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# PPEGMEMA-based cationic copolymers designed for layer-by-layer assembly†

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We have synthesized three PPEGMEMA-based cationic copolymers with similar amine contents but with systematic variation in the average length of the PEG side chains. The positively charged copolymers were paired with alginate to fabricate layer-by-layer assembled multilayered films. It was demonstrated that the polymeric design, in terms of the systematic variation in the average length of the PEG units, affects the polyelectrolyte multilayer growth mechanism and can be used to tune the structural properties and the water content of the layers. In addition, by partial cross-linking of the amine groups present in the copolymer backbone, disintegration of the film induced by pH changes was prevented. Finally, it was demonstrated how the cross-linked multilayered film can exhibit cationic, zwitterionic and anionic properties depending on the pH value and how these changes are associated with swelling, layer contraction and changes in water content.

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## 1. Introduction

Layer-by-layer (LbL) assembly has drawn great attention as a convenient and adaptable method to fabricate functional polymer-coated surfaces.<sup>1–7</sup> The designated polymeric components are sequentially adsorbed onto a solid substrate, driven by different intermolecular forces, such as electrostatic attraction,<sup>2,3,8–10</sup> hydrogen bonding,<sup>11–14</sup> and covalent bonding.<sup>15,16</sup> The electrostatic LbL assembly method has been extensively investigated due to its simplicity and high efficiency.<sup>1</sup> Accordingly, polyelectrolyte multilayered films can be readily prepared by alternate immersion of a charged substrate into solutions of oppositely charged polyelectrolytes with rinsing steps in between. To this end, the thicknesses and composition of the layer are controlled by the number of deposited layers, pH,<sup>10,17,18</sup> ionic strength,<sup>10,19,20</sup> temperature,<sup>21</sup> adsorption time,<sup>22</sup> and polymer molecular weight.<sup>23</sup>

The need for ionizable groups in the backbone of the polymers for electrostatic LbL assembly effectively hinders the incorporation of uncharged polymers into the polyelectrolyte multilayered films. It is, however, feasible to accommodate desired functionalities into the backbone of charged polymers by chemical modification. Several authors have implemented such an approach to tune the functionality of multilayered films obtained by electrostatic LbL assembly.<sup>24–26</sup> For instance, Sun and coworkers modified polyethyleneimine (PEI) with polyethylene glycol (PEG) and fabricated a multilayered film based

on PEGylated-PEI and hyaluronic acid. The obtained film demonstrated antifouling properties resulting from the grafted PEG units.<sup>24</sup> In another relevant work, Neoh and coworkers custom synthesized poly(isobutylene-*alt*-maleic acid) (PIAMA) derivatives bearing either alkyl- or PEG-carboxyl side chains, and LbL assembled the obtained polyanions with PEI.<sup>25</sup> It was shown that varying the side chains in the polyanion affects the wettability of the multilayered film; hence, such a film could be used to tune the protein adsorption and cell adhesion properties of the surface. In both the works mentioned above, the functional units (PEG or alkyl chains) are grafted onto a polyelectrolyte backbone, which has limitations in terms of tuning the properties of the polyion such as charge density and molecular architecture. A more versatile approach is to copolymerize neutral and charged monomers instead, where the former unit provides functional properties and the latter is responsible for the electrostatic interaction needed for LbL assembly. For instance, Schlenoff and coworkers copolymerized *N*-isopropylacrylamide (NIPAM, neutral monomer) with allylamine (cationic comonomer) as well as styrene sulfonic acid (anionic comonomer).<sup>27</sup> The synthesized PNIPAM-based copolymers were then paired to fabricate a thermoresponsive multilayered film. Based on this approach, one can thus control not only the chemical composition but also the charge density, molecular weight and chain architecture of the components in the multilayered film.

Poly(polyethylene glycol methyl ether methacrylate) (PPEGMEMA) is a methacrylate derivative of polyethylene glycol that has been vastly studied as a potential coating material due to its hydrophilic nature and high water content, which leads to lubricating<sup>28,29</sup> and protein-repellent properties.<sup>30–33</sup> A common method to fabricate PPEGMEMA-coated surfaces is surface-

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initiated polymerization.<sup>34–37</sup> However, this method has several disadvantages that can restrict its applicability. It is generally challenging to achieve a thick coating using surface-initiated polymerization, which is a less versatile and scalable method that requires special substrate treatment and reaction control. In contrast, LbL assembly can be conducted through various methods, *e.g.*, dip-coating, spray-coating and spin-coating, which not only covers a broader range of substrates but can also render considerably thicker coatings.<sup>7</sup> Therefore, an alternative way to fabricate PEGMEMA-based coatings is to incorporate PEGMEMA units into charged chains so that electrostatic LbL assembly can be utilized.

Adopting this approach, we have synthesized a PEGMEMA-based cationic copolymer in which PEGMEMA (neutral monomer) was randomly copolymerized with aminoethyl methacrylate (cationic comonomer). Three PEGMEMA copolymer samples with comparable degrees of polymerization and amine content, but with different average lengths for the PEG side chains were prepared and then paired with alginate as a reference anionic polymer. This approach provided a method to study how systematic variation of the polymer architecture affects the multilayer buildup process and the multilayer structure. For this purpose, the assembly of the multilayered films was simultaneously monitored with a quartz crystal microbalance with dissipation monitoring (QCM-D) and spectroscopic ellipsometry. Accordingly, information for the thickness, conformation and water content of the films was extracted. Finally, the film structure was stabilized by cross-linking the amine groups, which preserved the pH-responsivity of the film while preventing its charge-induced disintegration.

## 2. Experimental section

### 2.1. Materials

Di(ethylene glycol) methyl ether methacrylate (DEGMEMA, 95%), poly(ethylene glycol) methyl ether methacrylate (PEGMEMA) with number average molecular weights ( $M_n$ ) of 300 and PEGMEMA with  $M_n$  of 500 g mol<sup>-1</sup> were purchased from Sigma Aldrich. All three monomers were treated with neutral alumina column chromatography to remove the inhibitors before use. Sodium alginate (NaAlg, medium viscosity  $\geq 2.000$  cP, 2% (25 °C)), (2-Boc-amino)ethyl methacrylate (BocAMA, 99%), diethyl *meso*-2,5-dibromoadipate (98%), glutaraldehyde (50% aqueous solution), trifluoroacetic acid (TFA, 99%), (3-aminopropyl)triethoxysilane (APTES, 99%) and copper(i) chloride (CuCl, >99%, washed sequentially with acetic acid and ethanol before use) was purchased from Sigma Aldrich and used as received unless otherwise stated. Tris(2-dimethylaminoethyl)amine (Me6TREN, 99%) was purchased from Alfa Aesar, USA. All the solvents used in this work were obtained from Sigma Aldrich and of HPLC grade. A pH 3 citric acid/sodium phosphate buffer (1.62 g citric acid, 0.56 g sodium phosphate dibasic dehydrate in 500 mL water) was used for the preparation of the polymer solutions as well as for rinsing in between each deposition step. Solutions of 100 ppm (w/v) PEGMEMA and alginate were prepared for LbL assembly and filtered through 0.44  $\mu$ m regenerative cellulose

syringe filters to remove possible aggregates and dust. Ultra-pure water (Milli-Q Plus system, resistivity of 18.2 M $\Omega$ -cm, TOC value less than 5 ppb.) was used to prepare all the solutions.

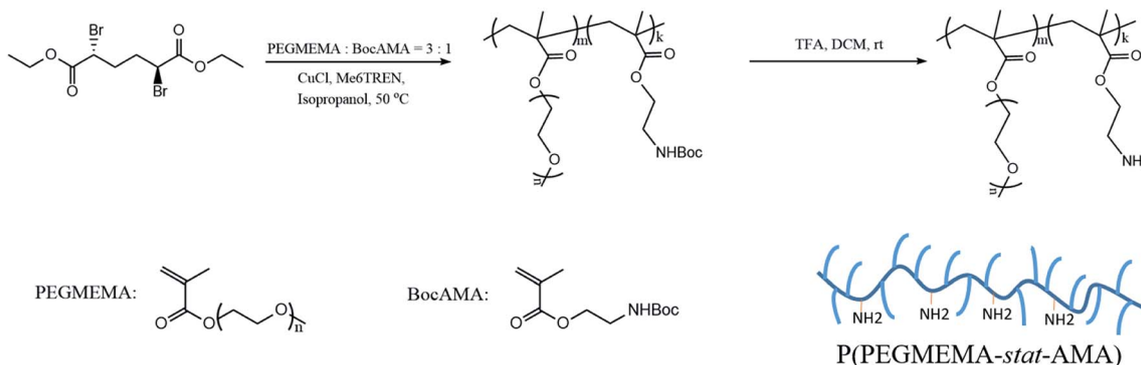
### 2.2. Synthesis of poly(poly(ethylene glycol) methyl ether methacrylate)-*stat*-poly(aminoethyl methacrylate) (P(PEGMEMA-*stat*-AMA))

Atom transfer radical polymerization (ATRP) is a well-developed living radical polymerization (LRP) technique, which has been widely utilized in preparation of polymers with narrow molecular weight distribution and macromolecular engineering.<sup>38</sup> PEGMEMA copolymers with randomly distributed primary amine groups were synthesized by ATRP with a protection-deprotection strategy according to previous reports.<sup>14,39,40</sup> ATRP in the presence of primary amine groups (the cationic group) is challenging due to possible complex formation between the amine groups and Cu(i), which may readily inactivate the catalyst; thus, a two-step protocol was employed to overcome this problem. First, PEGMEMA was copolymerized with protected BocAMA, yielding a P(PEGMEMA-*stat*-BocAMA) statistical copolymer. In the second step, Boc groups were hydrolyzed under acidic condition (TFA/DCM), yielding the targeted P(PEGMEMA-*stat*-AMA) copolymer (Scheme 1).

The detailed ATRP experimental procedure is as follows. For the synthesis of the protected P(PEGMEMA-*stat*-BocAMA), diethyl *meso*-2,5-dibromoadipate (initiator, 7.2 mg, 0.02 mmol), PEGMEMA (monomer, 3 mmol), (2-Boc-amino)ethyl methacrylate (monomer, 229.27 mg, 1 mmol) and Me6TREN (ligand, 9.2 mg, 0.04 mmol) were dissolved in 5 mL isopropanol. Dimethylformamide (0.1 mL) was added to the reaction mixture as an internal standard for subsequent calculation of the monomer conversion. The reaction mixture was bubbled with nitrogen gas for 30 min, after which the prewashed copper(i) chloride (1.23 mg, 0.0125 mmol) was added. The polymerization was triggered by immersing the reaction flask into a 50 °C water bath. After 16 hours, the reaction was quenched by adding 5 mL of water into the solution and by exposure to air. A small volume of the solution (0.1 mL) was collected for <sup>1</sup>H NMR measurement to determine the reaction conversion. The polymer solution was then purified *via* dialysis (regenerative cellulose tubing with MWCO 6–8 kDa, Spectrum Laboratories, Inc.) against water for 3 days. Finally, residual water was removed *via* lyophilization to obtain the pure P(PEGMEMA-*stat*-BocAMA) copolymers. Fig. 1 presents the <sup>1</sup>H NMR spectra for one PEGMEMA-based copolymer before and after the deprotection process. Accordingly, the peak corresponding to *t*-butyl (1.43 ppm) in the Boc protecting group is eliminated after the deprotection process.

In the deprotection process, the obtained P(PEGMEMA-*stat*-BocAMA) copolymers (300 mg) were dissolved in 4 mL dichloromethane followed by the addition of 0.5 mL of trifluoroacetic acid. The reaction mixture was stirred at room temperature for 3 hours, after which dichloromethane was removed by evaporation. The viscous polymer residue was redissolved in 5 mL of water, and the pH was adjusted to 7 with





Scheme 1 Synthesis procedure used for the P(PEGMEMA-*stat*-AMA) polymer.

1 M NaOH solution. The polymer was purified by dialysis and subsequent lyophilization, yielding a light-yellow viscous liquid.

All the P(PEGMEMA-*stat*-AMA) copolymers in this work were synthesized with the same procedure described above. In order to obtain polymers with a similar degree of polymerization (DP), the monomer to initiator ratio is fixed at 200 : 1 for all polymerizations, and the reaction time is set to be the same value of 16 hours to achieve high conversion (>80%) for all three polymerization processes. To determine the DPs of the polymers obtained, samples of the reaction mixture were collected before and after the polymerization process, and  $^1\text{H}$  NMR spectra (Bruker 400 MHz NMR spectrometer) for both samples were taken using deuterated DMSO as the solvent. The integral ratios of  $\delta$  6.0 ppm (monomer double bond (vinyl))/ $\delta$  7.9 ppm (DMF) before ( $r_1$ ) and after ( $r_2$ ) the polymerization were used to obtain the monomer conversion ( $c$ ) via  $c = 1 - r_2/r_1$ . The DP values were calculated accordingly and shown in Table 1.

The average number of ethylene oxide units in PEGMEMA monomers were tuned, in order to obtain P(PEGMEMA-*stat*-AMA) copolymers with different PEG side chain lengths. Overall

three P(PEGMEMA-*stat*-AMA) copolymers, namely P((EO)<sub>2.5</sub>-MEMA-*stat*-AMA), P((EO)<sub>4.5</sub>-MEMA-*stat*-AMA) and P((EO)<sub>9</sub>-MEMA-*stat*-AMA), with average PEGMEMA monomer molecular weights of 211, 300 and 500, corresponding to an average 2.5, 4.5 and 9 ethylene oxide units in the PEG chains in each monomer were synthesized, respectively. To this end, PEGMEMA with average molecular weights of  $M_n = 300$  and  $M_n = 500$ , were used for the synthesis of P((EO)<sub>4.5</sub>-MEMA-*stat*-AMA) and P((EO)<sub>9</sub>-MEMA-*stat*-AMA), respectively, while a mixture of PEGMEMA with average molecular weights of  $M_n = 188$  and  $M_n = 300$  in the ratio of 4 : 1 was used for the synthesis of P((EO)<sub>2.5</sub>-MEMA-*stat*-AMA). For convenience, from now we will refer to these three copolymers as s-PPEGMEMA (for the shortest PEG units), m-PPEGMEMA (for the medium PEG units) and l-PPEGMEMA (for the longest PEG units), with s, m and l referring to short, medium and longer PEG side chains, respectively.

To determine the number average molecular weight ( $M_n$ ), weight average molecular weight ( $M_w$ ), and polydispersity index (PDI) of the copolymers, asymmetric flow field-flow

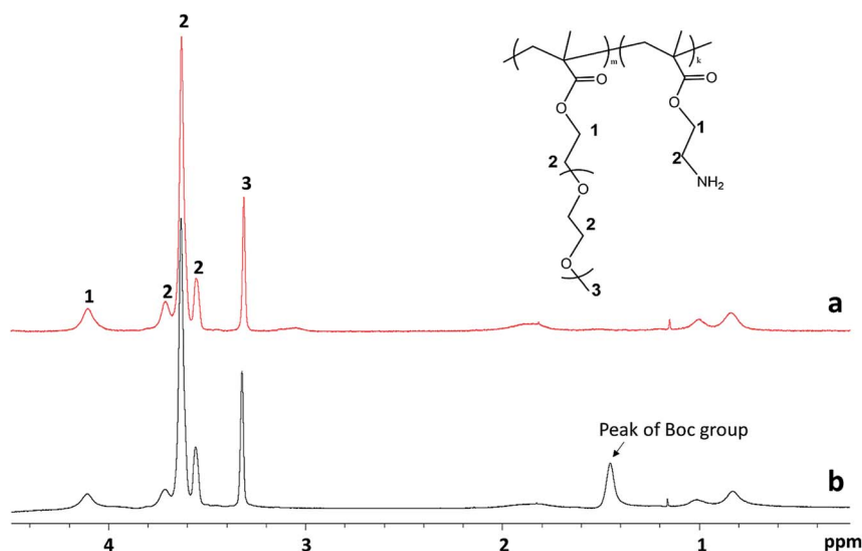


Fig. 1 NMR spectra for m-PPEGMEMA (a) after and (b) before deprotection to remove Boc groups.



Table 1 Properties of the synthesized P(PEGMEMA-stat-AMA) copolymers

	Polymer Composition <sup>a</sup>				Mn <sup>b</sup> (kDa)	PDI <sup>b</sup>
	P(EO) <sub>9</sub> MEMA	P(EO) <sub>4.5</sub> MEMA	DEGMEMA	AMA		
l-PPEGMEMA	124	0	0	41	35.9	2.66
m-PPEGMEMA	0	135	0	45	38.3	1.71
s-PPEGMEMA	0	29	115	48	38.2	1.43

<sup>a</sup> determined by <sup>1</sup>H NMR. <sup>b</sup> determined by AF4.

fractionation (AF4) was performed using a Wyatt Eclipse instrument with UV (Agilent 1230 infinity, Agilent), refractive index (Optilab rex, 633 nm, Wyatt) and multi-angle light scattering (MALS) (Dawn Heleos-II, 662 nm, Wyatt) detectors. A frit-inlet channel equipped with a regenerated cellulose membrane (MWCO 5 kDa, Millipore) and a W350 spacer was used as the separation channel. The samples were analyzed with a constant detector flow of 0.5 mL min<sup>-1</sup>, and a cross flow that decreased exponentially from 3 mL min<sup>-1</sup> to 0 mL min<sup>-1</sup> in 20 min. A 50 mM PBS buffer at pH 7.4 was prepared and filtered with a 0.1 μm membrane (Millipore) immediately before use as the eluent. The samples were dissolved in the PBS buffer with a concentration of 5 mg mL<sup>-1</sup>, and the injection volume was set to be 100 μL. The refractive index increment ( $dn/dc$ ) values for the copolymers were determined with an Optilab rex detector. For the molecular weight calculation,  $dn/dc$  values of 0.135, 0.120 and 0.118 were measured for s-PPEGMEMA, m-PPEGMEMA, and l-PPEGMEMA, respectively. Astra software (Wyatt, version 7.1.3.15) was utilized for data analysis and determination of the molecular weight from Debye plots.<sup>41</sup>

### 2.3. Quartz crystal microbalance with dissipation monitoring (QCM-D)

LbL assembly and the pH-responsiveness of PPEGMEMA/alginate multilayered films was monitored with QCM-D (Q-Sense E1, Biolin Scientific, Gothenburg, Sweden) using silica-coated sensors (QSX 335, Biolin Scientific). In a typical QCM-D experiment, an alternating voltage is applied to a quartz crystal sensor, which gives rise to oscillation of the sensor at its fundamental resonance frequency ( $F$ ). The resonance frequency is related to the mass of the oscillating sensor as well as any coupled mass. Additionally, the decay in the sensor oscillation is monitored upon repeated stopping of the drive generator output. Accordingly, the dissipation factor ( $D$ ) is determined as the ratio of the dissipated energy to the total stored energy.

The sensor was first aminated with APTES before the LbL assembly.<sup>42</sup> To do so, the sensor was rinsed with copious amounts of ethanol and water, dried, and then plasma-treated (PDC-32G plasma cleaner, Harrick Plasma) in water vapor with a constant pressure of 0.5 Torr for 1 min. Afterwards, the sensor was exposed to APTES and toluene vapor for 18 hours by placing it in a vacuumed desiccator with a 50% (v/v) APTES/Toluene solution. Hereafter, the sensor was rinsed with copious amounts of toluene and ethanol and dried with compressed air. The sensor was then immediately mounted

into the QCM-D module (QSense ellipsometry module, QELM 401, Biolin Scientific, Gothenburg, Sweden), and the measurement was started at 23 °C under a 150 μL min<sup>-1</sup> flow rate of citric acid/phosphate buffer with pH 3. After obtaining a stable baseline for all the harmonics, alginate and PEGMEMA solutions were alternately loaded and rinsed with buffer to obtain 7 bilayers in total. Glutaraldehyde solution (1%, w/w) was then loaded for 2 hours to cross-link the multilayered film, after which the chamber was rinsed with pH 3 buffer to remove the unbounded glutaraldehyde. To test the pH stability, 30 mM NaCl solutions with pH 3 and 5.6 were loaded, following which three consecutive pH cycles were conducted. Finally, the pH-responsiveness of the film was examined through titration of NaCl solutions with pH values ranging between 2 to 9. In all cases, the pH was adjusted with HCl/NaOH and NaCl was the only added electrolyte.

For thin, uniform and rigid films, the frequency shifts show a linear dependence to the adsorbed mass per unit area according to the Sauerbrey equation.<sup>43</sup> The Sauerbrey equation, however, only provides a valid estimation of the ratio if dissipation and normalized frequency shifts ( $(\Delta D_n)/(-\Delta f_n/n)$ ) are significantly smaller than  $4 \times 10^{-7}$  Hz<sup>-1</sup>; otherwise, the adsorbed mass will be underestimated.<sup>44</sup> For soft and highly hydrated polymeric films, as in the present case, the measured shifts (for different harmonics) in the resonance frequency and dissipation factor can be related to the thickness and viscoelastic properties of the film through modeling.<sup>44</sup> Here, the viscoelastic Voigt model, where the adhered layer is represented by a film of uniform thickness and density with distinct viscous and elastic components can provide a better estimation of the film properties. The measured frequency and dissipation shifts are then related to the properties of the film and medium by:<sup>45</sup>

$$\frac{\Delta f}{f} = -\frac{d_f \rho_f}{d_q \rho_q} \left( 1 - \eta_0 \rho_0 \times \frac{(\eta_f / \rho_f) \omega^2}{(\mu_f^2 + \omega^2 \eta_f^2)} \right)$$

$$\Delta D = \frac{d_f}{d_q \rho_q} \left( \eta_0 \rho_0 \times \frac{\mu_f \omega}{\mu_f^2 + \omega^2 \eta_f^2} \right)$$

where  $\omega$  is the angular frequency of oscillation,  $\eta_0$  and  $\rho_0$  are the viscosity and density of the medium, respectively, and  $d_q$  and  $\rho_q$  are the thickness and density of the quartz crystal resonator, respectively. The density and viscosity of the medium together with the density of the hydrated film are estimated and treated



as fixed parameters to avoid overparameterization. The thickness ( $d_f$ ), viscosity ( $\eta_f$ ) and shear modulus ( $\mu_f$ ) of the polymeric film are thus obtained from Voigt modeling. The instrument software (Dfind, Biolin Scientific) was employed for analysis of the data. The density of the hydrated film ( $\rho_f$ ) was estimated to be  $1050 \text{ kg m}^{-3}$  (see ESI, Section S2†). The density and viscosity of water at  $23^\circ\text{C}$  (from the software library) was used for the medium.

#### 2.4. Spectroscopic ellipsometry

As mentioned above, the QCM-D studies of the LbL assembly and tests of the pH responsiveness of the films were conducted in an ellipsometry module allowing for simultaneous QCM-D and ellipsometry measurement. Spectroscopic ellipsometry (M2000, JA Woollam Co., USA) is an optical method in which changes in the polarization state of light upon interaction with a sample are measured. Two parameters, *i.e.*,  $\psi$  (amplitude ratio) and  $\Delta$  (phase shift) are used to quantify the polarization change. The measured parameters correlate with the optical properties and thickness of the sample components:<sup>46</sup>

$$\tan(\psi)\exp(i\Delta) = \frac{r_p}{r_s} = \rho(\theta_0, \lambda, n_s, n_{\text{amb}}, n_j, k_j, d_j)$$

Here, the subscripts p and s refer to the parallel and perpendicular directions with respect to the plane of incidence, respectively, and  $r_p$  and  $r_s$  represent the reflection coefficients. The  $r_p$  to  $r_s$  ratio is a complex function of the angle of incidence ( $\theta_0$ ), wavelength ( $\lambda$ ), optical functions of the substrate ( $n_s$ ) and the ambient ( $n_{\text{amb}}$ ), as well as the optical functions ( $n_j, k_j$ ) and thickness ( $d_j$ ) of the film.

The QSense ellipsometry module (QELM 401, Biolin Scientific, Gothenburg, Sweden) allows *in situ* ellipsometry measurements on the QCM-D sensor through two optical windows. Accordingly,  $\psi$  and  $\Delta$  spectra were collected (wavelength range: 250–1000 nm, angle of incidence of  $70^\circ$ ) before adsorption of the first layer (bare sensor data) and at the end of each rinsing step. The instrument software (CompleteEASE, JA Woollam Co., USA) was employed to model the  $\psi$  and  $\Delta$  data. The optical model employed was composed of three layers representing a uniform hydrated polymer film, a silica coating, and a thick optically opaque titanium substrate. First, the bare sensor was modeled as a pseudosubstrate, and the fitted parameters were fixed in the model (see ESI, Section S3†). Considering the low ionic strength of the buffer solutions, the optical data for pure water at  $23^\circ\text{C}$  from the software library were used for the medium.<sup>47</sup> The multilayered film was regarded as a transparent and homogeneous layer with no adsorption ( $k = 0$ ); then, the refractive index ( $n$ ) was described by the Cauchy equation:<sup>48</sup>

$$n(\lambda) = A + \frac{B}{\lambda^2}$$

In the first modeling approach, the hydrated multilayered film was considered as a single-component film characterized by a thickness and two effective optical constants ( $A_{\text{eff}}, B_{\text{eff}}$ ). In

the second modeling approach, the multilayered film was considered as a two-component layer consisting of a dry polymer ( $A = 1.5$  and  $B = 0.005$ )<sup>49,50</sup> and water. In addition to film thickness, the volume fraction of water ( $f_w$ ) was then estimated according to the Bruggeman Effective Medium Approximation (BEMA):<sup>51</sup>

$$0 = f_w \frac{n_w^2 - n^2}{n_w^2 + 2n^2} + (1 - f_w) \frac{n_p^2 - n^2}{n_p^2 + 2n^2}$$

where  $n$  is the refractive index of the hydrated film,  $f_w$  is the volume fraction of water, and  $n_w$  and  $n_p$  are the refractive indices of the water and dry polymer, respectively. Additional fitting options including surface roughness, grading, and thickness nonuniformity were tested, but only the latter was found to significantly improve the fitting quality. To assess the quality of modeling, the MSE value, uniqueness of the thickness, correlation between the fitted parameters and correctness of the refractive indices were checked (see ESI, Section S3†).

## 3. Results and discussion

### 3.1. P(PEGMEMA-*stat*-AMA) cationic copolymers for multilayer assembly

As described in the experimental section, three PEGMEMA-based cationic copolymers, with varying average PEG side chain lengths were prepared by ATRP, as shown in Scheme 1; a summary of the properties of the synthesized copolymers is provided in Table 1. The conversion of each polymerization was determined from  $^1\text{H}$  NMR data, based on which the actual degree of polymerization of each monomer was obtained. Overall, the degree of polymerization for all three copolymers was comparable, with a slight variation observed due to a minor difference in the conversion. The number average molecular weight and polydispersity of the copolymers was characterized by AF4 measurement (see Table 1). Here, it is seen that the s-PPEGMEMA has a relatively narrow molecular weight distribution, while the PDI value was observed to increase as the average PEG chain length increased. The latter observation indicates less control of the polymerization for longer PEG side chains.

### 3.2. LbL assembly

Fig. 2 demonstrates the QCM-D data in terms of the shifts in the oscillation frequency ( $F$ ) and dissipation factor ( $D$ ) resulting from LbL assembly of PPEGMEMA/alginate multilayered films. The negative shift in oscillation frequency correlates to the effective mass (polymer film plus hydrodynamic water content) coupled to the sensor. The dissipation factor, on the other hand, represents the ability of the adhered film to dissipate the oscillatory energy; consequently, it is considered as a semi-quantitative measure of the film conformation and viscoelasticity. Accordingly, a soft and hydrated film is characterized by a relatively large positive shift in the dissipation factor, while a rigid and compact film produces relatively small shifts in the dissipation factor.

According to Fig. 2, the measured dissipation shifts for 14 layers (7 bilayers) for s-PPEGMEMA, m-PPEGMEMA and l-



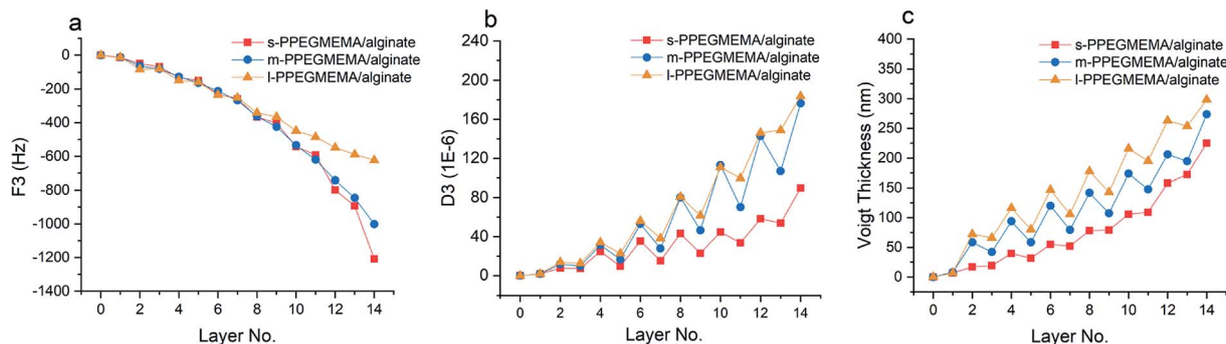


Fig. 2 QCM-D frequency shift (a), dissipation shift (b) and the corresponding Voigt thickness (c) for LbL assembly of s-PPEGMEMA (red square), m-PPEGMEMA (blue circle) and l-PPEGMEMA (yellow triangle) with alginate in pH 3 citric acid/phosphate buffer.

PPEGMEMA samples are approximately  $90 \times 10^{-6}$ ,  $180 \times 10^{-6}$  and  $183 \times 10^{-6}$ , respectively. Such large dissipation values suggest that the multilayered films are all swollen and highly hydrated. On the other hand, the frequency shifts for 14 layers for the s-PPEGMEMA, m-PPEGMEMA, and l-PPEGMEMA samples are around  $-1200$ ,  $-1000$  and  $-620$  Hz, respectively. Accordingly, increasing the average length of the PEG side chains leads to a relatively larger dissipation shift (more viscoelastic film) and a relatively smaller frequency shift (smaller coupled mass). The model was then fitted to the frequency and dissipation data to estimate the Voigt thickness of each PPEGMEMA/alginate multilayered film as a function of the deposition number (see Fig. 2c and Table 2).

According to Fig. 2c, it appears that the LbL assembly of s-PPEGMEMA/alginate multilayered films follows a nonlinear growth mechanism. However, the multilayer growth seems to be inclined towards a linear growth mechanism when increasing the length of the PEG side chains in the copolymers. It has been suggested that the diffusion of polyelectrolyte chains “in and out” of the film during the LbL buildup can give rise to nonlinear growth.<sup>1,52–54</sup> Accordingly, one could speculate that the bulkiness of the PEG side chains can hinder segmental diffusion within the film, and, as a consequence, promote a linear growth mechanism.

In addition, while the frequency data show a monotonic gain, the dissipation data demonstrate a zigzag trend. Accordingly, adsorption of PPEGMEMA copolymers gives rise to a relatively large increment in dissipation, whereas the deposition of alginate produces a considerable decrement in dissipation. This can be interpreted based on the conformation of

the interfacial top layer that has been previously observed.<sup>55</sup> When the PPEGMEMA copolymer is the top layer, a highly swollen and hydrated conformation is expected for the interfacial layer, which thus explains the large dissipation gain. On the other hand, adsorption of alginate seems to promote partial collapse of the sublayer, subsequently producing a relatively rigid and less hydrated interfacial layer.

Spectroscopic ellipsometry data can provide further information for the structural features of the multilayered films. Fig. 3 illustrates the  $\psi$  and  $\Delta$  spectra for the bare sensor as well as polymer-coated sensor after each deposition step. For s-PPEGMEMA, the LbL deposition is characterized by marked spectral oscillations in both the  $\psi$  and  $\Delta$  spectra. These changes are more pronounced specifically after the 10<sup>th</sup> deposited layer, where a secondary peak evolves in the UV range of the  $\psi$  spectra and a significant shift in the  $\Delta$  spectra is found. This observation is in line with the QCM-D data, suggesting a non-monotonic growth in thickness. For m-PPEGMEMA, the changes in the  $\psi$  and  $\Delta$  spectra are relatively less pronounced, which can imply a relatively smaller thickness and (or) higher water content in the film. For l-PPEGMEMA, the evolution of the  $\psi$  and  $\Delta$  spectra is even less significant as no trace of a secondary peak is found in the  $\psi$  spectra and a relatively smaller shift in the  $\Delta$  spectra is demonstrated, which together suggests a thinner and (or) more hydrated layer compared to the layers obtained with s-PPEGMEMA and m-PPEGMEMA. In addition, the shifts in  $\psi$  and  $\Delta$  following LbL deposition indicate a monotonic trend, which suggests a linear growth mechanism.

Table 2 Summary of the QCM-D and ellipsometry data obtained for the PPEGMEMA/alginate multilayered film

polymer	Voigt Thickness (nm)	Cauchy Thickness (nm)	BEMA Thickness <sup>a</sup> (nm)	BEMA Water Content <sup>a</sup> (%)	Dry Thickness <sup>b</sup> (nm)
s-PPEGMEMA	225.1	200.7 ± 0.3	198.9 ± 0.3	77.3 ± 0.1	48.9 ± 0.1
m-PPEGMEMA	273.8	180.6 ± 0.6	177.1 ± 0.6	83.3 ± 0.1	29.4 ± 0.1
l-PPEGMEMA	297.6	163.3 ± 2.7	156.1 ± 2.6	91.5 ± 0.2	17.6 ± 0.1

<sup>a</sup> Fitted with the BEMA method, Bruggeman model. <sup>b</sup> Fitted with the Cauchy method, with mean A, B and C parameters for PPEGMEMA and alginate Cauchy parameters.



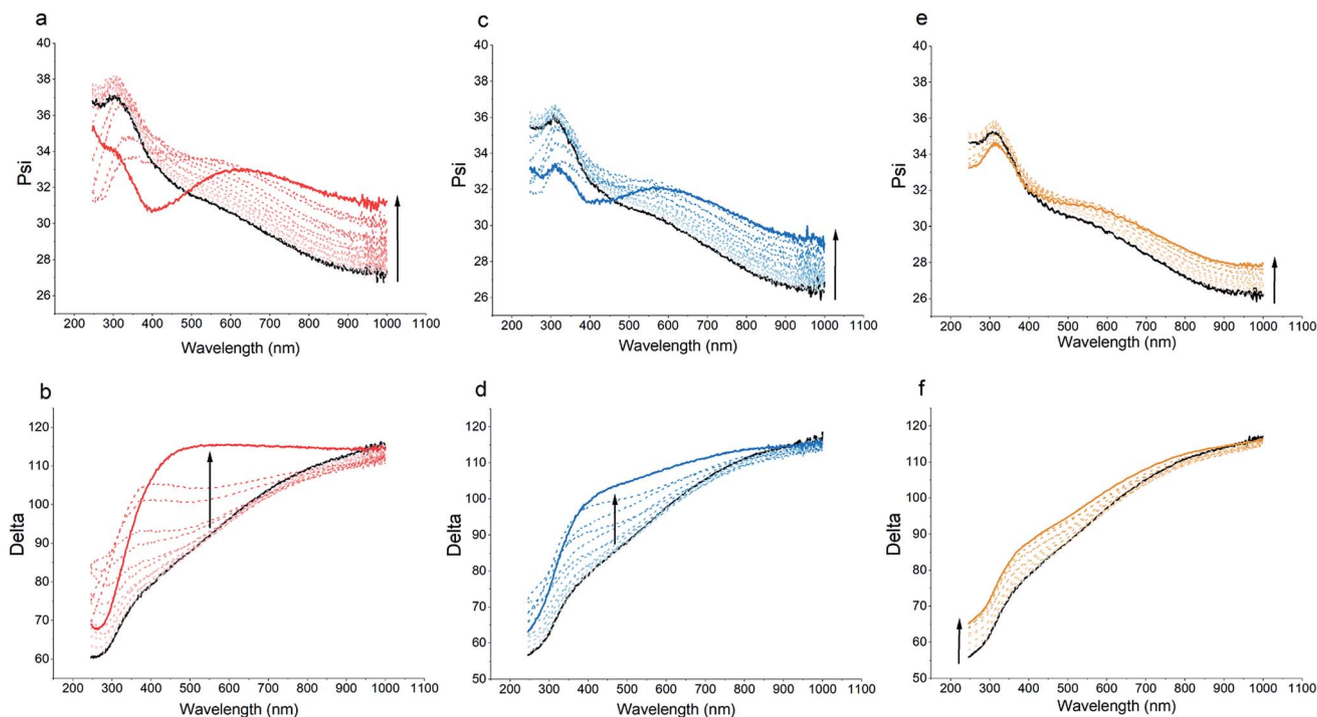


Fig. 3 Ellipsometric Psi (top row) and Delta (bottom row) for LbL assembly of s-PPEGMEMA (a and b), m-PPEGMEMA (c and d) and l-PPEGMEMA (e and f) with alginate in pH 3 citric acid/phosphate buffer (the black solid curve represents the bare sensor and the arrows hint at LbL deposition).

To obtain more quantitative information for the structure of the films, the spectroscopic ellipsometry data were examined using a single-component Cauchy and a two-component BEMA model. According to both modeling approaches, increasing the average length of the PEG side chains leads to a decrement in the total hydrated film thickness from approximately 200 nm for the s-PPEGMEMA/alginate multilayered film to approximately 160 nm for the l-PPEGMEMA/alginate multilayered film, while the water content within the film increases roughly from 77% for the s-PPEGMEMA/alginate multilayered film to 91% for the l-PPEGMEMA/alginate multilayered film. This difference in the water content of the films is reflected in the refractive index dispersion of the hydrated layers (see Fig. 4). As a general observation, the refractive indices for all three layers are closer to that of water than of the dry polymer, suggesting a large amount of water in the films. In addition, increasing the average PEG side chain length gives rise to a decrement in the refractive index, suggesting that the water content is relatively large. The increase in water content obtained upon increasing the average PEG side chain length can also explain the larger QCM-D dissipation shift for this layer. In addition, the dry thicknesses of the s-PPEGMEMA, m-PPEGMEMA, and l-PPEGMEMA samples are found to be approximately 49, 29 and 18 nm, respectively. Accordingly, it can be concluded that increasing the average length of the PEG side chains decreases the amount of adsorbed polymer chains, which can be attributed to a larger steric hindrance for LbL deposition.

It is noteworthy that a discrepancy in film thickness was observed between Voigt modeling and the optical thickness (Table 2). We attribute this discrepancy to the inhomogeneous

nature of the PPEGMEMA/alginate multilayered film. In particular, while the inner layers of the obtained film are relatively impact, the outer PPEGMEMA layer is highly swollen and hydrated. The sparsely distributed PPEGMEMA chains in the outermost layer of the film significantly affect the dissipation value and add to the overestimation of the Voigt thickness. This

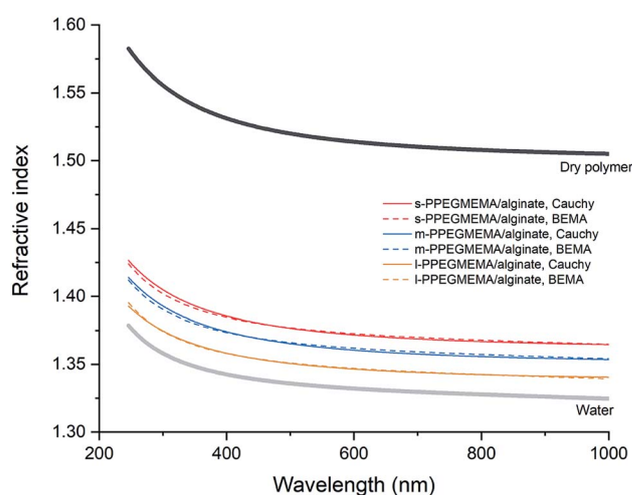


Fig. 4 Refractive indices obtained from single-component Cauchy (solid line) and two-component BEMA (dashed line) models for 3 bilayers of s-PPEGMEMA/alginate (red), m-PPEGMEMA/alginate (blue) and l-PPEGMEMA/alginate (yellow) multilayered films. Refractive indices for water (bold gray) and dry polymer (bold black) are provided for comparison.



effect is much less significant to ellipsometry modeling, where an outer layer with low polymer density will contribute with a low weight to the optical thickness.

### 3.3. Stabilization of the PPEGMEMA/alginate film by cross-linking

As electrostatic interactions are the driving force for the multi-layer buildup, the pH at which the multilayered film is fabricated should roughly correspond to a state of charge neutrality. Accordingly, a subsequent change in pH will create a charge imbalance that will cause excessive swelling and disintegration of the film (see ESI, Fig. S17†). Chemical cross-linking, on the other hand, is an effective method to enhance the pH-stability of polyelectrolyte multilayered films. Glutaraldehyde, as a common cross-linker for proteins and polysaccharides, can form a dynamic covalent bond between free amine groups on the PPEGMEMA copolymer through the Schiff base reaction.<sup>56–58</sup> Herein, we selected the m-PPEGMEMA/alginate multilayered film to investigate the effect of cross-linking and pH-stability.

Fig. 5 demonstrates the structural changes in the multilayered film resulting from the cross-linking process. With respect to the QCM-D data (Fig. 5a), a significant decrement in the dissipation of around  $-80 \times 10^{-6}$  accompanied by a small frequency shift of around  $-70$  Hz, is observed. While the change in frequency can be considered insignificant compared to the total frequency shift during the layer buildup (see Fig. 2a), the relatively large decrease in dissipation suggests structural collapse and enhanced rigidity of the film after cross-linking.

According to the ellipsometry data, chemical cross-linking is accompanied by a decrease in the optical thickness from 237 nm to 210 nm, confirming the film shrinkage, as inferred by the dissipation data. In the same line, a decrease in water content from 85% to 81% is demonstrated. The film shrinkage could be attributed to the consumption of a portion of the amine groups, and, hence, charge regulation within the film, as well as the conformational immobilization caused by the chemical cross-links.

### 3.4. pH-responsive behavior of the m-PPEGMEMA/alginate multilayered film

Fig. 6 demonstrates the stability of the multilayered film upon three consecutive pH cycles between 3 and 5.6. According to the QCM-D data (Fig. 6a), increasing the pH from 3 to 5.6 leads to a gain in dissipation and a decline in frequency, which both imply swelling and hydration of the polymer film. By lowering the pH back to 3, the frequency and dissipation shifts demonstrate conformational collapse and dehydration of the film. During the three pH cycles, no significant layer disintegration and mass loss was observed, indicating enhanced pH stability of the film after cross-linking. It is, however, worth mentioning that a slight structural hysteresis (irreversible frequency and dissipation shifts) is found after each pH cycle, which could be attributed to conformational restructuring within the film that can also cause irreversible ionization of amine and carboxyl groups.<sup>59–61</sup>

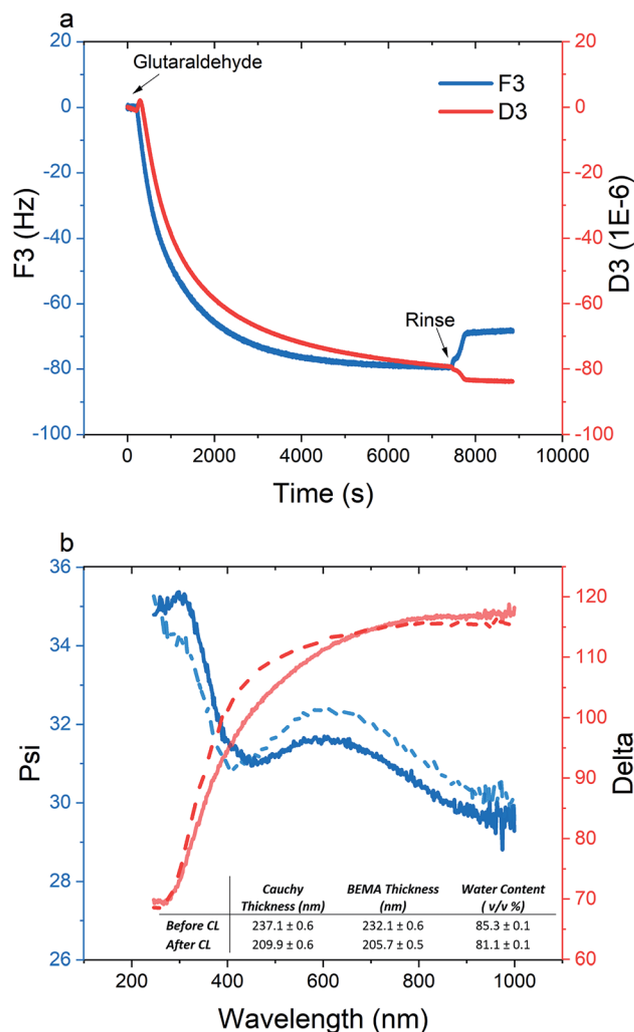


Fig. 5 (a) Frequency (blue) and dissipation (red) shifts for the cross-linking step of a m-PPEGMEMA/alginate multilayered film with 1% glutaraldehyde; (b) Psi (blue) and Delta (red) before (dashed line) and after (solid line) cross-linking. The inset shows the modeling result obtained from the Cauchy and BEMA methods.

Ellipsometry results (Fig. 6b) represent the same effect as concluded from the QCM-D data. During pH cycling, the Cauchy thickness exhibited a zig-zag trend in agreement with the frequency and dissipation shifts. Accordingly, the optical thickness obtained with the BEMA model fluctuates between approximately 210 nm at pH 3 and 225 nm at pH 6, while the estimated water content at the same time also varies systematically.

To further investigate the pH-responsiveness of the PPEGMEMA/alginate multilayered film, a pH titration was performed (see Fig. 7). For the QCM-D data presented in Fig. 7a, it can be seen that increasing the pH from 2 to 4 results in a relatively large dissipation shift of around  $-50 \times 10^{-6}$ , suggesting shrinkage of the film. In contrast, further increasing the pH from 4 to 9 is characterized by a relatively large gain in the dissipation of approximately  $140 \times 10^{-6}$ , indicating



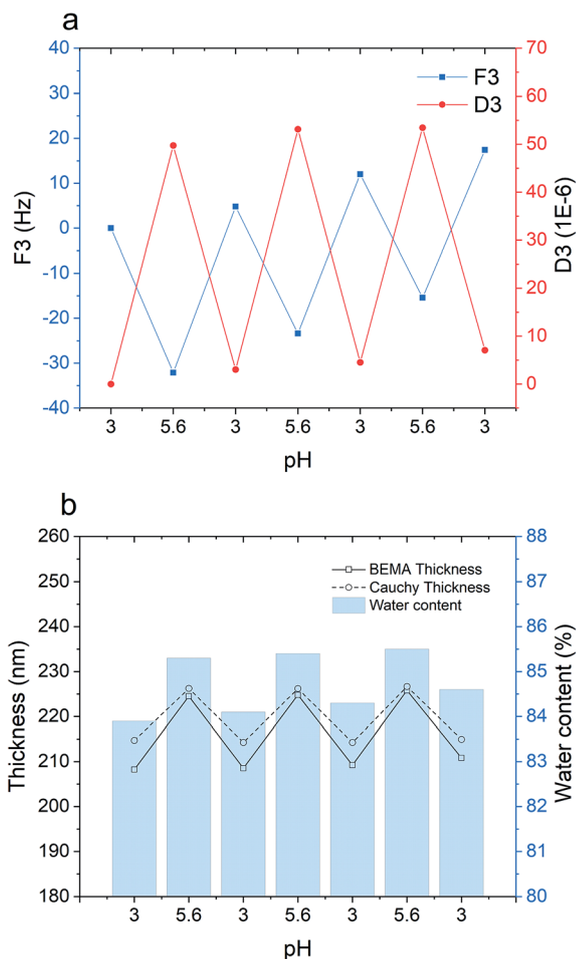


Fig. 6 m-PPEGMEMA/alginate multilayered film stability as a function of pH during repeated changes between pH 3 and 5.6: (a) frequency and dissipation changes monitored with QCM-D; (b) Cauchy (dashed line, circle) and BEMA thicknesses (solid line, square) as well as water content (violet bar) obtained *via* the BEMA model from ellipsometry.

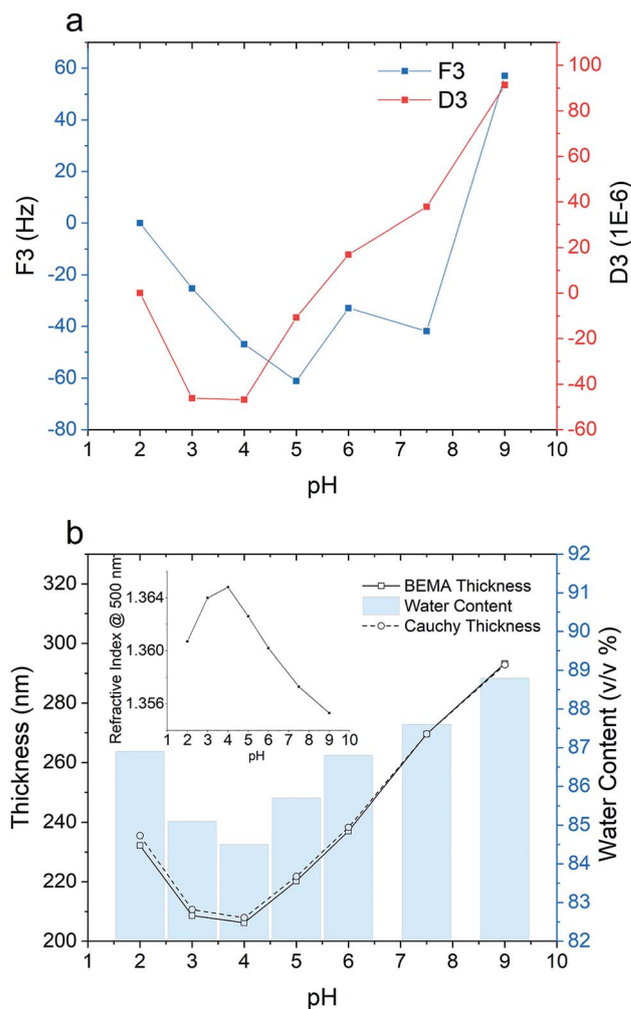


Fig. 7 pH responsiveness of a m-PPEGMEMA/alginate multilayered film; (a) frequency and dissipation changes obtained from QCM-D; (b) Cauchy thickness (circle, dashed line), BEMA thickness (square, solid line) and water content (blue bar). The inset shows how the refractive index changes with the pH variation.

a reswelling of the film. For the frequency shift, a similar variation is observed although the trend is less clear.

In the ellipsometry data (see Fig. 7b), increasing the pH from 2 to 4 is associated with a decline in the BEMA thickness from 235 nm to 208 nm, while further increasing the pH results in a gain in thickness from 208 to 290 nm. The estimated water content followed the same trend as the film thickness, *i.e.*, decreasing from 87 to 85% when the pH is changed from 2 to 4 and increasing from 85 to 89% when the pH is changed from 4 to 9.

The swollen-collapsed-swollen conformational change of the multilayered film when the pH is first changed from 2 to 4 and then subsequently changed to 9 can be explained based on pH-dependent charge regulation within the multilayered film. The collapsed state in the pH range of 3–4 could be attributed to the charge neutral state within the multilayered film. Under such conditions, the number of charged amine and carboxyl groups within the film are roughly comparable. Decreasing the pH to 2 is associated with protonation of the amine groups on the

PPEGMEMA copolymer and the carboxylate groups on the alginate, creating a net positive charge within the film. This leads to the accumulation of counterions ( $\text{Cl}^-$ ) within the film that can produce an osmotic pressure difference, and, hence, swelling of the film. On the other hand, increasing the pH to 9 is accompanied by deprotonation of the carboxyl groups and amine groups, which produces a net negative charge in the film. Accumulation of  $\text{Na}^+$  counterions then results in an osmotic swelling. Thus, the swollen-collapsed-swollen conformational change is due to the cross-linked PPEGMEMA/alginate multilayered film exhibiting cationic, zwitterionic and anionic states as a function of pH. To this end, we note that the charging state of the multilayered film, as well as the possibility to change between different charging states, is of high importance for the film functionality and for possible applications. While both the hydration level and the sign of the surface potential will be decisive for the films ability to either attract or repel specific biomolecules or cells,<sup>62</sup> the responsive nature of the film can be



used for control release of encapsulated drugs,<sup>63</sup> for response membranes<sup>64</sup> or in sensor technologies.<sup>65</sup> For these three different states, it must, however, be considered that the swelling capacity of the PPEGMEMA/alginate multilayered film is governed by the balance between the osmotic pressure of the solution and the entropic penalty of stretching the polymer chains between the cross-linking points.

## 4. Conclusion

In this work, three PPEGMEMA-based cationic copolymers with different average PEG side chain lengths were paired with alginate *via* electrostatic LbL assembly. *In situ* QCM-D and ellipsometry measurements were used to follow the multilayer buildup process for the three different systems, and the result shows that the thickness and structure of the multilayered film is closely correlated to the average PEG side chain length in the PPEGMEMA-based cationic copolymers. Specifically, as the average PEG side chain length increases, two main observations are noted. First, it is found that the multilayer builds up less effectively with a reduced increase in added mass during the alternating deposition of polymers. Second, the film is found to become more dissipative and exhibit a higher water content. Both of these effects are interpreted as being directly related to the average PEG side chain length, which, due to the high hydrophilicity of the PEG units, leads to a highly hydrated layer and to steric hindrance, which limits the amount of polymer adsorbed in each deposition step.

After buildup, the obtained PPEGMEMA/alginate multilayered films were cross-linked with glutaraldehyde to exhibit pH responsiveness without disintegration of the layer due to charge imbalance in the layer. It was further demonstrated how the PPEGMEMA/alginate multilayered films can be in a cationic, zwitterionic or anionic state depending on the pH, and how the transitions between these states lead to structural changes in the layer.

## Conflicts of interest

There are no conflicts to declare.

## Acknowledgements

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