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Aqueous solutions of random poly(methyl methacrylate-co-acrylic acid): effect of the acrylic acid content†

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Aqueous solutions from two types of PMMA-AA polymers (PMMA-25AA and PMMA-50AA) are investigated. The aim is to outline the effect of AA content on the bulk solution and air/solution interface properties. The experiments include dynamic light scattering, surface tension and interfacial rheology measurements. The drainage kinetics and stability of microscopic foam films are also investigated. It is established that at similar conditions (pH = 10, temperature 20 $^{\circ}$ C) the polymer molecules have almost the same mean molecular weight and size distribution of the bulk globules. Dynamic and equilibrium surface tension measurements reveal systematically higher values for PMMA-50AA as compared to PMMA-25AA. These outcomes are related to higher bulk electrophoretic mobility in the PMMA-50AA case and the overall more stretched configuration of the polymer at the air/solution interface. Surface dilational rheology characteristics are particularly sensitive to the polymer structural peculiarities: while no significant changes are registered in the case of PMMA-25AA, the solutions of PMMA-50AA exhibit a pronounced maximum in surface dilational elasticity for the concentration $\sim 1 \times 10^{-4}$ mol L⁻¹. This observation is a clear sign of specific bulk and/or interfacial (structure) transition which has to be investigated in further studies. Microscopic foam films provide additional evidence for the effect of finetuning the AA content of the copolymer. All microscopic films are stable which is due predominantly to the overwhelming electrostatic repulsion effects. The obtained results add new knowledge to the structure-property relationships of the PMMA-AA based aqueous formulations. They give valuable hints for further fine-tuning opportunities of these systems, that have high innovative potential for various applications.

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Introduction

The adsorption of amphiphilic polymers at interfaces and their performance in the bulk of fluid media are of great importance for understanding and tuning the properties and the stability of colloidal systems. Furthermore, study of the structure–property relationships can provide valuable information of fundamental interest and can determine the range of technical applications of these polymers. That is why the interfacial and bulk solution behavior of amphiphilic polymers has attracted the attention of researchers for many years. ¹⁻⁶

The polymer self-assembly in aqueous media is a topic of high interest on its own.^{7–13} Among other major factors (electrostatic and van der Waals interactions, hydrogen bonding, *etc.*), the hydrophobic interactions are of key importance for the onset of energetically favorable inter- and intra-associations of

the amphiphilic polymers in water. Aside from the onset of typical spherical polymer micelles (hydrophobic core/ hydrophilic shell(s)), 11-13 more complex structures based on loops and cross-linking of the chains can occur in the solution bulk. Besides, if a coordinated network of cross-links is built between the macromolecules, the viscosity of the system increases. As the hydrophobic portion of the macromolecule increases, second and third order polymer architectures are possible to form (unimolecular flower-like micelles, collapsed unimolecular micelles, etc.).8 Thus the self-association phenomena in aqueous media are always related to the type and chemical structure of amphiphilic polymers. The macromolecule self-assembly depends also on factors such as pH, ionic strength, temperature, polymer concentration, etc. Despite the conceptual similarity to common surfactant selfassemblies, the polymer micelles are distinguished by their structural rigidity, larger size, lower critical micellization concentration (CMC) and kinetically frozen (static) equilibrium in the solution as opposed to the dynamic exchange of structural units with the medium in the case of low molecular mass amphiphiles. Polymeric macromolecules can leave the associates only when the conditions in the solutions are changed.8-10

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Self-assemblies of amphiphilic polymers have found a broad range of applications. Among the most popular examples are the so-called polymer hydrogels which can be used as drug/protein carriers for controlled or prolonged release and *in situ* curing gels in tissue engineering. They are actually hydrophilic networks which swell upon changing certain conditions, such as solution pH, ionic concentration, temperature, charge distribution, magnetic field, enzyme presence, *etc.*¹⁴⁻¹⁷

Poly(methyl methacrylates) (PMMA) are among the most studied polymers since the beginning of 20th century because of their specific performance. The transparency and thermal stability of the polymer determines important applications as a lighter alternative to glass, transparent coatings, bone cement for medical and dental purposes, *etc.*^{18,19} Recent investigations on Langmuir adsorption layer equilibrium properties and dilational rheology behavior have reported evidence of quasi 2D-fluid to soft-glass transition at air–solution interface (with transition temperature ~298.15 K).^{20,21} The basic outcome of these studies is that the air–water interfacial region acts as a poorer solvent for the polymer. This conclusion has to be accounted for when combined bulk and interfacial studies are performed and when smart fluid phase formulations are developed based on PMMA types of polymers.

The modifications of PMMA not only improve the properties of PMMA systems but also may imply the onset of new characteristics. This has evoked considerable scientific and industrial interest in PMMA-based polymers. Thus, the copolymerization of PMMA with acrylic acid (AA) has been studied intensively and a variety of materials with useful performance features have been obtained. 19,22-26 For instance, random PMMA-AA copolymers of various compositions have been used to prepare hollow crosslinked gel particles with potential medical application for regeneration of soft tissues.26 It has been established that when the molar percentage of AA in PMMA-AA decreases, pH at which the copolymer can be dissolved in water shifts to higher values.²² It has also been established that the AA-component increases the interaction between PMMA-AA and charged montmorillonite nanoparticles. 19 The study of the impact of AA on the aggregation behavior of poly(n-butyl acrylate)-g-poly(acrylic acid) Pn BuA-g-PAA reveals that the lower molar fraction of the side chains corresponds to a higher degree of micellization of the grafted copolymer.7

Recently novel random copolymers – PMMA–AA – have been synthesized by free radical polymerization. ^{26,27} Despite the limitations, such as the broad molecular weight distribution and uncertainty in the positions of the side chains, ¹⁴ this polymerization methodology is widely used because of the relatively easy synthetic procedure. The specific properties of this type of amphiphilic polymer are expected to be dependent on the types and relative contents of hydrophobic and hydrophilic portions, as well as on the sequence of the monomer distribution. ⁸

In the present study, the interfacial and bulk behavior of two specific modifications of the random copolymers PMMA-25AA and PMMA-50AA are investigated in aqueous solutions. The aim is to understand and evaluate the effect of the AA content on the aqueous solution properties of these copolymers.

2. Materials

2.1. Synthesis and characterization of the materials

All chemicals are purchased from Sigma-Aldrich, unless otherwise stated, and used as received. The PMMA-25AA and PMMA-50AA polymers are synthesized via free radical polymerization of the appropriate mixtures of MMA and AA, using 2,2'-azobis(2methylpropionitrile) (AIBN) as an initiator.26,27 The structural formula of the resulting copolymers is given in Fig. 1. The abbreviation codes are derived from the molar percentages of each component. For example, PMMA-50AA contains 50 mol% MMA and 50 mol% AA units. The copolymers are characterized by GPC and ¹H NMR. The GPC analyses have given monomodal molar mass distributions with polydispersity values of 1.69 and 1.92 (Fig. S1, ESI[†]). The ¹H NMR spectra have been used for the determination of the copolymers' structural compositions by examining the relative ratios of signal intensities of the methoxyl protons of MMA groups and the methylene protons of AA groups (Fig. S2, ESI[†]). The experimental values obtained by ¹H NMR are in good agreement with the theoretical values (Table 1).

2.2. Preparation of polymer aqueous solutions

Due to its hydrophobic backbone, the copolymer PMMA-AA is poorly soluble in aqueous media. An experimental procedure has been developed to overcome this problem. The key approach is to perform the experiments at relatively high pH of 10.

The following procedure is applied to both PMMA–25AA and PMMA–50AA in order to eliminate possible differences in the results caused by variations in method of preparation. First, PMMA–AA is mixed with a buffer of pH = 10 ± 0.02 (Na₂B₄O₇/NaOH, provided by Fluka) at constant magnetic stirring of 500 rpm (J. P. Selecta) and temperature 20 °C. The duration of the mixing process depends on the copolymer concentration: 15 minutes for the low concentration range of $C_{\rm p} = 10^{-7}$ to 10^{-6} mol L⁻¹ and 1–1.5 hours for $C_{\rm p} = 10^{-4}$ mol L⁻¹ and higher polymer concentrations. The stirring is performed in a closed vessel to avoid evaporation. Second, the solution is placed into an ultrasonic bath (Bandelin electronic GmbH & Co. KG) and sonificated at a frequency of 35 kHz for 15 minutes. Finally, the solution is filtered through a 0.45 μ m sterile microfilter (Millex) using a sterile syringe.

The concentration range of PMMA-50AA in the experiments varies within $C_{\rm p}=1\times10^{-6}$ to 5×10^{-4} mol L⁻¹, while that of PMMA-25AA is between $C_{\rm p}=1\times10^{-7}$ to 1×10^{-4} mol L⁻¹. The upper concentration limits are determined by the conditions for the onset of turbidity because turbid solutions could not be

Fig. 1 Structure of the poly(methyl methacrylate-co-acrylic acid) (PMMA-AA) copolymers used in this study.

Table 1 Characteristic molecular parameters for PMMA-25AA and PMMA-50AA

			mol% MAA	
Composition	$M_{\rm w}^{a} \left({\rm g \ mol}^{-1} \right)$	PDI^a	Theor.	Exper.b
PMMA-25AA	28 730	1.92	27	25
PMMA-50AA	31 130	1.69	55	50

^a Determined by GPC. ^b Determined by ¹H NMR.

reliably investigated. It is established that at $C_{\rm p}=1\times10^{-3}$ mol L⁻¹ for PMMA-50AA and $C_{\rm p}=5\times10^{-4}$ mol L⁻¹ for PMMA-25AA, the aqueous solutions become turbid. In all experiments the temperature is kept at 20 \pm 0.1 °C and the pH-value is maintained at pH = 10.

3. Methods

3.1. Dynamic light scattering

"Nano ZS" device (Malvern Instruments Ltd.) is used for the measurement of the sizes and the electrophoretic mobility of the copolymer structures in the solution bulk. The light source is 'red' laser (633 nm) and the regime of backward scattering angle (173°) is applied. This angle is considered as suitable for size measurements in the range of 0.6 nm to 6 µm. In addition, backwards scattering has the advantage of decreasing the effect of multiple scattering and larger dust particles (the large particles scatter mainly in forward direction). The device combines the methods of Dynamic Light Scattering (DLS) for the determination of the diffusion coefficient and the hydrodynamic diameter of particles, Electrophoresis28 and Laser Doppler Velocimetry29 to obtain the zeta potential and the electrophoretic mobility. Size distribution data by intensity of the scattered light and by number are obtained. The initial results present the intensity size distribution and the other quantities are recalculated using Mie theory.30 It is worthy to mention the basic assumption set in the DLS technique - the model recalculations are applied for the simplest conjecture about the spherical shape of the particles. As long as the particle size is small enough to correspond to the limitations of the Rayleigh approximation31 this method provides reliable results. A detailed characterization of the DLS technique can be found elsewhere in the literature.31

3.2. Profile analysis tensiometer (PAT)

The values of the dynamic and equilibrium surface tension, as well as the surface dilational elasticity and viscosity are recorded by Profile Analysis Tensiometer – PAT (Sinterface, Berlin, Germany). The bubble regime is chosen as more suitable for the dilute solution measurements. Dynamic surface tension is followed within a period of 24 hours. For the determination of the surface dilational rheology low-frequency oscillations (0.005–0.2 Hz) of the air bubble are performed. The resulting response of the surface tension is recorded and Fourier analysis is applied to calculate the corresponding values of the surface

dilational elasticities and viscosities. The temperature is maintained at 20 \pm 0.1 $^{\circ}C$ by a thermostat.

3.3. Microinterferometric thin liquid film technique

The microinterferometric method of Scheludko–Exerowa allows the investigation of the film drainage and stability, determination of film thicknesses and their time evolution. To our case, the so-called 'common cell' is used and the experiments are carried out at constant capillary pressure. The copolymer solutions are placed in the measuring cell and kept in a thermostatic chamber for 2 hours before the beginning of the measurements. The visualization of the foam films is ensured by a digital camera connected to a high quality inverted microscope (Axiovert 200 MAT, Carl Zeiss). For every copolymer concentration, at least two experimental sets are obtained. Each set consists of about 60–70 foam films and their time evolution is recorded and analyzed.

Results and discussion

4.1. Size and electric properties of PMMA-25AA and PMMA-50AA in the bulk of aqueous solutions

In order to characterize the properties of the copolymers in the bulk of aqueous solutions, we have studied the size distribution and the electrophoretic mobility applying the methods of DLS, Electrophoresis and Laser Doppler Velocimetry. The DLS measurements are performed at the highest concentrations of the copolymers that are used in the surface tension and microscopic foam films experiments. The data are presented in Table 2. The electrophoretic mobilities confirm that PMMA-50AA is about twice as charged as PMMA-25AA and the sign of the charge is negative.

The size distributions by intensity and by number of PMMA-25AA and PMMA-50AA are compared in Fig. 2a and b.

The size of PMMA-50AA in the bulk of aqueous solutions is slightly smaller than that of PMMA-25AA. An additional peak appears in intensity distributions of PMMA-50AA at ~255 nm (Fig. 2a). This course of the intensity curve might be related to possible deflection of the polymer species from the globular conformation due to the higher density of the electric charges in the macromolecule and the resulting tendency toward uncoiling. However, the fraction of the species possessing a hydrodynamic diameter within this range is insignificant and does not sensibly show up in the distribution by number (Fig. 2b). The results are in conformity with the GPC data about the similar mean molecular mass distributions of both polymers as presented in Fig. S1 of the ESI.†

The slightly larger size of PMMA–25AA units in aqueous solutions as compared to PMMA–50AA, might be related to the lower number of AA units in the macromolecules and the slightly longer (hydrophobic) backbone, while the molecular weights of both copolymers (M_w) remains quite similar (see Table 1). It is also probable that a mixture of monomers and micelles and/or premicelles might be present in PMMA–25AA solutions (two peaks in Fig. 2b). For the case of PMMA–50AA solution the monomodal size distribution by number clearly shows that no self-assemblies are detected at these concentration values (Fig. 2b).

Table 2 Size (hydrodynamic diameter) by intensity and by number and electrophoretic mobility of PMMA-25AA and PMMA-50AA in aqueous solutions

Copolymer	Size by intensity, nm	Size by number, nm	Electrophoretic mobility, $\mu m \ cm \ V^{-1} \ s^{-1}$
PMMA-25AA PMMA-50AA	15.6 ± 9.5 $3.122 \pm 1.39 \ (peak 1)$ $255 \pm 144 \ (peak 2)$	$4.19 \pm 3.53 \\ 2.77 \pm 1.71$	$-1.7 \pm 0.68 \\ -3.45 \pm 0.55$

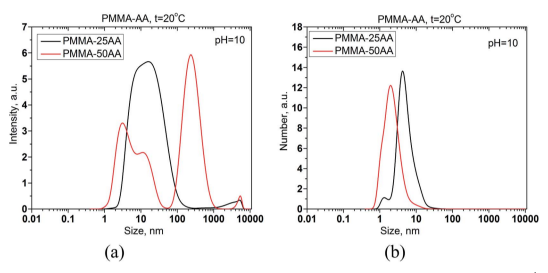


Fig. 2 Mean size distribution by intensity (a) and by number (b) for PMMA-25AA and PMMA-50AA at concentrations of 1×10^{-4} mol L $^{-1}$ and 5×10^{-4} mol L $^{-1}$, correspondingly. Temperature is 20 °C, pH = 10.

An additional factor that can influence the dimensions is the critical micellization concentration (CMC) of polymers, at which micelles presence should be expected in the solution. The parameter has been studied for both PMMA-25AA and PMMA-50AA and the results are presented in 4.2.

4.2. Adsorption layer properties of PMMA-25AA and PMMA-50AA at air/solution interface

The dynamic surface tension data of PMMA-25AA and PMMA-50AA solutions, measured over the course of 24 hours, are represented in Fig. 3a and b, respectively. Both polymers

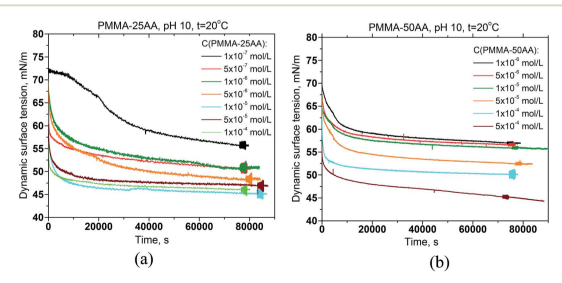


Fig. 3 Dynamic surface tension of aqueous polymer solutions vs. time for different polymer concentrations (a) PMMA–25AA, $C_p = 1 \times 10^{-7}$ to 1 \times 10⁻⁴ mol L⁻¹; (b) PMMA–50AA, $C_p = 1 \times 10^{-6}$ to 5 \times 10⁻⁴ mol L⁻¹. Temperature is 20 °C; pH = 10.

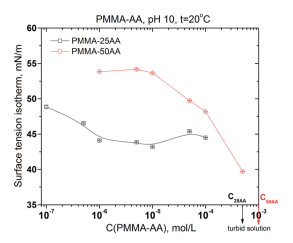


Fig. 4 Equilibrium surface tension of aqueous solutions of PMMA–25AA and PMMA–50AA vs. concentration. Temperature is 20 $^{\circ}$ C; pH = 10

possess surface activity. The dynamic surface-tension values drop down substantially at the initial stages of the measurements (up to 1–1.5 h), followed by a smooth and gradual decrease. This is a typical behavior for amphiphilic polymers and is usually interpreted as diffusion governing the initial stages of the adsorption process, followed by a reorganization of the adsorbed polymer at the air/solution interface. Here a specific run of the curve is recorded for the lowest experimental concentration (1 \times 10 $^{-7}$ mol L $^{-1}$) of PMMA–25AA (Fig. 3a) where a prolonged induction time of $\sim\!5500$ s precedes the major drop in the surface tension values. The result is probably related to the extremely low number density of the macromolecules in the solution.

Additional information about the interfacial properties of the solutions may be extracted by plotting the equilibrium surface tension values against the polymer concentration. Fig. 4 shows a comparison of the surface tension isotherms for both polymers: PMMA–25AA and PMMA–50AA. In the case of PMMA–25AA the bulk CMC-value is at $C_{\rm p} \sim 1 \times 10^{-6}$ mol L $^{-1}$; at higher concentrations a plateau region is registered. This might be interpreted as the adsorption layer at the air/solution interface being densely packed so that the addition of new amounts of polymer results only in bulk self-assembly. The solution becomes turbid at $C_{\rm p} \sim 5 \times 10^{-4}$ mol L $^{-1}$.

The curve for PMMA–50AA is characterized by a distinct plateau at concentrations $C_{\rm p}\sim 1\times 10^{-6}$ to 1×10^{-5} mol L⁻¹. At higher polymer quantities ($C_{\rm p}>1\times 10^{-5}$ mol L⁻¹), the equilibrium surface tension of PMMA–50AA sharply decreases as $C_{\rm p}$ increases. Considering the solution becoming turbid at $C_{\rm p}\sim 1\times 10^{-3}$ mol L⁻¹, it is not possible to determine the CMC-value in this case. So, the results show that generally PMMA–25AA has higher surface activity as compared to PMMA–50AA.

To relate this data to the size measurements (4.1.), it should be pointed out that at the particular concentrations of polymers in the solutions used in DLS experiments, PMMA-25AA has long reached its CMC but PMMA-50AA has not. This might have contributed to the slightly bigger hydrodynamic radius of PMMA-25AA despite the aqueous medium is poorer solvent for it compared to PMMA-50AA.

The interfacial rheology has also been studied in order to obtain more information for the properties of the adsorption layer at the air/solution interface. The surface dilational elasticities of PMMA-25AA and PMMA-50AA are presented as a function of the frequency of oscillation of the bubble surface area (Fig. 5a and 6a) and against the copolymers' concentration (Fig. 5b and 6b). The frequency range used in our experiments is within 0.005-0.2 Hz. In Fig. 5a it is shown that for each C_p the dilational elasticity rises slightly upon the frequency increase, reaching plateau. On the other hand, the elasticity values are only weakly dependent on the polymer concentration (Fig. 5b).

The values of surface dilational elasticities are slightly lower in the case of PMMA-50AA as compared to the PMMA-25AA solutions; this parameter does not vary essentially against the

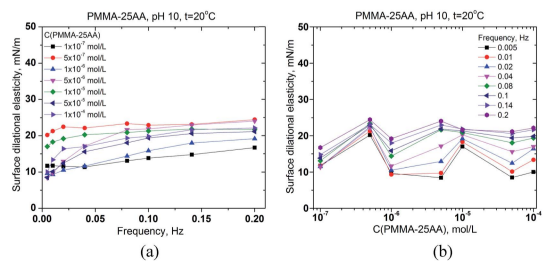


Fig. 5 Surface dilational elasticity of aqueous solutions of PMMA-25AA (a) vs. the frequency of oscillation of the air bubble for different polymer concentrations in the range 1×10^{-7} to 1×10^{-4} mol L⁻¹; (b) vs. the polymer concentration for different oscillation frequencies in the range of 0.005-0.2 Hz. Temperature is 20 °C, pH = 10.

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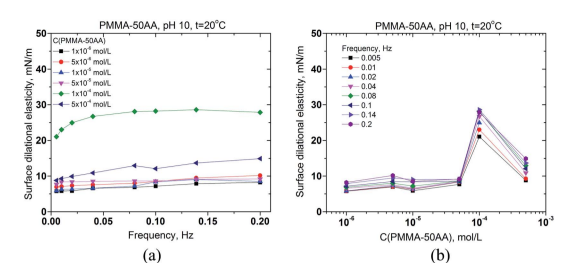


Fig. 6 Surface dilational elasticity of aqueous solutions of PMMA–50AA (a) vs. the frequency of oscillation of the air bubble for different polymer concentrations in the range 1×10^{-6} to 5×10^{-4} mol L⁻¹; (b) vs. the polymer concentration for different oscillation frequencies in the range of 0.005–0.2 Hz. Temperature is 20 °C, pH = 10.

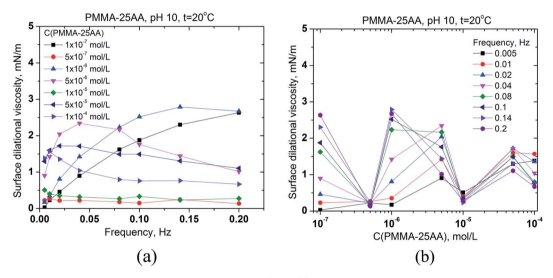


Fig. 7 Surface dilational viscosity of aqueous solutions of PMMA–25AA (a) vs. the frequency of oscillation of the air bubble for different copolymer concentrations in the range 1×10^{-7} to 1×10^{-4} mol L⁻¹; (b) vs. the copolymer concentration for different oscillation frequencies in the range of 0.005–0.2 Hz. Temperature is 20 °C, pH = 10.

frequency. Concerning the concentration dependences of the dilational elasticity, the run of the curves is definitely smoother than in the case of PMMA–25AA. However, there is a remarkable exception for $C_{\rm p}=1\times 10^{-4}~{\rm mol~L^{-1}}$ (Fig. 6a) where the dilational elasticities are sharply enhanced and there are distinct maxima in the course of the concentration curves at $C_{\rm p}=1\times 10^{-4}~{\rm mol~L^{-1}}$ (Fig. 6b). At this polymer quantity the respective values are of a similar order of magnitude as in the case of PMMA–25AA for $C_{\rm p}=1\times 10^{-4}~{\rm mol~L^{-1}}$ (compare Fig. 5b and 6b). The result might support the hypothesis that at this particular polymer concentration complexation (gelation) events occur in the solution bulk.

The surface dilational viscosities of PMMA-25AA and PMMA-50AA are also plotted against the bubble oscillation frequencies and vs. the polymer concentration (Fig. 7 and 8). Obviously the

values are quite low and of the same order of magnitude for both polymers, with being just slightly lower in the case of PMMA-50AA.

The results demonstrate that both polymers impose predominantly an elastic behavior of the adsorbed layers at the air/solution interface.

4.3. Microscopic foam films from aqueous solutions of PMMA-25AA and PMMA-50AA

Characteristic snapshots of foam films from aqueous solutions of PMMA-25AA and PMMA-50AA and their time evolution for different concentrations are presented in Fig. 9 and 10.

The microscopic foam films from both PMMA-25AA and PMMA-50AA, are stable (their lifetimes are longer than 5 minutes) within the investigated concentration ranges.

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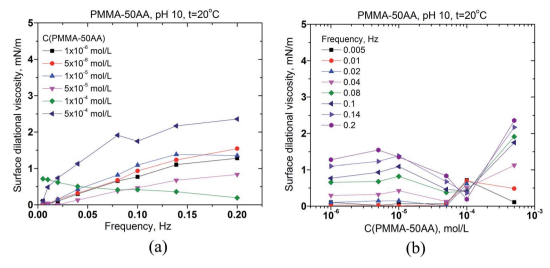


Fig. 8 Surface dilational viscosity of aqueous solutions of PMMA-50AA (a) vs. the frequency of oscillation of the air bubble for different copolymer concentrations in the range 1×10^{-6} to 5×10^{-4} mol L⁻¹; (b) vs. the copolymer concentration for different oscillation frequencies in the range of 0.005-0.2 Hz. Temperature is 20 °C, pH = 10.

Dimple formation is observed in all films from PMMA-25AA except for the lowest concentration investigated ($C_p = 1 \times 1$ 10⁻⁷ mol L⁻¹) (Fig. 9a). The dimples are slowly squeezed out until a significantly smaller white area/spot is left (Fig. 9b-d). At $C_p = 1 \times 10^{-7} \text{ mol L}^{-1}$ of PMMA-25AA, common black films (CBFs) are obtained (Fig. 9a). Besides, little black dots appear at the initial stages of the drainage process. The dots grow fast, develop into spots and within ~ 50 s the whole film area becomes black.

In the case of PMMA-50AA, the film drainage goes smoothly through the onset of channels (Fig. 10) until the so-called "gray" films of homogeneous thickness are obtained. No significant dimples are observed except at the early stages of the film drainage process at $C_p = 5 \times 10^{-5} \text{ mol L}^{-1}$, but in the latter case they are again squeezed out quickly (Fig. 10d).

The drainage kinetics is presented by the changes of film thickness against the time of film thinning. The results are shown in Fig. 11a and b, respectively for both copolymers.

All films drain until reaching equilibrium thickness. The drainage time is essentially lower for PMMA-25AA (Fig. 11a). At $C_{\rm p} > 5 \times 10^{-7} \; {\rm mol} \; {\rm L}^{-1} \; {\rm for} \; {\rm PMMA-25AA} \; {\rm there} \; {\rm is} \; {\rm no} \; {\rm significant}$ change in the film drainage times. In the case of PMMA-50AA solutions, the film drainage curves run more gradually with time for all investigated concentration.

The comparison between the runs of equilibrium film thickness versus the concentration for PMMA-25AA and PMMA-50AA, respectively, is presented in Fig. 12.

The film equilibrium thickness generally increases with time for PMMA-50AA, while it remains in the range of \sim 30-40 nm for the concentrations $C_p > 5 \times 10^{-7} \text{ mol L}^{-1}$ of PMMA-25AA.

5. Discussion

The investigation of aqueous solutions from two types of PMMA-AA copolymers (PMMA-25AA and PMMA-50AA) shows

that the acrylic acid content of the co-polymers has a significant effect on the bulk and air/solution interface properties of the investigated aqueous systems.

First, the surface activity of the polymer macromolecules is related to their amphiphilic nature and depends predominantly on the hydrophobic/hydrophilic (PMMA/AA) molar ratio. Because the mean molecular weight of both polymers is similar (see Fig. S1 in the ESI†), the data from surface tension measurements of the solutions are determined by the molar ratio of the hydrophilic AA against the MMA portions of the respective macromolecules. Therefore, as is to be expected, the results show higher surface activity of PMMA-25AA at the air/ solution interface and the CMC-value is reached at a lower polymer concentration. The surface tension values of PMMA-50AA remain generally higher and a significant shift in CMC (if any) to higher values is expected. The polymer concentration at the onset of turbidity in the solution bulk is also shifted to higher values for PMMA-50AA, as compared to PMMA-25AA solutions (Fig. 4). The surface dilational rheology measurements exhibit no specific characteristics and are typical for amphiphilic substances. The only exception is the surface dilation elasticity of solutions from PMMA-50AA at $C_p = 1 \times 1$ 10^{-4} mol L⁻¹ where there is a well-defined maximum. This concentration performance is evidence for additional structural reorganization in the bulk of the solutions.

Second, the data from the microscopic foam film measurements give ample evidence about the electrostatic stabilization of the films. In our experiments pH is maintained at 10. As the two film interfaces with the adsorbed layers get closer during the film drainage, the negatively charged AA groups provoke the onset of electrostatic repulsive forces and lead to the formation of stable microscopic foam films of equilibrium thicknesses. One interesting observation is that the equilibrium film thickness in the case of PMMA-25AA is systematically lower than that for the same molar concentrations of PMMA-50AA (Fig. 12).

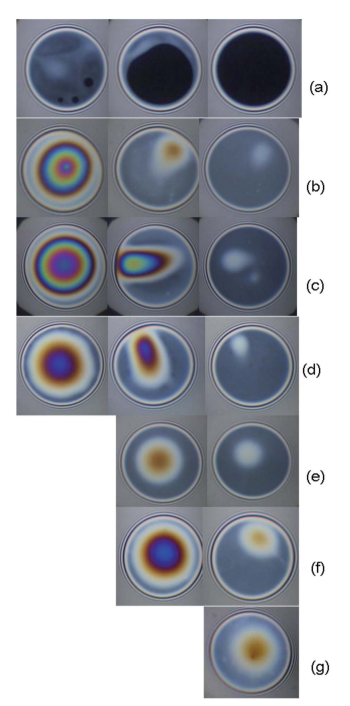


Fig. 9 Characteristic snapshots illustrating the time evolution of microscopic foam films from aqueous solutions of PMMA–25AA for different polymer concentrations. Temperature is 20 °C, pH = 10, the film radius is 100 μm . (a) 1 \times 10 $^{-7}$ mol L $^{-1}$: 38.52 s, 48.4 s, 52.4 s to \sim 5 min; (b) 5 \times 10 $^{-7}$ mol L $^{-1}$: 22.52 s, 59.12 s, 4.57 min; (c) 1 \times 10 $^{-6}$ mol L $^{-1}$: 24 s, 53.36 s, 5.27 min; (d) 5 \times 10 $^{-6}$ mol L $^{-1}$: 5.23 min, 5.43 min, 6.44 min; (e) 1 \times 10 $^{-5}$ mol L $^{-1}$: 5.11 min, 5.25 min; (f) 5 \times 10 $^{-5}$ mol L $^{-1}$: 1.16 min, 4.32 min; (g) 1 \times 10 $^{-4}$ mol L $^{-1}$: 6.06 min.

While there is no significant difference in the mean sizes of the polymer globules, in the cases of PMMA-25AA and PMMA-50AA (Fig. 2b) and the mean molecular weight (Table 1), the relative amount of the charged AA chains is about 50% lower for PMMA-25AA (Table 2). Therefore, the electrostatic repulsive

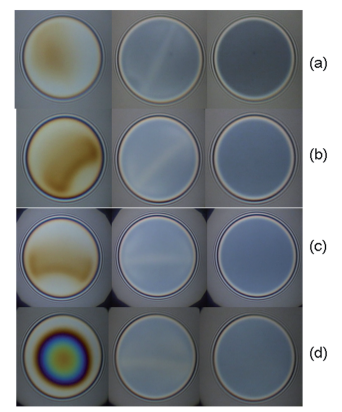


Fig. 10 Characteristic snapshots illustrating the time evolution of microscopic foam films from aqueous solutions of PMMA–50AA for different polymer concentration. Temperature is 20 °C, pH = 10, the film radius is 100 μm . (a) 1×10^{-6} mol L $^{-1}$: 11.44 s, 34.57 s, 5.24 min; (b) 5×10^{-6} mol L $^{-1}$: 14.56 s, 36.16 s, 5.19 min; (c) 1×10^{-5} mol L $^{-1}$: 12.56 s, 37.52 s, 5.08 min; (d) 5×10^{-5} mol L $^{-1}$: 13.46 s, 43.44 s, 5.09 min.

interactions between the adsorbed polymer molecules at the film surfaces is less than in the case of PMMA-50AA. Consequently, the interfaces of the films formed from PMMA-25AA solutions experience weaker repulsion forces and, therefore, this would result in thinner foam films. In contrast, PMMA-50AA should supposedly acquire more stretched configurations in the adsorbed state because of the electric repulsions between the AA groups of the macromolecule. This invokes stronger electrostatic repulsive interactions between the two adsorbed layers of PMMA-50AA due to the denser electrostatic charges at the film interfaces which keep them at larger distance apart (thicker films).

The above explanations presume that the adsorbed polymer layers are densely packed but in a stretched configuration and with similar number density of the adsorbed macromolecules within the investigated concentration range. This might also be deduced from the dynamic surface tension results and the performance of the surface dilational elasticities against the polymer concentration (Fig. 5b and 6b). However, the equilibrium surface tension isotherms advance another factor as well, namely the formation of polymer (pre)micelles in the case of PMMA-25AA (Fig. 4). This may lead to the onset of bulk charged entities which also have an impact on the specific film

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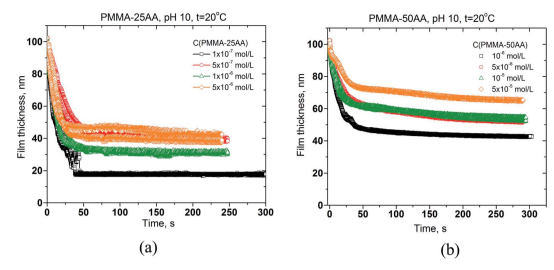


Fig. 11 Time evolution of the mean thickness of foam films from aqueous solutions of (a) PMMA-25AA, and (b) PMMA-50AA, for various polymer concentrations. Temperature is 20 °C; pH = 10; the film radius is 100 μ m.

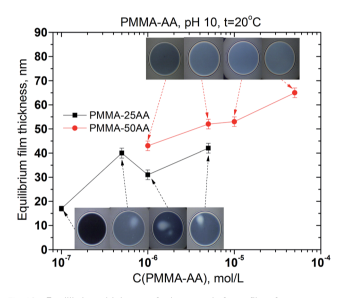


Fig. 12 Equilibrium thickness of microscopic foam films from aqueous solutions of PMMA–25AA and PMMA–50AA *versus* polymer concentration. Temperature is 20 °C, pH = 10, the film radius is 100 μ m. Insets: characteristic snapshots of the films at $\sim\!\!5$ min after their formation.

thicknesses of the equilibrium films. These micelles reduce the number density of the charged entities in the equilibrium films from the PMMA-25AA solutions and are swept into the film meniscus region in the course of the film drainage. So, the electrostatic interactions in the foam films are further reduced and the film-thickness values are lower than in the case of PMMA-50AA. As already stated, CMC cannot be registered for the PMMA-50AA solutions within the investigated concentration range. Due to the more extended configuration of the macromolecules, however, gel-like structures might also form both in the film bulk and/or at the film interfaces. Evidence for such a possibility is the sluggish drainage process of foam film drainage as shown in Fig. 11. Unlike the PMMA-25AA case, the film thinning proceeds slowly, passing through stages of regular

thickness within the draining films and even after \sim 5 min the drainage still moves ahead towards attaining the resulting equilibrium foam films which are without thickness irregularities (no white spots and/or channels).

Concluding remarks

The investigation of aqueous solutions from two types of PMMA-AA copolymers – PMMA-25AA and PMMA-50AA – results in the outline of the following relationships:

- It is established that under similar conditions (pH = 10, temperature 20 °C) the polymer molecules have almost the same mean molecular weight and size distribution of the bulk globules. For the polymer with lower AA content (*i.e.* PMMA–25AA), CMC of the amphiphilic polymer is established to be at $C_{\rm p} \sim 1 \times 10^{-6}$ mol L⁻¹, while no CMC can be found for aqueous solutions of PMMA–50AA.
- Dynamic surface tension measurements reveal systematically higher values for PMMA-50AA as compared to PMMA-25AA. Equilibrium surface tension data are also systematically higher for the case of PMMA-50AA as compared to the polymer with the lower AA content, PMMA-25AA. These outcomes might be related to higher bulk electrophoretic mobility in the case of PMMA-50AA and the overall more stretched configuration of the polymer at the air/solution interface. The electrostatic repulsions prevent the formation of denser adsorption layers in the case of PMMA-25AA.
- Surface dilational rheology characteristics are particularly sensitive to the polymer structural peculiarities and result in the most striking difference between the two polymers: while no significant changes are registered in the case of PMMA–25AA, the solutions of PMMA–50AA exhibit a pronounced maximum in surface dilational elasticity for the concentration of $C_{\rm p}\sim 1\times 10^{-4}$ mol L⁻¹. This fact is a clear sign of specific bulk and/or interfacial (structure) transition which has to be investigated in further studies.

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• Microscopic foam films provide additional evidence for the effect of regulating the AA content of the copolymer. All microscopic films are stable which is due predominantly to the overwhelming electrostatic repulsion effects. However, the structure of PMMA-25AA seems more suitable in view of foam film stabilization: the equilibrium foam films are thinner and a lower quantity of polymer is needed in order to attain film stability.

The obtained results add new knowledge to the structure– property relationships of the PMM-AA based aqueous formulations. They give valuable hints for further fine-tuning opportunities of these systems, that have high innovative potential for various applications.

Abbreviations

PMMA Poly(methyl methacrylate)

AA Acrylic acid

DLS Dynamic light scattering
CMC Critical micellar concentration
GPC Gel permeation chromatography

1H NMR Proton nuclear magnetic resonance

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