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Anthony Phimpachanh, Joseph Chamieh, Laurent Leclercq, Simon Harrisson, Mathias Destarac, et al.. Characterization of Diblock Copolymers by Capillary Electrophoresis: From Electrophoretic Mobility Distribution to Distribution of Composition. *Macromolecules*, 2020, 53 (1), pp.334-345. 10.1021/acs.macromol.9b01978 . hal-02462075v2

HAL Id: hal-02462075

<https://hal.umontpellier.fr/hal-02462075v2>

Submitted on 9 Dec 2020

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Characterization of diblock copolymers by capillary electrophoresis: From electrophoretic mobility distribution to distribution of composition

Anthony Phimpachanh^{1,3}, Joseph Chamieh², Laurent Leclercq², Simon Harrisson⁴, Mathias Destarac⁴,
Patrick Lacroix-Desmazes³, Corine Gérardin³, Martin In^{1*}, Hervé Cottet^{2*}

¹L2C, Univ Montpellier, CNRS, Montpellier, France

²IBMM, Univ Montpellier, CNRS, ENSCM, Montpellier, France

³ICGM, Univ Montpellier, CNRS, ENSCM, Montpellier, France

⁴IMRCP, University of Toulouse, CNRS UMR5623, Toulouse, France

* Corresponding authors: herve.cottet@umontpellier.fr & martin.in@umontpellier.fr

Abstract

Free solution capillary-electrophoresis (CE) is a powerful separation technique for the characterization of diblock copolymers. In this work, four series of double-hydrophilic anionic and cationic block copolymers, namely, poly(acrylamide)-*block*-poly(acrylic acid) (PAM-*b*-PAA), poly(acrylamide)-*block*-poly((3-acrylamidopropyl)trimethylammonium chloride) (PAM-*b*-PAPTAC), poly(ethylene oxide)-*block*-poly(acrylic acid) (PEO-*b*-PAA) and poly(poly(ethylene glycol) methyl ether acrylate)-*block*-poly(acrylic acid) (P(PEGA)-*b*-PAA), were synthesized by reversible addition-fragmentation chain transfer (RAFT) polymerization and characterized by CE. The electrophoretic mobility distributions of the copolymers were transformed into distributions of composition ratio by introducing a retardation parameter, X_{exp} , that represents the hydrodynamic drag retardation due to the neutral block of the copolymer. A linear correlation between X_{exp} and the ratio of the degrees of polymerization of each blocks was experimentally established and was consistent with the model of electrophoretic mobility of composite macromolecules with hydrodynamic coupling. Finally, the comparison of the distributions between the different copolymer families was significantly improved by considering the distributions in composition ratio compared to the electrophoretic mobility distributions, since it takes into account the differences in solvation, expansion and drag force according to the chemical nature of the blocks.

1 Introduction

The characterization of diblock copolymers by separation techniques is challenging but of primary importance to check their purity and to characterize their distributions in size and in chemical composition.¹ Block copolymers prepared by reversible deactivation radical polymerization frequently contain homopolymer impurities.^{2, 3} These include dead chains from termination reactions during polymerization of the first block, as well as the products of side-reactions such as chain transfer to solvent or monomer during polymerization of the second block. The growth of the second block in copolymer synthesis is often evidenced by a decrease of the elution time in size-exclusion chromatography (SEC) as a result of the increased hydrodynamic radius^{4, 5}. However, this only holds if the second block contributes significantly to the hydrodynamic radius of the diblock copolymer. Getting more quantitative information can be challenging especially in aqueous SEC. Different solvation properties between the blocks of a block copolymer can lead to coelution of polymers of different mass in SEC, resulting in inaccuracy in the obtained molar masses.⁶ Additionally, interactions with the stationary phase³ may lead to HPLC-type elution which is dependent on the chemical composition of the polymer as well as its size. For diblock copolymer SEC, particular elution conditions are generally required and size distributions should be expressed in terms of hydrodynamic radius (and not molar mass) due to the difference in chemical composition / solvation of the two blocks⁶. The proportion of each monomer in a copolymer can be obtained by liquid chromatography under critical conditions LCCC, also known as LC-PEAT, for the point of exclusion-adsorption transition for neutral blocks^{7, 8}. The critical conditions for LCCC (or LC-PEAT) are usually difficult to find and are very sensitive to small changes in mobile phase composition and/or temperature.

An alternative separation technique for charged copolymers is free solution capillary electrophoresis (CE)^{2, 3, 7-9}. The electrophoretic separation of charged homopolymers from diblock copolymers is generally easily obtained in free solution CE. Moreover, for self-assembling diblock copolymers, CE can also separate micelles from unimers^{2, 7, 8} and allows studying the impact of added surfactant on the copolymer micelles^{2, 7}. In the presence of cationic blocks, experimental difficulties

arise from polymer adsorption onto the wall of silica based capillaries. The characterization of cationic diblock copolymers requires the use of a neutrally coated³, or positively charged capillary¹⁰. Another challenging issue in the characterization of diblock copolymers by CE is to extract the distribution in composition of the copolymers from the electropherogram. Raw electropherograms can be transformed into distributions of effective mobility, or of any other related parameter, provided that the relationship between effective mobility and the considered parameter is known¹¹. The determination of polymer dispersity via the variance of the chemical composition distribution¹¹ or via the calculation of the ratio of moments of the distribution has also been studied¹. A key point to achieve such electropherogram transformation is to have a reliable relation between the effective mobility of the diblock copolymer and the degrees of polymerization of each block and thus, to the chemical composition of the copolymer³. The electrophoretic mobility of a diblock copolymer is generally expressed as a weighted average of the mobilities of different subunits constituting the copolymer¹². The choice of the subunits and the corresponding weights have been described in the literature as depending on the conformations of each block and on the hydrodynamic coupling regime between the two blocks¹²⁻¹⁵. More recently, Chubynsky and Slater studied in more detail the “end-effect” (i.e. the fact that the ends of the copolymer chain are more hydrodynamically exposed to the solvent)¹⁶ and the effect of polymer stiffness on the electrophoretic modeling¹⁷. The electrophoretic models of composite objects^{12, 13} which are relevant for diblock copolymers, were also applied to end-labeled free solution electrophoresis (ELFSE)^{14, 15}, which consists in attaching a monodisperse neutral block (drag-tag) to a polydisperse biopolyelectrolyte (for instance, for DNA sequencing in free solution^{15, 18}), or conversely, in attaching a monodisperse polyelectrolyte to a polydisperse neutral polymer (for instance, for size-based neutral polymer characterization¹⁴). In this way, the dependence of electrophoretic mobility with the molar mass of the end-labelled composite object is obtained in free solution due to the variation of the charge-to-friction ratio.

Double-hydrophilic block copolymers (DHBC) are block copolymers containing two hydrophilic segments. DHBCs on their own are completely soluble in water and do not self-assemble in dilute conditions. DHBCs can still retain an amphiphilic character and this can lead to self-organization at the meso-scale in concentrated conditions¹⁹. They can undergo morphological transitions induced by

external stimuli²⁰ in dilute solution. The great development of reversible deactivation radical polymerization²¹⁻²⁴ in the last two decades allows tailoring the stimuli-responsiveness (e.g. to changes in pH, temperature, ionic strength, or light) of these polymers by controlling both the nature of monomers and the degree of polymerization of the blocks. When one block is a polyelectrolyte, DHBCs can undergo micellization by electrostatic complexation in the presence of an oppositely charged polyelectrolyte. These properties lead to a wide range of applications such as control of crystallization of inorganic compounds²⁵, drug delivery²⁶ or template for ordered mesoporous materials²⁷. For this last application of DHBC, the asymmetry ratio, defined as the ratio of degrees of polymerization of both blocks, is of crucial interest since it determines the structure of the DHBC-templated mesoporous materials.

It is the aim of the present work to characterize the composition of DHBC by CE, with particular attention to the asymmetry ratio. The approach has been applied to series of anionic and cationic DHBCs, namely, poly(acrylamide)-*block*-poly(acrylic acid) (PAM-*b*-PAA), poly(acrylamide)-*block*-poly((3-acrylamidopropyl)trimethylammonium chloride) (PAM-*b*-PAPTAC), poly(ethylene oxide)-*block*-poly(acrylic acid) (PEO-*b*-PAA) and poly(poly(ethylene glycol) methyl ether acrylate)-*block*-poly(acrylic acid) (P(PEGA)-*b*-PAA). These DHBC have been synthesized in aqueous medium by reversible addition-fragmentation transfer (RAFT) polymerization.

In the next section, different models for the electrophoretic mobility of composite objects such as block copolymers are briefly reviewed. In the third section, the synthesis of the copolymers and the experimental conditions of their characterization by CE are reported. The results of this work are presented in section 4, where we describe a method to transform the distribution of electrophoretic mobility into a distribution of the ratio of degree of polymerization of both blocks.

2 Mobility of diblock copolymers constituted of a polyelectrolyte and a neutral block

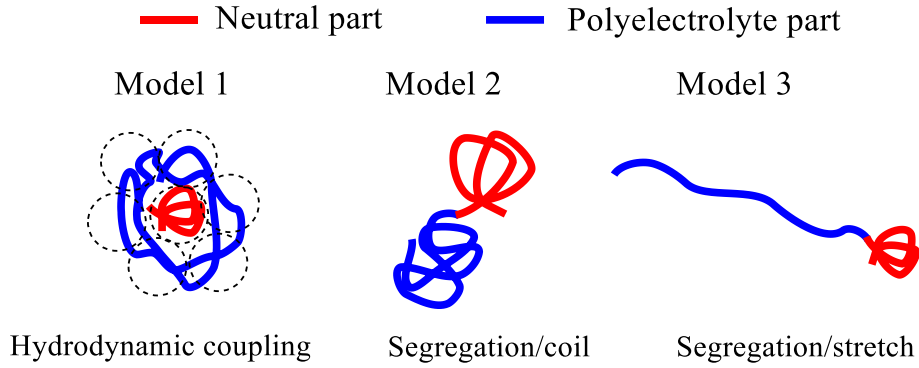


Figure 1. Representation of possible conformations of a double hydrophilic block copolymer composed of a neutral Gaussian coil (in red) linked to a polyelectrolyte block (in blue). In model 1, the polyelectrolyte chain is a coil in hydrodynamic interaction with the neutral Gaussian coil. In model 2, the polyelectrolyte and the neutral polymer separate into two coils. In model 3, the polyelectrolyte is fully stretched and has no hydrodynamic coupling with the neutral coil. Adapted from²⁸

In this section, we present different possible electrophoretic mobility models that are relevant for diblock copolymers composed of a polyelectrolyte part attached to a neutral polymer coil. These theoretical models were developed by Desruisseaux et al.²⁸, building on previous work by Long et al.^{12, 13} Figure 1 shows the different possible conformations that can be encountered for DHBC, corresponding to the different models presented below.

2.1 Model 1: hydrodynamic coupling

In Model 1, hydrodynamic coupling between the polyelectrolyte part and the neutral coil is taken into account. The polyelectrolyte block of the DHBC is composed of N_{blob} equivalent blobs of a size equivalent to the hydrodynamic radius of the neutral coil $R_h^{neutral}$. If $R_h^{neutral}$ is larger or equal to the Debye length, Long et al.²⁹ demonstrated that the electrophoretic mobility of the DHBC composite object composed of $N_{blob} + 1$ subunits of equal size, is given by the number-average of the electrophoretic mobilities calculated on all the equivalent blobs constituting the object. The electrophoretic mobility of the DHBC, $\mu_{ep,1}^{diblock}$, is thus given by²⁸:

$$\mu_{ep,1}^{diblock} = \frac{\sum_{i=0}^{N_{blob}} \mu_{ep}^i}{N_{blob} + 1} = \frac{N_{blob} \mu_{ep}^0 + \mu_{ep}^{neutral}}{N_{blob} + 1} = \frac{\mu_{ep}^0}{1 + \frac{\alpha}{DP_0}} \quad (1)$$

where μ_{ep}^i is the effective mobility of the i^{th} entity (or blob) constituting the DHBC, DP_0 is the degree of polymerization of the polyelectrolyte block, α is the number of charged monomers per blob, μ_{ep}^0 is the effective electrophoretic mobility of the polyelectrolyte part (alone) and $\mu_{ep}^{neutral}$ is the electrophoretic mobility of the neutral part ($\mu_{ep}^{neutral} = 0$). The number of equivalent blobs in the polyelectrolyte chain is given by $N_{blob} = \frac{DP_0}{\alpha}$. Note that N_{blob} (and $\mu_{ep,1}^{diblock}$) depends on the polyelectrolyte persistence length, and thus, on the ionic strength. Equation (1) neglects the so-called end-effect¹⁶. Please note that subscript and superscript 0 refer to the polyelectrolyte block, for consistency with ref 28.

2.2 Models without hydrodynamic coupling

2.2.1 Model 2: polyelectrolyte chain in coil conformation

Model 2 in Figure 1 corresponds to the segregation of the neutral polymer coil from the polyelectrolyte coil. In the absence of hydrodynamic coupling between the two parts, and if the polyelectrolyte chain does not stretch during electrophoresis (i.e. at sufficiently low electric field), the electrophoretic mobility of the DHBC, $\mu_{ep,2}^{diblock}$, is given by the average electrophoretic mobility of the two parts weighted by their hydrodynamic friction coefficient¹². Using Stokes equation for spherical objects, $\mu_{ep,2}^{diblock}$ is expressed as^{13, 28}:

$$\mu_{ep,2}^{diblock} = \frac{\sum_{i=0}^1 \gamma_i \mu_{ep}^i}{\sum_{i=0}^1 \gamma_i} = \frac{\mu_{ep}^0}{1 + \frac{R_h^{neutral}}{R_h^0}} \quad (2)$$

where γ_i is the friction coefficient of the i^{th} part constituting the DHBC, $R_h^{neutral}$ is the hydrodynamic radius of the neutral coil, R_h^0 is the hydrodynamic radius of the polyelectrolyte block.

2.2.2 Model 3: polyelectrolyte chain in fully stretched conformation

When the polyelectrolyte coil is stretched under the concomitant influence of the electric field and the opposed drag force due to the presence of the neutral coil, it can reach a fully extended conformation as depicted in Figure 1 (Model 3). Stokes law is no longer appropriate for the frictional coefficient of the polyelectrolyte chain, and the electrophoretic mobility of the DHBC, $\mu_{ep,3}^{diblock}$, is given by²⁷ :

$$\mu_{ep,3}^{diblock} = \frac{\sum_{i=0}^1 \gamma_i \mu_{ep}^i}{\sum_{i=0}^1 \gamma_i} = \frac{\mu_{ep}^0}{1 + \frac{2R_h^{neutral}}{b_0 DP_0} \ln DP_0} \quad (3)$$

where b_0 is the size of a charged monomer in the polyelectrolyte chain. Equation (3) assumes that the friction coefficient γ_{rod} of the stretched polyelectrolyte cylinder is averaged on all orientations relative to the flow direction, and is given by:

$$\gamma_{rod} = \frac{3\pi\eta b_0 DP_0}{\ln(DP_0)} \quad (4)$$

where η is the viscosity of the solvent.

Even if the electrical field is not strong enough to stretch the polyelectrolyte block (hydrodynamic segregation), the polyelectrolyte contour length may still be shorter than the persistence length of the polyelectrolyte. In this situation, the segregation between the neutral and the polyelectrolyte parts is sterically obtained, but the electrophoretic mobility is still described by equation (3)²⁷.

3 Experimental

3.1 Chemicals

Ammonium persulfate (APS, 98%) and sodium formaldehyde sulfoxylate dehydrate (NaFS, 98%) were purchased from Acros organics. Poly(ethylene glycol) methyl ether acrylate ($M_n = 480$ g/mol, 8.5 EO units on average) and (3-acrylamidopropyl)trimethylammonium chloride (APTAC) aqueous solution (75

wt.%) were purchased from Sigma Aldrich (Saint-Quentin-Fallavier, France) and used as received. Acrylic acid from the same suppliers was distilled under vacuum at room temperature. PEO $M_n=5000$ g.mol⁻¹, $\bar{D} = 1.04$ was purchased from. 4,4'-azobiscyanopentanoic acid (ACPA, Aldrich, 98%) and 2,2-Azobis(isobutyramidine) dihydrochloride (AIBA, Sigma-Aldrich, 97%) were used as received.

For EC experiments, background electrolytes were prepared in ultra-pure water purified on a Millipore system (Molsheim, France) from tris(hydroxymethyl)aminomethane (TRIS, 99,9%, Merck), 4-Morpholinoethanesulphonic acid (MES, >99%, Acros Organics), and 2-[Bis(2-hydroxyethyl)amino]-2-(hydroxymethyl)propane-1,3-diol (BIS-TRIS, >99%, Acros Organics). Anisic acid (99,5%), ammediol (99,5%) used as markers for detection were purchased from Sigma-Aldrich.

3.2 Double-hydrophilic block copolymer synthesis

This section describes the synthesis of the copolymers. The synthetic pathway, the size exclusion chromatograms and the ¹H NMR spectra are given in SI as well as the temporal electropherograms.

3.2.1 Synthesis of poly(acrylamide)-*b*-poly(acrylic acid)

Aqueous RAFT/MADIX polymerization of AA

Polyacrylamide macro RAFT/MADIX agent (PAM-Xa, $M_n = 5000$ g.mol⁻¹) was synthesized according the procedure described by Layrac et al..³⁰ Synthesis of PAM₇₀-*b*-PAA₂₀ was performed as follows³¹: PAM-based chain transfer agent (PAM-Xa,) (15.76 g, 3.029 mmol), acrylic acid (4.24 g; 58.77 mmol), AIBA (0.0821 g, 0,3 mmol) and water (42 g) (solids = 30.6%) were introduced in a round bottom flask. The mixture was degassed with argon at room temperature for 30 min and then placed in a thermostated oil bath at 65°C under argon for 2 hours. Conversion was quantitative, acrylic acid traces were eliminated by dialysis (MWCO 1000 Da) and pH-metric monitoring. The polymer solution was then freeze dried and a white powder was obtained. Four DHBC were synthesized according to this procedure: PAM₇₀-*b*-PAA₂₀, PAM₁₄₀-*b*-PAA₄₀, PAM₁₄₀-*b*-PAA₆₀ and PAM₁₄₀-*b*-PAA₈₀ (see Table 1).

3.2.2 Synthesis of poly(acrylamide)-*b*-poly((3-acrylamidopropyl)trimethylammonium chloride)

Aqueous redox RAFT/MADIX polymerization of APTAC

Synthesis of PAM₇₀-*b*-PAPTAC₃₀ was performed as follows: two aqueous solutions of NaFS (5%w) (1.23 g of solution, 0.4 mmol) and NaPS (5%w) (2.37 mg of solution, 0.5 mmol) were prepared. PAM-Xa macroxanthate (9.97 g, 1.99 mmol), APTAC monomer (13.3 g of solution, 0.057 mmol), and water (50 g) (solids = 25.4%) were introduced in a round bottom flask. The pH of the mixture was first adjusted at 2 with hydrochloric solution 1M. Then the mixture was degassed with argon at room temperature for 30 min and placed in a thermostated oil bath at 25°C under argon. Both solution of NaFS and NaPS were introduced in the round bottom flask and the reaction mixture was stirred for 3 hours. Monomer traces were eliminated with dialysis (MWCO 1000 Da) and conductivity monitoring. The polymer solution was then lyophilized and a white powder was obtained. This redox process at 25°C was developed after the paper of Sutton et al.⁹ to minimize the formation of dead chains. Four DHBC were synthesized according to this procedure: PAM₇₀-*b*-PAPTAC₃₀, PAM₇₀-*b*-PAPTAC₆₀, PAM₁₄₀-*b*-PAPTAC₆₀ and PAM₁₄₀-*b*-PAPTAC₁₂₀ (see Table 1).

3.2.3 Synthesis of poly(ethylene oxide)-*b*-poly(acrylic acid)

Aqueous RAFT polymerization of AA

Synthesis of PEO₁₀₅-*b*-PAA₂₀ was performed as follows: PEO₁₀₅-CTA (poly(ethylene oxide)-chain transfer agent) macro RAFT agent was obtained following the procedure published by Bathfield et al.³². PEO₁₀₅-CTA, (10.53 g, 2.19 mmol), ACPA (0.123 mg, 0,439 mmol), acrylic acid (5.5 g, 76 mmol) and deionized water (29.5 mL) (solids=35.4%) were introduced in a Schlenk tube equipped with a magnetic stirrer. The mixture was degassed by five freeze-evacuate-thaw cycles and then heated for 42 hours at 75°C under nitrogen in a thermostated oil bath. Final conversion = 72%. Monomer conversion was determined by ¹H NMR spectroscopy, using a Bruker 400MHZ spectrometer. Samples for analysis by NMR were prepared by adding 0.6 mL of D₂O to 0.1 mL of polymerization medium. Once the reaction was complete, the solvent was evaporated, and then the DHBC was dissolved in a minimum amount of dichloromethane before being precipitated twice in a large volume of cold diethyl ether. It was then recovered by filtration, and finally dried under vacuum overnight before analysis by SEC and ¹H-NMR. SEC was performed in DMF-LiBr after methylation³³ with trimethylsilyldiazomethane. Four DHBC

were synthesized according to this procedure: PEO₁₀₅-*b*-PAA₂₀, PEO₁₀₅-*b*-PAA₃₀, PEO₂₁₀-*b*-PAA₄₀ and PEO₂₁₀-*b*-PAA₅₀ (see Table 1). The degrees of polymerization of the commercial starting PEO have been determined by ¹H NMR³² and are presented in Table 1.

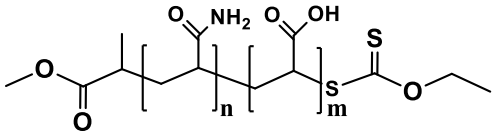
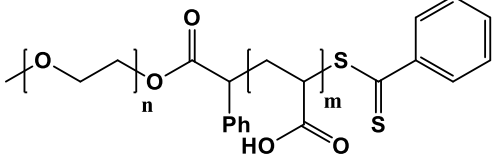
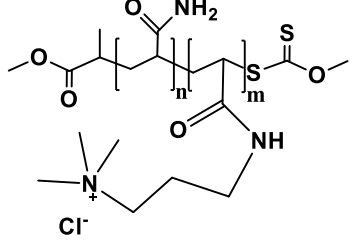
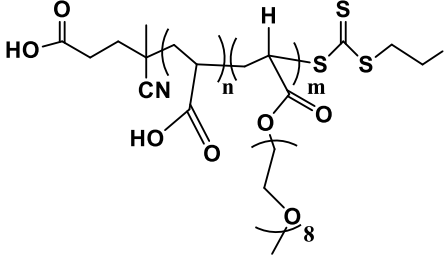
3.2.4 Synthesis of poly(acrylic acid)-*b*-poly(poly(ethylene glycol) methyl ether acrylate)

Aqueous RAFT polymerization of AA and PEGA

5,7-dithia-6-thio-4-methyl-4-cyanodecanoic acid (CTPPA) was obtained by reaction of ACPA with bis(propylsulfanylthiocarbonyl) disulfide according to literature.³⁴ Synthesis of PAA₂₁-*b*-P(PEGA)₁₂ was performed as follows: PAA-CTPPA chain transfer agent was synthesized according to the process described in literature³⁵: in a round bottom flask, CTPPA (0.8 g, 2.57 mmol, purity=89%), acrylic acid (3.9g, 54.1 mmol), ACPA (0,072 g, 0.26 mmol) and half of the amount of water (7.5 g) are stirred until dissolution of CTPPA. The remaining water (7.5 g) was introduced and the mixture was degassed with argon for 40 min. The mixture was then heated in an oil bath at 70°C for 5.5 h. For the synthesis of PAA-*b*-P(PEGA), ACPA (0.0715 g; 0.26 mmol) and poly(ethylene glycol) methyl ether acrylate (12.8 g, 26.4 mmol) were added to the PAA-CTPPA reaction medium and the mixture was degassed with argon for 40 min. The mixture was then heated in an oil bath at 70°C for 5 h. Conversion was followed by ¹H NMR. At the end of the polymerization, water was evaporated under reduced pressure and the polymer washed with diethyl ether. NMR sample preparation: 0.6 mL of D₂O was added to 0.1 mL of polymerization medium and quenched in liquid nitrogen. Two DHBC were synthesized according to this procedure: P(PEGA)₁₂-*b*-PAA₂₁ and P(PEGA)₂₂-*b*-PAA₄₅ (see Table 1).

Table 1: Presentation of the chemical structure and the different DHBC samples synthesized and studied in this work. The subscripts in the names correspond to the degree of polymerization of each block. $MW_{neutral}$ and MW_0 are the molar masses of the neutral and of the polyelectrolyte block respectively.

Type of DHBC	$MW_{neutral}-MW_0$ as determined by NMR	Chemical structure
PAM ₇₀ - <i>b</i> -PAA ₂₀	5k-1.4k	
PAM ₁₄₀ - <i>b</i> -PAA ₄₀	10k-2.8k	

PAM ₁₄₀ - <i>b</i> -PAA ₆₀	10k-4.4k	
PAM ₁₄₀ - <i>b</i> -PAA ₈₀	10k-5.6k	
PEO ₁₀₅ - <i>b</i> -PAA ₂₀	5k-1.4k	
PEO ₁₀₅ - <i>b</i> -PAA ₃₀	5k-2.2k	
PEO ₂₁₀ - <i>b</i> -PAA ₄₀	10k-2.8k	
PEO ₂₁₀ - <i>b</i> -PAA ₅₀	10k-3.6k	
PAM ₇₀ - <i>b</i> -PAPTAC ₃₀	5k-5k	
PAM ₇₀ - <i>b</i> -PAPTAC ₆₀	5k-10k	
PAM ₁₄₀ - <i>b</i> -PAPTAC ₆₀	10k-10k	
PAM ₁₄₀ - <i>b</i> -PAPTAC ₁₂₀	10k-20k	
P(PEGA) ₁₂ - <i>b</i> -PAA ₂₁	5.7k-1.5k	
P(PEGA) ₂₂ - <i>b</i> -PAA ₄₅	10.6k-3.2k	

3.3 Capillary electrophoresis

Instrumentation and method

Capillary electrophoresis experiments were performed on an Agilent 7100 capillary electrophoresis instrument with a diode array UV detector. Fused silica capillaries of 50/375 μm I.D./O.D. with polyimide outer coating (cat. no. TSP050375) were from Polymicro Technologies (Phoenix, AZ, USA). Capillary dimensions were 38.5 cm long (30 cm to detection window). New capillaries were conditioned by performing the following washes at 1 bar: 1M NaOH for 30 min and water for 15 min. The temperature of the capillary cartridge was set at 25 °C.

In the case of PAM-*b*-PAA, an electrolyte consisting of 20 mM MES and 14 mM ammonium dihydrogen phosphate pH 6.5 was used. 0.1 g/L anisic acid was added in the sample as a mobility marker. The same background electrolyte was used for PAM-*b*-PAPTAC, but with a different mobility marker (imidazole 0.1 g/L). In the case of PEO-*b*-PAA and P(PEGA)-*b*-PAA, an electrolyte constituted of 6 mM anisic acid and 12 mM BIS-

244 TRIS, pH 6.5 was used as buffer, with MES at 0.5 g/L as mobility marker in the case of P(PEGA)-*b*-
245 PAA.

246 All copolymers were dissolved in water at a concentration of 5 g/L. Samples were injected
247 hydrodynamically on the inlet side of the capillary by applying 30 mbar for 5 s. Separations were carried
248 out by applying a +20 kV voltage. For PAM-*b*-PAA, PEO-*b*-PAA and P(PEGA)-*b*-PAA and PAM-*b*-
249 PAPTAC, detection was realized at 192 +/- 2 nm (reference off).

250 For PAM-*b*-PAA, PEO-*b*-PAA and P(PEGA)-*b*-PAA, the capillary was rinsed between each run by
251 flushing the capillary for 2 min with the background electrolyte, 2 min with 0.1 M NaOH, 2 min with
252 ultra-pure water and 2 min with background electrolyte. For the analysis of the cationic polymer PAM-
253 *b*-PAPTAC, and in order to reduce the adsorption on the capillary wall, surface of the capillary was
254 modified using UltraTrol™ LN (Target Discovery, Inc., Palo Alto, CA), which is a commercial neutral
255 semi-permanent coating based on polyacrylamide derivatives. The coating procedure was performed
256 using the following successive flushes: methanol for 2 min at 1 bar, water for 2 min at 3 bar, 1 M NaOH
257 for 2 min at 3 bar, 0.1 M NaOH for 2 min at 1 bar, 1 M HCl for 5 min at 1 bar, water for 5 min at 1 bar,
258 UltraTrol™ LN solution for 5 min at 1 bar, wait for 5 min, water for 2 min at 1 bar. Prior to each analysis
259 of PAM-*b*-PAPTAC, the capillary was rinsed with the background electrolyte for 2 min at 1 bar.

260

261 **Electropherogram data treatment**

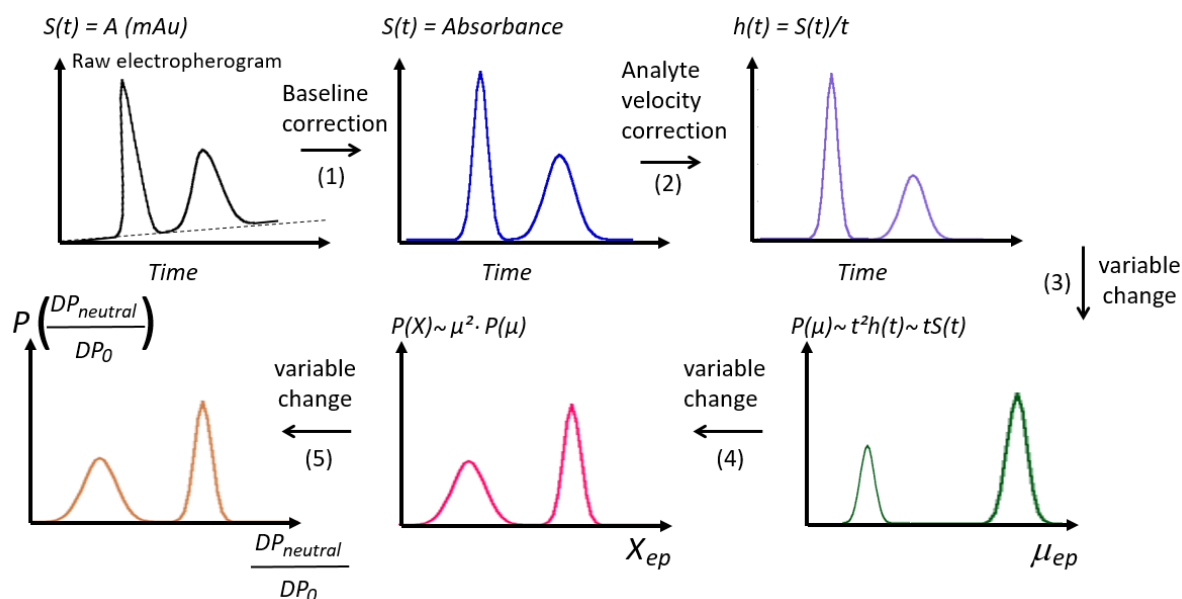


Figure 2. General scheme for changing a time-scale electropherogram into a mobility-scale and X_{exp} -scale distributions. The raw electropherogram is first corrected from baseline shift (1). The time-scale electropherogram is then corrected from the differences in analyte velocities (2). The time-corrected electropherogram is converted into the effective mobility-scale distribution (3). The mobility-scale electropherogram is changed into a X_{exp} -scale distribution (4) and finally to a compositional $\frac{DP_{\text{neutral}}}{DP_0}$ ratio (5).

$S(t)$ is the UV absorbance signal (in mAU). $h(t)$ is the time-corrected UV absorbance. $P(\mu_{\text{ep}})$ is the effective mobility distribution. $P(X_{\text{exp}})$ is the distribution in X_{exp} (see section 4.2) and $P(\frac{DP_{\text{neutral}}}{DP_0})$ is the distribution in $\frac{DP_{\text{neutral}}}{DP_0}$.

$\frac{DP_{\text{neutral}}}{DP_0}$. Adapted from¹¹ for the characterization of diblock copolymers.

Scale transformations. In this section, the transformations of the electropherograms into distributions of the parameter of interest are described following a previously published protocol¹¹. Briefly, experimental raw time-scale electropherograms, were first corrected from any baseline shift using Origin (Origin 2016, OriginLab, USA) as depicted in step 1, Figure 2. For quantitative purpose, the absorbance signal $S(t)$ was next divided by the migration time (t) to correct the differences in analyte

velocity (Figure 2, step 2)¹¹. Next, the time-corrected electropherogram $h(t)$ was changed into an effective mobility distribution $P(\mu_{ep}) = t \times S(t)$ (Figure 2, step 3)¹¹, which requires the transformations of both the x and y axis¹¹. Note that μ_{ep} is obtained by equation (5):

$$\mu_{ep} = \frac{lL}{V} \left(\frac{1}{t} - \frac{1}{t_{eo}} \right) \quad (5)$$

where l is the effective capillary length, L is the total capillary length, t is the migration time, t_{eo} is the EOF marker migration time and V is the separation voltage.

Moments of the electrophoretic mobility distribution The average effective mobility of the diblock copolymer $\overline{\mu_{ep}^{diblock}}$ was obtained by integration of the peak of the copolymer in the effective mobility scale according to:

$$\overline{\mu_{ep}^{diblock}} = \frac{\int P(\mu_{ep}) \mu_{ep} d\mu_{ep}}{\int P(\mu_{ep}) d\mu_{ep}} \approx \frac{\sum_i P(\mu_{ep,i}) \mu_{ep,i} (\mu_{ep,i+1} - \mu_{ep,i})}{\sum_i P(\mu_{ep,i}) (\mu_{ep,i+1} - \mu_{ep,i})} \quad (6)$$

where integration is carried out over the peak. In practice the integration is done numerically and the i index represents the digitized experimental data points. The summation is carried out over values of $P(\mu_{ep,i})$ greater than the median of the base line added to its standard deviation. Calculation of $\overline{\mu_{ep}^{diblock}}$ was performed using Excel 2016 (Microsoft, USA), following the discrete form of equation (6).

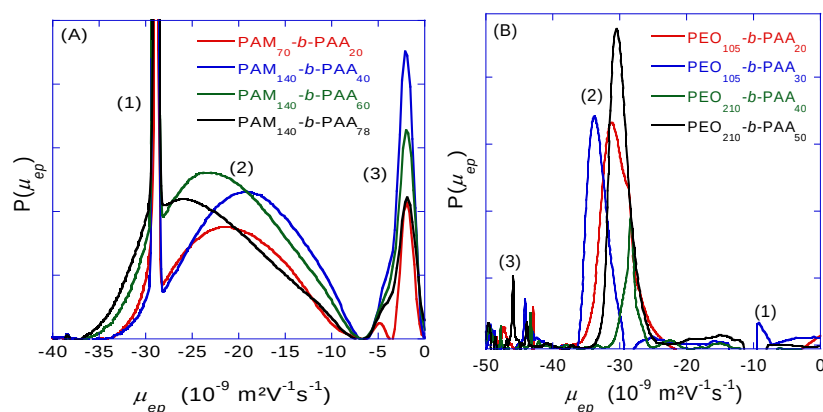
Variance of the diblock electrophoretic mobility was obtained by the following equation:

$$\sigma_{\mu}^2 = \frac{\int P(\mu_{ep}) \left(\mu_{ep} - \overline{\mu_{ep}^{diblock}} \right)^2 d\mu_{ep}}{\int P(\mu_{ep}) d\mu_{ep}} = \frac{\sum_i P(\mu_{ep,i}) (\mu_{ep,i} - \overline{\mu_{ep}^{diblock}})^2 (\mu_{ep,i+1} - \mu_{ep,i})}{\sum_i P(\mu_{ep,i}) (\mu_{ep,i+1} - \mu_{ep,i})} \quad (7)$$

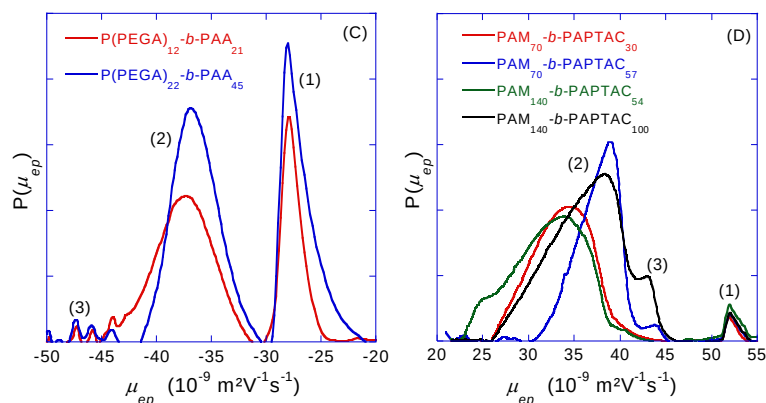
4. Results and discussion

4.1. Electrophoretic separation of the DHBC

The main goal of the CE characterization is to provide information about the chemical composition distribution of the DHBC and about the purity of the DHBC in terms of possible presence of homopolymers. The separation of the DHBC by CE requires an appropriate background electrolyte, depending on the nature of the copolymer. For copolymers absorbing in UV (i.e. those with a PAM neutral block), direct UV detection was possible and a background electrolyte based on 20 mM MES and 14 mM ammonium diiodate at pH 6.5 was used, with a UV detection at 192 nm. For DHBC copolymers that do not absorb UV enough to ensure sensitivity (i.e. PEO-*b*-PAA or P(PEGA)-*b*-PAA), an indirect detection mode based on a 6 mM anisic acid and 12 mM BIS TRIS at pH 6.5 was used. At this pH about 60% of the carboxylic acid groups of the PAA are ionized, and this ensures appropriate selectivity of separation between PAA homopolyelectrolyte and the DHBC. Uncoated fused silica capillary was used for the characterization of all anionic DHBC. Semi-permanent UltraTrolLN neutral coating was used for the characterization of the cationic PAM-*b*-PAPTAC DHBC, to avoid any copolymer adsorption on the capillary surface. To correct the apparent mobility from the electroosmotic mobility, a mobility marker (anisic acid for PAM-*b*-PAA, MES for P(PEGA)-*b*-PAA, and imidazolium for PAM-*b*-PAPTAC) of known effective mobility ($\mu_{ep, MES} = -28$ TU (where TU, Tiselius Unit, stands for $10^{-9} \text{ m}^2 \text{ V}^{-1} \text{ s}^{-1}$) and $\mu_{ep, imidazolium} = 52$ TU) was co-injected. For PEO-*b*-PAA, the electroosmotic mobility was estimated from the electroosmotic flow (EOF) peak. The distributions of effective mobility (DEM) of PAM-*b*-PAA, PEO-*b*-PAA, P(PEGA)-*b*-PAA and PAM-*b*-PAPTAC are displayed in Figure 3.



305



306

307 **Figure 3.:** Distributions of effective electrophoretic mobility obtained for PAM-*b*-PAA (A), PEO-*b*-
 308 PAA (B), P(PEGA)-*b*-PAA (C), and PAM-*b*-PAPTAC (D).-Electrophoretic conditions: fused silica
 309 capillary (A, B, C) or coated with UltraTrolLN™ (D), 50 μm I.D. × 38.5 cm (effective length, 30 cm).
 310 Electrolytes: 20 mM MES, 14 mM ammediol, pH 6.5 (A, D); 6 mM anisic acid, 12 mM BIS TRIS, pH
 311 6.5 (B, C). Applied voltage: +20 kV. Hydrodynamic injection: 30 mbar, 5 s. Direct (A, D) or indirect
 312 (B, C) UV detection at 192±2 nm. Temperature: 25 °C. Samples: 5 g/L DHBC. Assignment of the
 313 peaks: PAM-*b*-PAA (A): anisic acid (1), DHBC (2), PAM homopolymer (3); PEO-*b*-PAA (B): system
 314 peak (1), DHBC (2), PAA oligomers (3); P(PEGA)-*b*-PAA (C): MES (1), DHBC (2), PAA oligomers
 315 (3); PAM-*b*-PAPTAC (D): Imidazolium (1), DHBC (2), PAPTAC (3). The degree of polymerization of
 316 each block is specified on the graph.

317 The DEM in the series PAM-*b*-PAA (Figure 3A) show three peaks, two sharp at -29 TU (peak 1) and -2
 318 TU (peak 3) and one broad (peak 2) between -7 and -37.5 TU, the latter being assigned to the copolymer
 319 of interest. The peak at -29 TU is assigned to anisic acid (electrophoretic mobility marker) and the one
 320 at -2 TU corresponds to homopolymer of PAM. The non-zero electrophoretic mobility of the PAM
 321 homopolymer is explained by the incorporation of the negatively charged initiator 4,4'-azobis(4-
 322 cyanopentanoic) acid. The mass proportion of this PAM population of dead chains has been quantified
 323 by external calibration based on time-corrected peak areas using direct injections of PAM solutions of
 324 known concentration in the same condition as the DHBC. The proportion of PAM homopolymer
 325 amounts to 21wt% in the solid form polymer sample for PAM₇₀-*b*-PAA₂₀, 15% for PAM₁₄₀-*b*-PAA₄₀

and PAM₁₄₀-*b*-PAA₆₀ and 8% for PAM₁₄₀-*b*-PAA₇₈. The copolymer peak is broad and, as expected, the DEM shifts further from zero as the proportion of charged monomer increases in the composition of the copolymers. This can be verified by comparing the average electrophoretic mobility value $\overline{\mu_{ep}^{diblock}}$ (given in Table 2) which varies between -19.5 TU and -24.1 TU from PAM₁₄₀-*b*-PAA₄₀ to PAM₁₄₀-*b*-PAA₇₈. The greater the average molar mass of the polymer, the more dispersed its electrophoretic mobility, as demonstrated by the standard deviation σ_{μ} which varies from 3.9 TU for PAM-*b*-PAA 70-20, to 6.2 TU for PAM-*b*-PAA 140-60.

The DEM of PEO based copolymers are presented in Figure 3B for linear PEO and Figure 3C for PEO grafted polyacrylates (P(PEGA)). Three populations are observed in both series: several small peaks associated with large electrophoretic mobility at -45TU are assigned to short oligomers of PAA, the weight percent of which is estimated to be lower than 10%. The sensitivity of the UV detection is too low to conclude about the presence / absence of PEO or P(PEGA) in the DHBC. The least mobile species at -28TU in Figure 3C corresponds to the MES used as mobility marker. The peak at intermediate values of mobility corresponds to the DHBC. The electrophoretic mobility of PEO-*b*-PAA ($\overline{\mu_{ep}^{diblock}}$ ranging from -28 to -33 TU, Table 2) is significantly closer to zero than that of P(PEGA)-*b*-PAA copolymers ($\overline{\mu_{ep}^{diblock}}$ ranging from -36 to -38 TU, Table 2), although the molar masses are close. This is because P(PEGA), a comb-like polymer, is more compact than linear PEO of the same molar mass. As a consequence, the drag force due to the neutral block is lower for P(PEGA) than for PEO.

As for PAM-*b*-PAPTAC copolymer, PAPTAC homopolymer was detected at about +43TU, as a shoulder merged in the copolymer distribution, only for the $DP_{neutral}/DP_0$ equal to 140/100 and 70/57 samples. Figure 3D displays DEM ranged between +22 and +41 TU, with higher effective mobilities for the DHBC of highest charge. Comparison of PAM-*b*-PAA series with PAM-*b*-PAPTAC series illustrates the importance of the nature of the blocks on the drag effect of the neutral block. This effect is discussed in more detail in section 4.2.

As a general trend, electrophoretic mobilities of DHBC are always closer to zero than those of the homopolyelectrolyte which are: $\mu_{\text{PAA}} = -42$ TU; $\mu_{\text{PAPTAC}} = +44$ TU, and the electrophoretic mobility of the DHBC increases as the proportion of charged monomers in the DHBC increases (see Figure SI 20). In terms of EM dispersion, the least dispersed series is the P(PEGA)-*b*-PAA, with relative standard deviation of EM $\sigma_{\mu} / \overline{\mu_{ep}^{diblock}}$ between 6.5% to 9%, followed by the PEO-*b*-PAA ($\sigma_{\mu} / \overline{\mu_{ep}^{diblock}} \sim 5\%$ to 16%) and the most disperse series is the PAM-*b*-PAA series with 20% to 30% relative standard deviation of the electrophoretic mobility. This dispersion in mobility results from both the level of control of the polymerization, and from the spatial extension of the polymer in the solvent. It can be explained by the chemistry of the RAFT polymerization (chain transfer agent R-SC(S)Z where Z is the activating group and R is the leaving group), for which the polymerization of acrylates is better controlled by dithiobenzoates (Z: -SC(S)Ph) ($\mathcal{D} < 1.2$) than by ethyl xanthate (Z: -SC(S)OEt) ($\mathcal{D} > 1.3$). Besides, the 2-phenylacetate ester of PEO is a better homolytic leaving group than PAM.³⁶ Furthermore, the synthesis of PAM-*b*-PAA cumulates two successive RAFT/MADIX polymerizations of AM and AA, whereas the synthesis of PEO-*b*-PAA starts from a narrow PEO-CTA ($\mathcal{D} = 1.04$) (PEO obtained by anionic polymerization) to perform a single RAFT polymerization of AA. This is consistent with the higher dispersity of PAM-*b*-PAA compared to PEO-*b*-PAA³¹. In addition, in reversible-deactivation radical polymerization³⁷, at full conversion, the dispersity as defined by the ratio of the weight average molar mass over the number average molar mass, decreases when the *DP* increases³⁸:

$$\mathcal{D} = 1 + \frac{1}{DP} + \frac{1}{C_{ex}} \quad (8)$$

where *DP* is the targeted polymerization degree and C_{ex} is the degenerative chain transfer constant between dormant and active chains^{39, 40}, which is consistent with a higher dispersity for a shorter poly(acrylic acid) block in PEO-*b*-PAA.

To get a better description of the copolymer distribution, it would be interesting to get a distribution of a new parameter which is directly related to the chemical composition of the copolymer instead of the

electrophoretic mobility, which is not linearly dependent on the copolymer composition. This is the purpose of the two following sections.

	$DP_{neutral-}$ DP_0	$\mu_{ep}^{diblock}$ peak max (TU)	$\overline{\mu_{ep}^{diblock}}$ integration (TU)	σ_{μ} (TU)	X_{exp} peak max	$\overline{X_{exp}}$ by integration	$\sigma_{X_{exp}}$
PAM- <i>b</i> -PAA	70-20	-22	-19.9	3.89	0.61	1.12	0.45
	140-40	-19.1	-19.5	5.44	0.74	1.26	0.73
	140-60	-23.3	-21.4	6.19	0.58	1.05	1.0
	140-78	-25.3	-24.1	5.68	0.41	0.99	0.56
PEO- <i>b</i> -PAA	105-20	-31	-30.2	4.83	0.28	0.33	0.12
	105-30	-33.8	-33.2	2.45	0.18	0.21	0.06
	210-40	-28	-28.4	1.36	0.40	0.41	0.06
	210-50	-30.5	-30	1.97	0.30	0.34	0.64
P(PEGA)- <i>b</i> -PAA	21-11.5	-37.6	-38.6	3.61	0.059	0.043	0.09
	44.9-22.2	-37	-36.6	2.41	0.079	0.098	0.08
PAM- <i>b</i> -PAPTAC	70-30	34.6	33.9	3.18	0.19	0.25	0.12
	70-60	39.1	37.1	3.2	0.08	0.11	0.05
	140-60	34.2	32	4.1	0.21	0.32	0.39
	140-120	38.5	36.3	4.1	0.09	0.164	0.13

Table 2. Electrophoretic mobility $\mu_{ep}^{diblock}$ at peak maximum and average value $\overline{\mu_{ep}^{diblock}}$ obtained by peak integration, standard deviation of the electrophoretic mobility distribution σ_{μ} , X_{exp} value at peak maximum and average value $\overline{X_{exp}}$ obtained by peak integration, standard deviation of the X_{exp} distribution $\sigma_{X_{exp}}$ of all DHBC

studied in this work. Note that the peak of anisic acid was first subtracted before integration for the calculation of

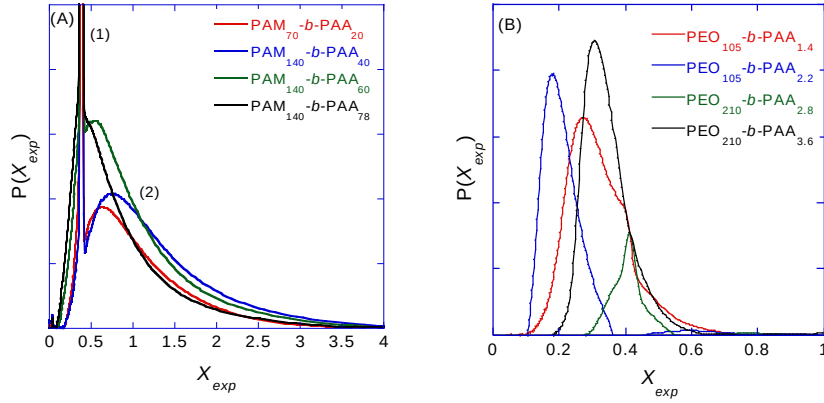
$\overline{\mu_{ep}^{diblock}}$ and σ_{μ} for PAM-*b*-PAA.

4.2. Change of variable from μ_{ep} to the retardation parameter X_{exp}

The drag effect of the neutral block can be expressed by the retardation parameter X_{exp} defined as :

$$X_{exp} = \frac{\mu_{ep}^0 - \mu_{ep}^{diblock}}{\mu_{ep}^{diblock}} = \frac{\mu_{ep}^0}{\mu_{ep}^{diblock}} - 1 \quad (9)$$

where μ_{ep}^0 is the electrophoretic mobility of the homopolyelectrolyte; $\mu_{ep}^{diblock}$ is the electrophoretic mobility of the copolymer. X_{exp} expresses the relative decrease of mobility due to the presence of the neutral block. It is positive and increases as the drag effect increases.



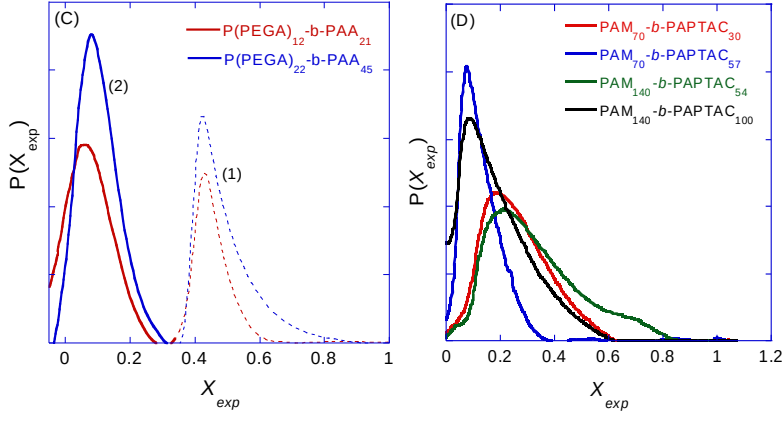


Figure 4. Distribution of retardation parameter X_{exp} for PAM-*b*-PAA (A) PEO-*b*-PAA (B), P(PEGA)-*b*-PAA (C), and PAM-*b*-PAPTAC (D). Experimental conditions as in Figure 3 X_{exp} was determined using eq. (8), eq. (12) and $\mu_{ep,PAA}^0 = -42\text{TU}$; $\mu_{ep,PAPTAC}^0 = 44\text{ TU}$. Assignment of the peaks: PAM-*b*-PAA (A): anisic acid (1), DHBC (2); P(PEGA)-*b*-PAA (C): MES (1), DHBC (2). The degree of polymerization of each block is specified on the graph.

The new experimental variable, X_{exp} , not only points out the friction due to the neutral block but is also more directly related to the composition of the DHBC. Introducing equation (9) in the various expressions of the electrophoretic mobility (equations 1 to 3), X_{exp} , can be expressed as a function of the ratio between the degrees of polymerization of the neutral $DP_{neutral}$ and the charged blocks DP_0 . Taking into account the hydrodynamic coupling (model 1), X_{exp} reads:

$$X_{model,1} = \frac{1}{N_{blob}} = \frac{\alpha}{DP_0} \quad (10a)$$

and can be further developed as a function of the Kuhn lengths of the neutral block, b_{K_1} , and of the polyelectrolyte, b_{K_0} :^{15, 41}

$$X_{model,1} = \alpha_1 \frac{DP_{neutral}}{DP_0} \quad (10b)$$

408 with $\alpha_1 = \frac{b_1 b_{K_1}}{b_0 b_{K_0}}$ (10c)

409 where b_1 is the neutral monomer size. The Kuhn statistical segment length (which is twice the persistence
 410 length) is a measure of the polymer stiffness. Parameter α_1 in Equation (10b) is a relative friction
 411 coefficient and it is non-dimensional. Since the polyelectrolyte is generally stiffer than the neutral block,
 412 α_1 is often much smaller than unity¹⁵.

413 As for model 2, it is clear from equation (2) that the X parameter is directly expressed as the ratio of the
 414 hydrodynamic radius of each block:

415 $X_{model,2} = \frac{R_h^{neutral}}{R_h^0}$ (11a)

416 which can be rewritten as a function of the degrees of polymerization of each block by:

417 $X_{model,2} = \frac{C_1 DP_{neutral}^{a_1}}{C_0 DP_0^{a_0}}$ (11b)

418 where C_1 (resp. C_2) and a_1 (resp. a_2) are, respectively, the prefactors and exponents for the neutral (resp.
 419 charged) block in the relationship between R_h and DP . Note that a_0 and a_1 are supposed to be close to
 420 0.5-0.6 for coil conformations, and slightly higher for more extended conformations.

421 As for model 3 (see equation (3)), the X parameter is directly expressed as:

422 $X_{model,3} = \frac{2 R_h^{neutral} \ln DP_0}{b_0 DP_0}$ (12a)

423 Injecting $R_h^{neutral}$ in equation (9a) leads to:

424 $X_{model,3} = \frac{2 C_1 DP_{neutral}^{a_1} \ln DP_0}{b_0 DP_0}$ (12b)

Finally, equations (10b) (11b) and (12b) demonstrates that, whatever the considered model, the X parameter is related to a compositional ratio between the neutral and the charged blocks with, however, different scaling dependences with the DP of each block, and with an additional logarithmic term in model 3. It is worth noting that in the case of hydrodynamic coupling (model 1), the newly introduced variable varies linearly with the ratio of degree of polymerization of the two blocks.

The distributions of the X_{exp} parameter are simply deduced from the distribution of electrophoretic mobility using the following equation (Figure 2, step 4)¹¹:

$$P(X_{exp}) = \left| \frac{1}{\frac{\partial X_{ep}}{\partial \mu_{ep}}} \right| P(\mu_{ep}) = \mu_{ep}^2 P(\mu_{ep}) \quad (13)$$

All the X_{exp} distributions are presented in Figure 4. Since μ_{ep} and X_{exp} are not linearly related, the change of the variable from μ_{ep} to X_{exp} modify the form of the distribution. The different moments characterizing the distribution of X_{exp} are calculated using similar relations as equations (6) and (7) and are reported in Table 2.

X_{exp} range varies between 0.1-3 for PAM-*b*-PAA, 0.1-0.7 for PEO-*b*-PAA, 0-0.3 for P(PEGA)-*b*-PAA and 0-0.8 for PAM-*b*-PAPTAC. The dispersion of the retardation parameter X_{exp} expressed as $\sigma_{X_{exp}}$ values (Table 2) follows the following order: P(PEGA)-*b*-PAA < PEO-*b*-PAA ~ PAM-*b*-PAPTAC < PAM-*b*-PAA. The dispersion of the retardation parameter cannot be interpreted as a dispersity in molar mass or in composition since the retardation parameter will change with these chemical features in a way that depends on the conformation of the blocks. So, a further step is needed to get the composition dispersion.

4.3. Change of variable from X_{exp} to chemical composition ratio

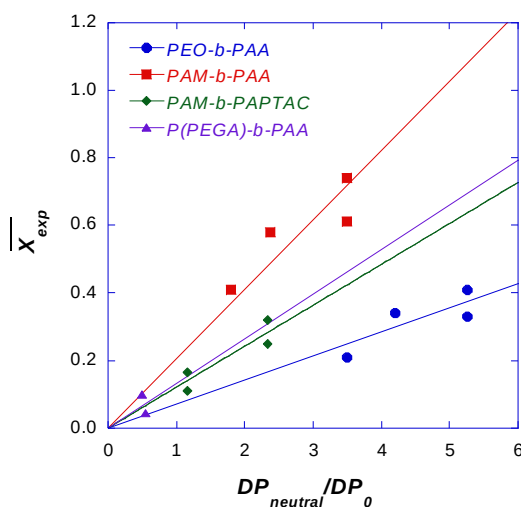


Figure 5. Plot of $\overline{X}_{exp}$ against $\frac{DP_{neutral}}{DP_0} \cdot \overline{X}_{exp}$ was determined by using eq.(6) after replacing μ_{ep} by

$X_{exp} \cdot \frac{DP_{neutral}}{DP_0}$ was obtained by NMR (see Table 1). Least-square linear regressions provide the following

experimental slopes: 0.071+/-0.005 for PEO-b-PAA; 0.12+/-0.01 for PAM-b-PAPTAC; 0.13 +/- 0.01 for P(PEGA)-b-PAA; 0.21+/-0.02 for PAM-b-PAA.

To go further in the interpretation and in the process of the experimental data, it is crucial to identify the model which is best adapted to describe the electrophoretic behavior of the DHBC investigated in this work. To assess the validity of model 1 (hydrodynamic coupling between blocks, see section 2.3), $\overline{X}_{exp}$ was plotted against $\frac{DP_{neutral}}{DP_0}$ in Figure 5 for the four DHBC families.

Table 3. Characteristic parameters of neutral and charged blocks constituting the DHBC studied in this work. M_w are expressed in g/mol.

	b_l (nm)	b_{K_1} (nm)	$R_h^{neutral}$ (nm) ^a	
PAM	0.25	0.6 ⁴²	$0.01447 \times M_w^{0.57}$	457
PEO	0.3 ⁴⁴	0.74 ⁴⁴	$0.02398 \times M_w^{0.53}$	459
	b_0 (nm)	b_{K_0} (nm)	R_h^0 (nm)	
PAA	0.25	2.5 ⁴⁶	$0.007906 \times M_w^{0.585}$	461
PAPTAC	approximated as PAA			462

^a From Mark-Houwink parameters using $R_h = \left(\frac{3[\eta]M}{10\pi N_A} \right)^{1/3}$, where $[\eta]$ is the intrinsic viscosity and N_A is the Avogadro number. from ref⁴⁶ (see Table 1 and Figure 13(a) herein).

The results are consistent with model 1 which predicts a linear dependence of the retardation parameter $\overline{X_{exp}}$ on the ratio of degree of polymerization $\frac{DP_{neutral}}{DP_0}$. The slopes determined from the graph in Figure 5 correspond to the parameter α_1 in equation (10b) which can be calculated from equation (10c). Experimentally, the numerical values of the slopes $\alpha_{1,exp}$ are in the range of ~0.1-0.2. Taking the characteristic parameters (Kuhn lengths, monomer dimensions) given in Table 3 leads to $\alpha_l=0.24$ (vs $\alpha_{1,exp}=0.21\pm0.02$ experimentally obtained) for PAM-*b*-PAA, $\alpha_l=0.36$ (vs $\alpha_{1,exp}=0.071\pm0.005$) for PEO-*b*-PAA, and $\alpha_l=0.24$ (vs $\alpha_{1,exp}=0.12\pm0.01$) for PAM-*b*-PAPTAC. Theoretical values of α_l are in a reasonably good agreement with the experimental ones, knowing the uncertainty on the persistence length (notably for the polyelectrolyte blocks) and monomer sizes. As for P(PEGA)-*b*-PAA, we only get an estimation of $\alpha_{1,exp}=0.13$, since the P(PEGA) Kuhn length is not available in the literature. From Figure 5, we can conclude that the linear correlation between $\overline{X_{exp}}$ and $\frac{DP_{neutral}}{DP_0}$ is confirmed and that model 1 (with hydrodynamic coupling between the two blocks) can be used to transform the X_{exp}

479 distributions into compositional $\frac{DP_{neutral}}{DP_0}$ distributions. As for models 2 and 3, they lead to poor

480 correlations between theoretical $X_{model,i}$ versus experimental $\overline{X_{exp}}$ values (see Figure 6).

481 The knowledge of α_l provides the last relation necessary to carry on the general scheme presented in
 482 Figure 2 to its end and which leads to the distribution of ratio of chemical composition. In practice, we
 483 used $\alpha_{l,exp}$ obtained in Figure 5 together with equation (10b) to transform the data of Figures 4A to 4D
 484 into the distributions presented in Figures 7A to 7D, using the following equation:

$$485 \quad P\left(\frac{DP_{neutral}}{DP_0}\right) = \frac{P(X)}{\frac{\partial\left(\frac{DP_{neutral}}{DP_0}\right)}{\partial X}} = \alpha_l P(X) \quad (14)$$

486 Since X_{exp} and $\frac{DP_{neutral}}{DP_0}$ are linearly correlated, the shapes of both distributions are similar. However,

487 reading $\frac{DP_{neutral}}{DP_0}$ axis, which corresponds to a compositional ratio, is more convenient for the practitioners

488 than keeping the X_{exp} scale. Moreover, and as previously anticipated, since the α_l coefficients are

489 different from one DHBC to another, the distribution in $\frac{DP_{neutral}}{DP_0}$ allows a better comparison between

490 them. On the whole, the dispersion of the composition ratio are in the order of: P(PEGA)-*b*-PAA <

491 PAM-*b*-PAPTAC ~ PEO-*b*-PAA < PAM-*b*-PAA.

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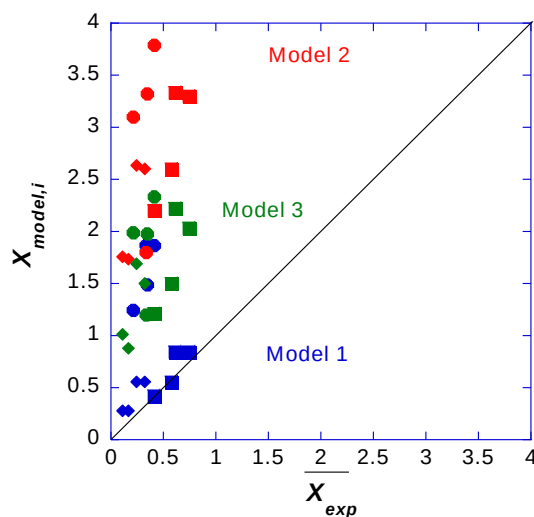
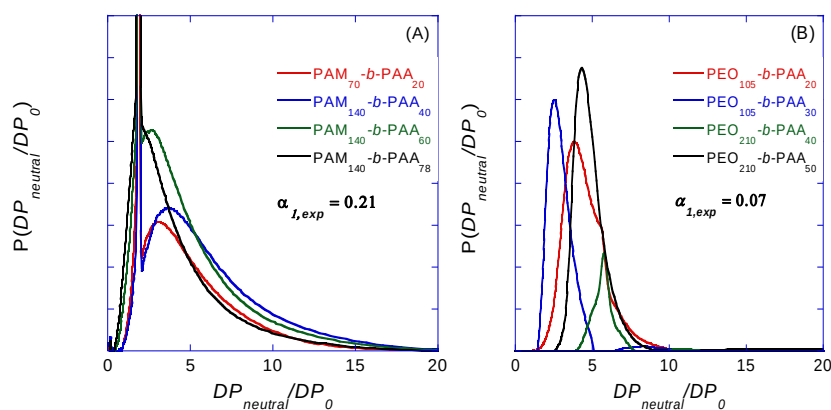


Figure 6. Comparison between $X_{model,i}$ and $\overline{X}_{exp}$ for the three different models using theoretical parameters given in Table 3. PAM-*b*-PAPTAC (♦); PEO-*b*-PAA (●); PAM-*b*-PAA (■). $\overline{X}_{exp}$ was determined by integration of the DHBC peak (in X scale). For all $X_{model,i}$ calculations, theoretical $DP_{neutral}$ and DP_0 were used. $X_{model,1}$ was determined according to equations (10b) and (10c), $X_{model,2}$ according to equation (11a) and $X_{model,3}$ according to equations (12a), with the characteristic numerical parameters given in Table 3.



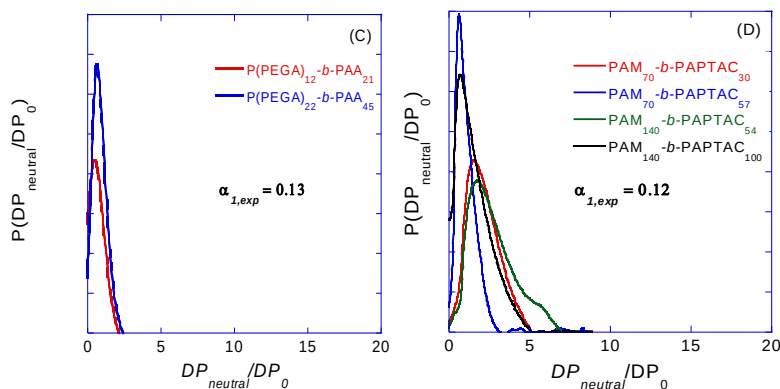


Figure 7. Distribution of composition in terms of the ratio of the degrees of polymerization of both blocks for PAM-*b*-PAA (A), PEO-*b*-PAA (B), P(PEGA)-*b*-PAA (C), and PAM-*b*-PAPTAC (D). Experimental conditions as in Figure 3. $\frac{DP_{neutral}}{DP_0}$ was determined using eq. (10b) and $P\left(\frac{DP_{neutral}}{DP_0}\right)$ was obtained using eq. (14). In (C), the MES peak has been removed. In (A), the sharp peak is a mobility marker (anisic acid) and was deleted before peak integration.

Conclusion

In an effort to make information provided by capillary electrophoresis more directly useful for polymer chemists, a protocol was proposed to convert electrophoretic mobility distributions of double hydrophilic block copolymers into distributions of chemical composition ratios. This ratio of composition is expressed as the ratio of the degrees of polymerization of each block $\frac{DP_{neutral}}{DP_0}$. To get this composition ratio, we have introduced the retardation parameter X which takes into account the drag force exerted by the neutral block on the polyelectrolyte. The distribution of X that characterizes a DHBC is readily obtained from the experimental electropherogram and the relation between X and the ratio of DP . The latter is available from different models for electrophoretic mobility of composite objects. A linear relation has been found experimentally between the retardation parameter X and the ratio of DP s, within each of the four families of DHBC studied in this work. This result is consistent with the model of electrophoretic mobility of Long *et al.* that takes into account hydrodynamic coupling, although the

prefactors are slightly overestimated. Our experimental findings not only support the theoretical prediction, but also facilitate the last step of data transformation, from distribution of X into distributions of ratios of DP .

The dispersions in composition are in the order of: P(PEGA)-*b*-PAA < PEO-*b*-PAA ~ PAM-*b*-PAPTAC < PAM-*b*-PAA. Therefore, we can conclude that in the DHBC families, the PAM block leads to broader composition ratio distributions compared to a PEO block, when associated to a PAA block. Similarly, the P(PEGA) block lead to less disperse composition ratio distributions compared to a PEO block, when associated to a PAA block. Finally, PAA associated with PAM leads to broader composition ratio distributions compared to PAPTAC associated with PAM. The relatively low composition dispersity of the PEO-PAA block copolymer is most likely due to the low dispersity of the PEO block, prepared by anionic polymerization, and the use of a dithiobenzoate chain transfer agent, which has a higher chain transfer constant and thus gives narrower molar mass distributions than the xanthate chain transfer agent used to prepare the PAM-PAA and PAM-PAPTAC block copolymers. The use of a trithiocarbonate chain transfer agent and a relatively short P(PEGA) block leads to a fairly narrow composition distribution for P(PEGA)-PAA block copolymers.

Finally, the transformation of electrophoretic mobility distributions into composition ratio distributions significantly improved the comparison of the distributions between the different copolymer families, since it takes into account the differences in expansion and drag force according to the chemical nature of the blocks.

Supporting Information.

Synthetic pathway, SEC and NMR characterizations, raw electropherograms are provided for all dibloc copolymers studied in this work.

543 Acknowledgements

544 This work was supported by the MESOPIC project funded by the Agence Nationale de la Recherche
545 (ANR) under grant # ANR-15-CE07-0005.

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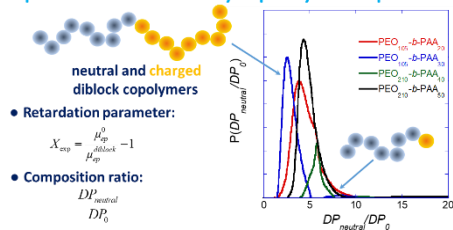
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Composition distributions by Capillary Electrophoresis



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