 min intervals. The changes in $R_h$ between the two pH values were completely reproducible, indicating that the association and dissociation of the complex caused by pH changes was reversible over many cycles. This result suggests that the complex may find applications as a pH-responsive controlled association–dissociation system.

![Figure 9](image1.png)

**Figure 9.** (a) Scattering intensity and (b) hydrodynamic radius ($R_h$) of the PNaSS$_{58}$–b–PAA$_{125}$/PNaSS$_{58}$–b–PNIPAM$_{115}$ complex at $C_p$ = 4.4 g/L in 0.1 M NaCl as a function of pH.

![Figure 10](image2.png)

**Figure 10.** The hydrodynamic radius ($R_h$) of the PNaSS$_{58}$–b–PAA$_{125}$/PNaSS$_{58}$–b–PNIPAM$_{115}$ complex at $C_p$ = 4.4 g/L in 0.1 M NaCl as a function of pH 3 (○) and pH 10 (△).

To confirm the formation of the PNaSS$_{58}$–b–PAA$_{125}$/PNaSS$_{58}$–b–PNIPAM$_{115}$ complex via hydrogen bonding interactions, urea was added to the mixture of PNaSS$_{58}$–b–PAA$_{125}$ and PNaSS$_{58}$–b–PNIPAM$_{115}$ in 0.1 M NaCl at pH 3. It is known that urea disturbs hydrogen bonding interactions [25]. Figure 11 shows the light scattering intensity for the PNaSS$_{58}$–b–PAA$_{125}$/PNaSS$_{58}$–b–PNIPAM$_{115}$ complex as a function of the concentration of urea. The light scattering intensity decreased as the concentration of urea increased from 0 to 3.0 mol/L, and the scattering intensity dropped below 0.1 MHz at 3.0 mol/L urea concentration. These observations are indicative of the complete dissociation of the complex caused by an excess amount of urea. Thus, the complex was confirmed to be formed via hydrogen bonding interactions.
The mixture at pH 10 also exhibited a slight decrease in %T change at 35–37 °C, which indicates the LCST. The mixture at pH 10 also exhibited a slight decrease in %T change at 35–37 °C. On the other hand, the mixture at pH 3 did not show any %T change in the temperature range from 20 to 80 °C. Ordinarily, the mechanism of LCST for PNIPAM can be explained as follows [26,27]. Below LCST, the PNIPAM chains are hydrated because the pendant amide groups form hydrogen bonding with water molecules, whereas above LCST, molecular motions prevail over the hydrogen bonding interactions resulting in the dehydration of the PNIPAM chains, thus leading to phase separation. Therefore, the PNIPAM chains dehydrated causing phase separation. In the case of the PNaSS58–b–PNIPAM115 complex at pH 3, the pendant amide groups in the PNIPAM block interact with the pendant carboxylic acid in the PAA block irrespective of the temperature. Hence, the pendant amide groups in the PNIPAM block are prevented from forming hydrogen bonds with water molecules. Therefore, the LCST of the complex at pH 3 cannot be observed. However, at pH 10, the mixture dissociated and hydrogen bonding interactions between the pendant amide groups in PNaSS58–b–PNIPAM115 and water molecules were formed at low temperature. Hence, the LCST can be observed for the mixture of PNaSS58–b–PAA125 and PNaSS58–b–PNIPAM115 at pH 10, although the decrease in %T at the LCST was small comparing to the PNaSS58–b–PNIPAM115 case. This is simply because the concentration of the NIPAM units in the former solution is lower than that in the latter solution.

**Figure 11.** The scattering intensity of the PNaSS58–b–PNIPAM115 diblock copolymer dissolves in water at room temperature, but it separates from aqueous solutions when heated above the lower critical solution temperature (LCST). Figure 12 shows the percent transmittance (%T) values monitored at 600 nm for 0.1 M NaCl aqueous solutions of PNaSS58–b–PNIPAM115 and a mixture of PNaSS58–b–PAA125 and PNaSS58–b–PNIPAM115 at pH 3 and 10, as a function of solution temperature. The diblock copolymer, PNaSS58–b–PNIPAM115, exhibited a significant %T change at 35–37 °C, which indicates the LCST.

**Figure 12.** Percent transmittance (%T) at 600 nm for 0.1 M NaCl aqueous solutions of PNaSS58–b–PNIPAM115 (△) and a mixture of PNaSS58–b–PAA125 and PNaSS58–b–PNIPAM115 at pH 3 (□) and pH 10 (○) as a function of solution temperature.
4. Conclusions

The diblock copolymers PNaSS$_{58}$–b–PAA$_{125}$ and PNaSS$_{58}$–b–PNIPAM$_{115}$ were prepared via RAFT-controlled radical polymerization using a PNaSS$_{58}$ macro–CTA. Both polymerizations proceeded by a controlled mechanism. A mixture of PNaSS$_{58}$–b–PAA$_{125}$ and PNaSS$_{58}$–b–PNIPAM$_{115}$ formed a water-soluble complex under acidic conditions. The formation of the complex was confirmed by various measurement techniques. The $^1$H NMR data indicated restricted motion of the PNIPAM block at pH 3 in the complex owing to hydrogen bonding interactions between the pendant carboxylic acids in the PAA block and the pendant amide groups in the PNIPAM block. The DLS and SLS data suggested that the PNaSS$_{58}$–b–PAA$_{125}$/PNaSS$_{58}$–b–PNIPAM$_{115}$ complex was spherical in shape. When urea was added to the complex aqueous solution, the complex dissociated. This observation indicates that the driving force for the formation of the complex is hydrogen bonding interactions. The $\%T$ data indicated that the LCST of the complex was not observed at pH 3, owing to the complex formation. However, the complex dissociated to a unimer above pH 4.0.

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