Aggregation Processes of a Weak Polyelectrolyte, Poly(allylamine) Hydrochloride

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Poly(allylamine) hydrochloride is a weak cationic polyelectrolyte that exhibits different aggregation properties at different solution pH values and aging times. Specifically, after several days aging in a pH 3 buffer, less than 1 mg/mL poly(allylamine) hydrochloride became turbid, and the hydrodynamic radius increased with a single diffusion mode. However, the hydrodynamic radius did not change at high concentrations. The dynamic processes of polymer aggregations at different pH values were verified by a light scattering and zeta-potential apparatus. The major interaction was caused by the capturing of counterions by the polyelectrolyte, which generates electrostatic, hydrophobic and cation–π interactions.

Key Words: Flocculation, Homogeneity parameter, Polyelectrolyte, Probe charge effect, pH dependence

Introduction

The study of polymer aggregation dynamics is very important in the formation of the polyelectrolyte multilayers through the layer by layer (LBL) and microcapsule templating method. The application of colloidal particles is often closely connected to the use of polyelectrolytes, which are widely used to stabilize or flocculate suspensions. For application in water purification, addition of small amounts of polymer results in attractive interactions, causing large floculants to be formed by bridging interactions. In the food and cosmetics industries, a suspension or emulsion is generally the end product. A bio-mimicking polymer with an NH₂ functional group, poly(allylamine) hydrochloride (PAH) has been vigorously studied for polystyrene sulfonate (PSS)/PAH multilayer formation due to the strong effect of salt concentration and pH on the conformational and mechanical properties. Composites consisting of a polystyrene (PS) core particle and PAH/PSS thin layer also showed potential as a drug delivery system. Although these composites have been widely used in several previous studies, little is known about the basic physical properties of weakly charged polyelectrolytes and hydrophobic anionic salts. The objective of this research was to reveal the intermolecular interaction and aggregation processes of a polyelectrolyte, PAH, with various polymer concentrations and aging times by dynamic light scattering (DLS) and zeta-potential measurements.

Experimental

Materials. PAH (Mw = 70,000) and potassium hydrogen phthalate were obtained from Aldrich. A pH 3 buffer was prepared from HCl and 0.1-M potassium hydrogen phthalate. A pH 10 buffer was prepared from NaOH and 0.025-M borax. NaCl was added to each solution to maintain the same salt concentration as 100 mM between pH 3 and pH 10 buffer solutions because ion, Na⁺, concentration is critical in the polyelectrolyte solutions.

Sample Preparation. Polyelectrolyte solutions were prepared by thoroughly mixing appropriate amounts (0.05-10 mg/mL) of PAH powder with pH 3, pH 10 buffer or DI-water. NaCl was used to maintain salt concentration of each buffer solution. Double distilled, salt free water from a MilliQ system was used as a preparative solution. The dissolved solutions were stirred for several days. DLS was used to observe changes in the physical properties over the course of aging time. After PAH was dissolved in each buffer, the solution was filtered through a 0.8-μm membrane directly into a pre-cleaned, light scattering cell (13 × 100 mm test tube). The measurements were performed daily for 30 days. For FT-IR analysis, the solution was dried in a vacuum and a dissolved solution was used for FT-Raman measurement.

Light Scattering Measurements. A Uniphase He-Ne laser with vertically polarized light of λ₀ = 632 nm was used. The detector optics employed optical fibers coupled to an ALV/SO-SIPD/DUAL detection unit with an EMI PM-28B power supply and ALV/PM-PD preamplifier/discriminator. The correlator was an ALV-5000/E/WIN multiple tau correlator with 288 exponentially spaced channels. The sampling time of the correlator ranged from 10⁻⁶ to 100 sec. The cylindrical scattering cell was located in an index bath matching solvent (decane) that was maintained at room temperature. The scattered light was passed through 400-μm and 200-μm pinholes. Two focal lenses and two Glan-Thomson polarizers were used. Each focal lens and polarizer pair were placed in front of the laser and pinhole. Each correlation function was gathered at six different angles from θ = 30° to 105°. For the static light scattering (SLS) experiments, intensities were measured from θ = 30° to 90° in vertically polarized incident and scattered beam (Vv) geometry.

Zeta Potential Measurements. The electrophoretic mo-
bility of PAH represented by globular coil or aggregation was determined by photon correlation spectroscopy on a Zetasizer Nano ZS (Malvern Instruments, Malvern, U.K.). Three rounds of assays were averaged. All measurements were performed at 25 °C. The mobility \( u \) was converted into \( \zeta \)-potential using the relation \( \zeta = \frac{uh}{e} \), where \( h \) is the viscosity of the solution and \( e \) is the permittivity of the water solution.

Results and Discussion

Dynamics of Aggregated PAH at Different pH Solutions. Figure 1 shows the images of PAH at different concentrations and age time. PAH and potassium hydrogen phthalate were chosen as the cationic polyelectrolyte and hydrophobic anionic salt, respectively. The aggregation behaviors depend on the nature of the ionic groups of the polyelectrolytes and salt. The salts in the pH 3 and 10 buffers were potassium hydrogen phthalate and borax, respectively. In this system the salt concentration was maintained and the only difference was the polymer concentration, achieved by adding PAH. The dynamic behavior of the polymer with different concentrations was studied by DLS and zeta-potentiometer.

In Figure 1, high degrees of aggregation (turbid state) were observed with long aging time at concentrations below 1 mg/mL PAH into pH 3 buffer solution. The diffusion coefficients of turbid solution from DLS steeply decreased compared to initial transparent state, and the peak width of hydrodynamic radius (RH) distribution was narrow because uniform hydrophobic domains were formed during the aggregation process.

Supplemental experiments were performed with 0.05, 0.1, 0.5, 1.0, 2.0, 4.0, 5.0, 10.0 and 20 mg/mL PAH into pH 3 and pH 10 buffer solutions. But the aggregated PAH (turbid state) was only observed in pH3 solution, but not in pH 10 solution.

The solution dynamics of aggregated PAH (turbid state in pH 3 solution) were investigated by light scattering and microelectrophoresis. PAH has different physical property or structural conformation between the solutions of turbid and transparent state despite the PAH solutions at the same concentration. The diffusion coefficients and zeta-potential of aggregated PAH at the specific concentration which is transformed into turbid state with long aging time were measured at pH 3, 5, 7 and 10 by dilution of turbid solution with individual pH buffer solutions (pH 3, 5, 7 and 10).

If PAH had an extended coil in the turbid solution, the diffusion coefficient of polymer into pH 5, 7 and 10 solutions would be increased due to charge screening, and it has similar tendency into pH 7 and 10 solutions due to neutral charge of functional group.

As seen in Figure 2, the diffusion coefficients with non-linear single exponential fit were determined to be \( 1.86 \times 10^{-8}, 0.40 \times 10^{-8}, 0.163 \times 10^{-8} \) and \( 1.405 \times 10^{-8} \) cm²/s for pH 3, 5, 7 and pH 10, respectively. The RH of aggregated PAH deduced by the Stokes-Einstein (S-E) equation were 138, 621, 1244 and 199 nm for pH 3, 5, 7 and pH 10, respectively. An RH of 138 nm corresponds to a primary aggregate, and 621 and 1244 nm correspond to secondary aggregates.

![Figure 1](image_url)  
Figure 1. The turbidity change of PAH at various concentrations (bottom) and aging time (right) in pH 3 buffer containing hydrophobic anionic salt.

![Figure 2](image_url)  
Figure 2. (a) Translational diffusion coefficients of aggregated PAH (turbid state, 1 mg/mL PAH into pH3 buffer after 14 days) with various pH solutions by DLS. An amount of 0.2 mL of aggregated PAH (turbid state) was mixed with 2 mL of each pH buffer solution into pre-cleaned light scattering cell. (b) Zeta potential for each pH solution. (The line is just for guide.)
formed by the small primary aggregates shown in Figure 2(a). This phenomenon was further elucidated by the microelectrophoresis experiment.

Figure 2(b) shows the magnitude of the zeta potential corresponding to the diffusion coefficient, which indicates the potential stability of the colloidal system. Although each turbid state had different $R_H$ and surface charge values, the aggregated PAH behaved as a particle with a specific domain. If all the particles in suspension have a large negative or positive zeta potential, the particles will tend to repel each other and flocculation does not occur. However, if the particles have low zeta potential values, there is no force to prevent the particles from coming together, resulting in flocculation. The most important factor that changes the zeta potential is pH. The aggregated PAH in pH 3 buffer has a highly positive zeta potential, suggesting primary aggregates were very stable due to repelling forces. If more alkali is added to the aggregated PAH, the positive charge neutralizes and the system becomes very unstable: the values of zeta-potential are 4 and 0 mV for pH 5 and 7, respectively. When the conditions reached a higher pH, the solution had a negative zeta potential, as low as $-14$ mV. As a result, the aggregated PAH at pH 3 and 10 had a stable colloidal system. The aggregated PAH at pH 5 and 7 had an unstable colloidal system with secondary aggregates, which were formed by bridging of the pH buffer. After a long aging time, the neutrally aggregated PAH precipitated and the stabilization was sustained at pH 3 and 10. The diffusion coefficient and zeta potential relationship proved that aggregated PAH (primary aggregate) has a specific domain and was affected by pH and ionic strength. Although PAH behaves like particles with aggregated domains, the aggregate maintains a high surface charge on the polyelectrolytes. A polyelectrolyte complex is primarily formed by the strong coulomb interaction between the oppositely charged polyelectrolyte and hydrophobic salt.

**Time Dependent Behavior: Light Scattering Study.** As observed with DLS measurement, the change of $R_H$ had not been observed into the PAH solution (every concentration which became turbid) without hydrophobic anionic salt, even over aging time of one month. In order to determine how the salt influences the equilibrium, the structural stability of 0.5 mg/mL PAH in pH 3 buffer was studied over time. The sample was analyzed by DLS up to 30 days after the original sample preparation. Figure 3(a) shows the correlation functions of PAH 0.5 mg/mL pH 3 buffer system with various aging times. The inset shows the relaxation time ($\tau$) distributions of the PAH solutions at a scattering angle of 90° in Vv geometry. The 0.05 mg/mL PAH in pH 3 buffer became turbid after five days. At higher concentration than 0.05 mg/mL, aging time longer than five days was required until being turbid. The 0.05, 0.25, 0.5, 1 and 2 mg/mL PAH concentrations become turbid after 5, 7, 14, 18 and 30 days, respectively. But the aging time to make turbid state is not exact time because the time period is so long. There are ± 2 day deviations in the aging time. The $R_H$ distribution from the CONTIN-plot was monodispersed in a concentrated state compared to the transparent state before aggregation. The polyelectrolyte and globular coils below 1 mg/mL have a broad $R_H$ distribution. In the transparent state, a single diffusion mode was observed; however, the distribution of decay times was very broad with broad $R_H$ distribution. The increasing $R_H$ was caused by aggregation of the smaller molecular mass salt. The salt influence also depends on the molecule type and mixing ratios. The degree
of aggregation was controlled by the addition of polymer during complex formation. The response time of the polyelectrolyte complexes to a salt is very different, and strongly dependent on the polyelectrolyte components used.\textsuperscript{15} It can not be fully explained why the aggregation time was long and turbid with respect to the multiplicity of this system, such as buffer ions, pH condition and NaCl salts. The change of conformation and interaction between potassium hydrogen phthalate and PAH will be discussed in terms of spectroscopic observations. In Figure 3(a), the single diffusion mode was corroborated by the sample turbidity as observed by DLS analysis.

The turbidity was not observed for concentrations greater than 2 mg/mL. The PAH 2 mg/mL solution became turbid after one month of aging. The same system was observed for three months and a turbid state was not observed in the 2 mg/mL solution. In Figure 3(b), the translational diffusion coefficient changed regularly with concentrations. The diffusion coefficients were $2.6 \times 10^{-7}, 2.77 \times 10^{-8}, 2.1 \times 10^{-8}$ and $1.5 \times 10^{-8}$ cm$^2$/s for 2, 1, 0.5 and 0.25 mg/mL PAH solutions, respectively. The $R_h$ were 9.5, 89, 117 and 166 nm for 2, 1, 0.5 and 0.25 mg/mL PAH solutions, respectively. Generally, in polyelectrolyte solutions, two modes (fast and slow) may be observed. The physical origin of the slow diffusive process is not yet understood; however, it may be a result of the diffusion of large domains in the solution. The large domains can be formed by a number of molecules “incompactly connected” by electrostatic interactions. The PAH system was a transparent state. As mentioned above, the transparent state has a very broad $R_h$ distribution that contains fast (coil) and slow modes (connected molecule). SLS and DLS experiments can elucidate the $R_h$ ($\rho$ parameter) ratio which indicates whether the polyelectrolyte presents as a random coil (linear, polydispersed chain) or a domain. The literature value of $\rho$ calculated for an ideal random coil system (linear, polydispersed chain) is 2.05. For systems with gyration radii of 149, 179 and 212 nm, and the $R_h$ values of 89, 117 and 166 nm for 1, 0.5 and 0.25 mg/mL aggregated PAH, $\rho$ was 1.67, 1.53 and 1.3, respectively. The experimental values were smaller than the theoretical, indicating the aggregated PAH (turbid state) exists as a structure with a well-defined domain rather than a random coil.\textsuperscript{16-18}

**The Interaction between PAH and Hydrophobic Salt.**

As the aging time increased, PAH formed self assemblies with potassium hydrogen phthalate. FT-IR and Raman spectra of the transparent and turbid state show some clues to solve this problem. The change from the transparent to turbid state was due to the potassium hydrogen phthalate, not NaCl. The dynamic behavior of polyelectrolytes is much more complicated than that of the uncharged polymers due to the multiplicity of interactions present in the polyelectrolyte solution, the nature of which is not fully understood. However, the polyelectrolyte effect was supposed to appear due to conformational changes of the molecules in the system. The turbid state only occurred at low concentrations (below 1 mg/mL). Although potassium hydrogen phthalate is a small ionic molecule, the concentration was greater than that of PAH. The objectives of these experiments were to obtain information about the interactions between PAH and potassium hydrogen phthalate. The PAH solutions with various concentrations were prepared and incubated at room temperature until they became turbid. The objective was to observe the difference between a transparent and turbid solution with 0.5 mg/mL PAH. Figure 4 shows the pH 3 buffer, (a), and aggregated PAH solution, (b, c), spectra compared to that of the standard potassium hydrogen phthalate.

The important PAH absorption bands indicating degree of ionization were found in the strong broad absorption band range of 2800-3250 cm$^{-1}$ associated with the stretching mode of the NH$_3^+$ group. The PAH bands indiscernibly overlap with bands associated with the CH$_2$ asymmetric and symmetric stretching modes.\textsuperscript{19} Changes in C=O carboxylate ion symmetric stretching were observed in the range of 1300-1450 cm$^{-1}$, specifically near 1385 cm$^{-1}$. In Figures 4(b) and (c), the 1385 cm$^{-1}$ peak disappeared, due to ionic interaction of the carboxylate ion of the potassium hydrogen phthalate and amine group of PAH. The ratio of two peaks near 1700 cm$^{-1}$ was different in Figures (b) and (c) due to a red shift.\textsuperscript{20,22} For the turbid sample, the intensity of the right peak increased with respect to the left one. A peak shift was observed near 1600 cm$^{-1}$ in ATR (transparent solution - 1687 cm$^{-1}$; turbid solution - 1665 cm$^{-1}$). The Raman spectrum of the solution was measured to obtain reliable data. For the transparent solution, specific interaction between PAH and potassium hydrogen phthalate was not expected, resulting in only a potassium hydrogen peak similar to FT-IR. It was difficult to measure the FT-Raman due to low intensity of the strongly turbid sample, such that when a small amount of PAH was added, the color changed from turbid to slightly transparent. Figure 5 shows the FT-Raman spectra of PAH with step-wise addition of solution. The most important peak was near 1767 cm$^{-1}$ which appeared only in the turbid state that was unrelated to PAH. A buried peak was expected to appear through peak shifting by interaction of the carboxylate group of the potassium hydrophobic salt and PAH interactions.
gen phthalate and amine group of PAH. The appearance of a new Raman peak at 1767 cm$^{-1}$ indicated the PAH solution could be chemically changed by potassium hydrogen phthalate, due to the hydrogen bonding with the amine group of PAH.

**Segregation of Aggregated PAH.** As previously mentioned, the aggregated PAH acted as a charged particle with a specific domain due to a high concentration of the hydrophobic potassium hydrogen phthalate. The primary aggregate from the pH 3 buffer altered with time and concentration. The change of solution dynamics was elucidated by addition of PAH powder to the aggregated PAH solution.$^{23}$

Figure 6 shows the correlation function of aggregated PAH (1 mg/mL, pH 3) in 6, 12, and 48 hours after adding PAH powder (extended coil state) of various concentrations in which the $R_H$ value decreased by electrostatic repulsion between the aggregated PAH and individual PAH chains.

Upon adding PAH, the $R_H$ was unchanged until 12 mg PAH, at which concentration the diffusion coefficient steeply decreased and the solution became transparent. At other concentrations, the decrease of diffusion coefficient depended on aging time. The forming rate of aggregated PAH was very slow, elucidating the self-assembly of the hydrophobic salt, potassium hydrogen phthalate, and electrostatic repulsion was a very slow process. With 4 and 5 mg/mL, the $R_H$ was unchanged and the zeta potential was as high as 30 mV. The zeta potential value was unchanged over long aging times; however, above 8 mg/mL, the diffusion coefficient increased and divided to a fast and slow mode upon addition of PAH. The zeta potentials of the samples with 4 and 5 mg/mL were 36.1 and 34.85 mV, respectively. These values were similar to initial aggregated PAH solution. Upon increasing the amount of PAH, the zeta potential decreased to 23.6 mV (8 mg/mL) and 23.72 mV (12 mg/mL), which were similar to the transparent state. Therefore, adding PAH to the aggregated PAH solution caused a change from the turbid to transparent state. In the transparent state, the zeta potential was approximately 20 mV without dependence on concentration. After adding 8 mg/mL PAH, the diffusion coefficient was unchanged. After 24 hours, the $R_H$ began to

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**Figure 5.** FT-Raman spectra of (a) PAH 50 mg/mL (transparent state), with (b) 5 mg/mL and (c) 9 mg/mL in 1 mg/mL turbid PAH, (d) PAH 1 mg/mL (transparent state) in pH 3 buffer, and (e) 10 mg/mL PAH in DI-water (from top to bottom).

**Figure 6.** (a) Correlation function of aggregated PAH (turbid state, 1 mg/mL PAH into pH 3 buffer after 14 days) in six hours after adding PAH powder with various concentrations. The inset shows the relaxation time ($\tau$) distributions of aggregated PAH at a scattering angle of 90° in Vv geometry. The terminal shows the relaxation time ($\tau$) distributions of aggregated PAH at a scattering angle of 90° in Vv geometry. (b) Correlation function of aggregated PAH 24 hours after adding PAH polyelectrolyte of various concentrations. The inset shows the relaxation time ($\tau$) distributions of aggregated PAH at a scattering angle of 90° in Vv geometry. (c) Correlation function of aggregated PAH 48 hours after adding PAH polyelectrolyte of various concentrations. The inset shows the relaxation time ($\tau$) distributions of aggregated PAH at a scattering angle of 90° in Vv geometry.
decrease, and after 48 hours, the aggregated PAH was segregated to individual and globular coils. The scattering intensity data in Figure 6 shows the supplemental expectation of the aggregated PAH. For the highly monodispersed or aggregated PAH, the scattered intensity increased and the aggregated PAH had a very high light scattering intensity, which steeply decreased at a specific concentration. To this end, the aggregated PAH was dispersed by the like charged polyelectrolyte. Figure 7 shows the corresponding relaxation time ($\tau$) distributions at a scattering angle of 90° in Vv geometry after adding (a) 5 mg PAH, (b) 9 mg PAH, and (c) 12 mg PAH. The relation time was obtained every 6, 12, 24, and 48 hours. We could clearly observe the shift of the decay times by changing the concentration and aging time.

Table 1 shows the radius distributions of Figure 7 corresponding to the scattering intensity data. Adding 5 and 12 mg/mL PAH powder did not change both radius distribution and scattering intensity. But the change by adding 9 mg/mL PAH powder was proceeded slowly.

**Conclusion**

The dynamic behavior of polymer aggregation with potassium hydrogen phthalate and NaCl was investigated as a function of pH and aging time. PAH showed broad diffusion in dilute solutions. Upon increasing aging time, potassium hydrogen phthalate contributed to aggregation, and the diffusion coefficient steeply decreased with specific aging time. Although the system was complex, the alterable factors were limited to polymer concentration and aging time. The growth of aggregation under kinetic conditions over long aging times was difficult to understand and it was believed that the aggregated size initially formed by a shear field of stirring. When the shear field was removed by potassium hydrogen phthalate, the particles further aggregated, potentially due to hydrogen bonding or ionic interactions between the anionic hydrophobic salt and cationic polyelectrolyte. Evidence for the aggregation is due to the charge interactions. Although the flocculants were very strongly connected after long aging times, the rate of the formation was faster at low polymer concentrations. Notably, the 2 mg/mL PAH solution became turbid after one month. The conformation of aggregated PAH could be obtained from the $R_g/R_H$ ($\rho$ parameter) ratio by static and dynamic light scattering. The ratio values were smaller than ideal random coils indicating the aggregated PAH (turbid state) existed as a well-defined domain rather than random coil.

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