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Imprinting of Metal Receptors into Multilayer Polyelectrolyte Films: Fabrication and Applications in Marine Antifouling

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Abstract

Polymeric films constructed using the layer-by-layer (LbL) fabrication process were employed as a platform for metal ion immobilization and applied as a marine antifouling coating. The novel Cu^{2+} ion imprinting process described is based on the use of metal ion templates and LbL multilayer covalent cross-linking. Custom synthesized, peptide mimicking polycations composed of histidine grafted poly(allylamine) (PAH) to bind metal ions, and methyl ester containing polyanions for convenient cross-linking were used in the fabrication process. Two methods of LbL film formation have been investigated using alternate polyelectrolyte deposition namely non-imprinted LbL_A, and imprinted LbL_B. Both LbL films were cross linked at mild temperature to yield covalent bridging of the layers for improved stability in sea water environment. A comparative study of the non- imprinted LbL_A films and imprinted LbL_B films for Cu^{2+} ion binding capacity, leaching rate and stability of the films was performed. The results reveal that the imprinted films possess enhanced affinity to retain metal ions due to preorganization of imidazole bearing histidine receptors. As a result the binding capacity of the films for Cu^{2+} could be improved by seven fold. Antifouling properties of the resulting materials in marine environment have been demonstrated against settlement of barnacle larvae, indicating that controlled release of Cu ions was achieved.

Keywords: Antifouling; polyelectrolyte; histidine; imidazole; peptide mimicking polymer; ion imprinting, metal selective materials, electrostatic layer by layer;

Introduction

The ongoing quest for complex, advanced materials systems necessitates the development of various thin polymeric films for applications in a variety of fields including membrane technology,¹⁻³ microelectronics,⁴ optical sensing,⁵ and biomedical applications.⁶⁻⁷ Particular attention is directed toward the management of the molecular payloads within the film e.g. selective loading, and controlled delivery or multiplexed release.⁸⁻⁹ The laver-by-laver (LbL) electrostatic fabrication process utilizing alternate deposition of positively and negatively charged polyions on a substrate¹⁰ is a simple yet versatile method for preparation of thin polyelectrolyte multilayer coatings.¹¹⁻¹⁴ Polymeric films with enhanced affinity toward metal ions are needed for applications in ion capture and separation,^{1-3, 15} catalysis,¹⁶⁻¹⁷ protein binding, in metal affinity chromatography¹⁸ and antimicrobial materials.¹⁹ The selective adsorption and separation of metal ions from aqueous waste also received an increasing consideration in environmental protection in recent years.²⁰ For example, numerous adsorbents have been developed for the removal of copper ions from industrial wastewater and in other applications.²¹⁻ ²² However, these adsorbents still suffer from drawbacks such as limited selectivity and low reusability.Molecular-imprinting techniques (MIPs) have become acclaimed in many fields, such as chemosensor fabrication, molecular separation, and catalysis and drug delivery.²³⁻²⁷ Ion imprinted polymers can be considered as a sub category within MIPs^{22, 27} and refer to materials with improved affinity toward metals by pre-organization of metal receptors within polymeric matrix in presence metal ion templates. This process is typically achieved by the reduction of receptor mobility through various ways such as bulk polymerization, precipitation

polymerization, and suspension polymerization. Since the receptor sites are embedded deeply into the polymeric matrix, these imprinting techniques often produce materials exhibiting high selectivity but low rebinding capacity and poor site accessibility to target ions.²⁸ In order to conquer these drawbacks, thin film surface-ion-imprinted polymers were proposed as promising solution. Films obtained by surface ion-imprinted techniques exhibits good characteristics, such as complete removal of templates and fast adsorption kinetics. ²⁹⁻³³Small molecules and ions have been loaded into non-cross-linked LbL fabricated thin electrostatic film systems using both the traditional solution LbL method^{16, 34} as well as spray deposition approaches.³⁵ The films obtained show not only high affinity to metals but also possess reversible loading/unloading capability.³⁶ Enhanced selectivity of such films toward the ions loaded is attributed to reduced mobility of polyions upon deposition in the electrostatic multilayer.³⁷ This affinity however can potentially be further increased by the reduction of receptor mobility, for example through covalent cross-linking of the LbL layers. The ion imprinting through covalent crosslinking of multilayer systems in presence of metal templates was not presented until now. Studies, which employed cross-linking of LbL systems in presence of ions have focused only on moderately coordinating polyethylenimine/polyacrylic acid (PEI/PAA) couple. Enhanced affinity of the films toward specific metals upon cross-linking has not been demonstrated, nor discussed, in the corresponding reports.³⁸⁻³⁹ Finally, direct imprinting of metals into polyelectrolyte film by immobilization of metal templated receptors was demonstrated only for a single PEI layer (non LbL) using an epichlorohydrin crosslinker.⁴⁰⁻⁴¹ However a single polymer layer is inferior compared to LbL coatings in terms of substrate coverage and number of receptors available. Reported systems used also rather poorly coordinating PEI receptors.

Protein complexes with metal ions play an important role in various biological processes.⁴² A common structural feature of such protein-metal assemblies⁴³ are peptide motifs typically arranged in a special sequence of residues to ensure the strong interactions between the metal ions and donor atoms from the peptide backbones or functional side groups.^{29, 44} These motifs are frequently bearing residues such as histidine (His), since nitrogen of the imidazole moiety is known for its high affinity to metals.⁴⁵⁻⁴⁶ It is used as a metal coordinating component of polymeric systems and it can be grafted on various polymeric backbones mimicking peptide

architecture.⁴⁷⁻⁴⁹ Hence in this work we shall describe a system in which this specific coordinating capacity of histidine is utilized.

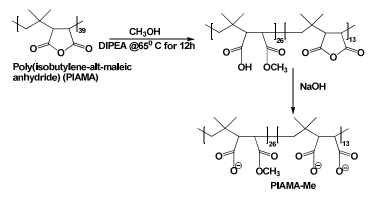
Marine fouling has been a serious problem for maritime technologies, from ship hull protection to off-shore stationary structures and high-value added underwater sensing and communications.⁵⁰⁻⁵¹ Marine fouling has been controlled traditionally through the use of antifouling paints with toxic constituents or biocides.⁵² Before its ban tributyltin (TBT) was the active agent in antifouling paints and used extensively in the maritime sector for over thirty years.⁵³ As an alternative, copper based coatings are ten-twelve folds less harmful than those containing TBT,⁵⁴⁻⁵⁵ however still remain problematic and are criticized for excessive metal pollution. Copper is used as an antifouling agent with a leaching rate on the order of 10-15 µg/cm² per day. Hence a surface of one square meter can be kept clean of biological growth by annual leaching of approximately 30 grams of copper.⁵⁶ Those levels are suspected as toxic to aquatic organisms, through accumulation in filter feeders, such as mussels, and to damage larval stages of aquatic invertebrates and fish species.⁵⁷ Designing and fabricating new, effective and environmentally friendly coating systems as alternatives to TBT-based antifouling paints poses an important challenge for the scientific community. The ideal antifouling coating with embedded metal ions would prevent marine growth as well as maintain a long performance life while it would fulfill the expectations of the increasingly strict environmental regulations.

In this work we investigate the concept of improving of thin LbL polymeric film affinity to Cu^{2+} ions by reducing mobility of metal receptors in presence of guest molecules. This concept is typically referred to as metal ion imprinting.²² Utilizing properties of the polymer grafted imidazole bearing histidine receptor, we synthesized a peptide mimicking material with high affinity to copper. Two alternative approaches for the fabrication of metal attracting films were compared. In the first imprinting approach, thin LbL films composed of custom synthesized histidine grafted polyallyamine were cross-linked in presence of Cu^{2+} ions to immobilize receptors in a conformation suitable for the metal ion coordination. In the second non-imprinting approach, the films were cross-linked without the presence of ions to provide a reference material. The structure and properties of both films are extensively characterized. Finally, antifouling applications of the copper ion selective LbL films are demonstrated.

Experimental section

Materials

Poly(isobutylene-alt-maleic anhydride) (PIAMA, Mw: 6 kDa), poly(allylamine hydrochloride) (PAH, Mw: 58 kDa), L-histidine methyl ester dihydrochloride, 3-aminopropyltrimethoxysilane (APTMS), N,N-diisopropylethylamine (DIPEA), copper(II) nitrate trihydrate (Cu(NO₃)₂x3H₂O), sodium chloride, sea salts and sodium hydroxide (all from Sigma Aldrich), N,N-dimethylformamide (DMF), dimethylsulfoxide (DMSO), toluene, methanol, acetone and isopropanol (all from Tedia) were used directly as received without further purification. Dialysis membrane tubing (MWCO: 3.5 kD) was received from Fisher Scientific. Silicon wafers (Latech Scientific Supply Pte. Ltd) were 0.6 mm thick, one side polished and with a natural silicon dioxide layer. QSX 303 Silicon dioxide 50 nm quartz crystal microbalance (QCM) chips were obtained from Analytical Technologies Pte Ltd. Deionized (DI) water with 18 M Ω cm⁻¹ resistivity was obtained from a Millipore Nanopure system.



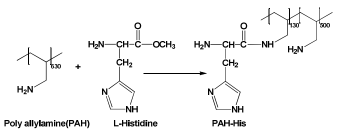
Scheme 1. The synthesis of PIAMA-Me.

Synthesis

Synthesis of PIAMA-Ester (PIAMA-Me): PIAMA-Me synthesis was achieved following a modification of the method described previously (Scheme 1).⁵⁸ To the solution of poly(isobutylene-*alt*-maleic anhydride) (1.0 g, 6.5 mmol) in 300 mL of methanol, DIPEA (1.2 mL, 6.9 mmol) was added and the mixture was stirred for 16 h at 50 °C. After evaporation of methanol and DIPEA, the material was suspended in water using a small excess of NaOH with respect to the carboxylic groups in the polymer backbone. The polymer solution was dialyzed

against 0.01 M NaOH and subsequently against pure water for a few days. The purified polymer aqueous solution was concentrated by rotary evaporator and finally freeze dried to yield the solid polymer. To properly identify the composition of the polymers obtained, NMR spectra were compared with NMR of poly (isobutylene-*alt*-maleic anhydride) opened by a treatment with a stoichiometric amount of NaOH to carboxyl groups. NMR calculated Mn: 8 kDa. ¹H NMR integrated for a single repeating unit: (D₂O) $\delta_{\rm H}$: 0.5-1.36 ppm (6 H, m), 3.67 ppm (2 H, s). IR: 1860 cm⁻¹, 1780 cm⁻¹, 1720 cm⁻¹, 1580 cm⁻¹.

Synthesis of PAH-Histidine(PAH-His): To a solution of L-histidine methyl ester (2.63 g, 11 mmol) in 20 mL of DI water, 1 g of PAH (7.4 mmol of the repeating units) was added in small portions (Scheme 2). The solution was stirred for 1 h at room temperature and small portions of 5M NaOH were added until the solution pH reached 10. The solution was freeze dried for 72 h. 2 mL of DMSO was added to the freeze dried polymer-histidine mixture and kept under vacuum at 65 °C for 72 h. After evaporation of DMSO, the remaining polymer was dissolved in water and dialyzed against diluted HCl for 12 h and against pure water for several days. The polymer solution was concentrated by rotary evaporator and freeze dried to yield a white solid powder (1.7 g, yield 81.7 %). NMR calculated Mn: 80 kDa. ¹H NMR integrated for a single repeating unit: (D₂O) $\delta_{\rm H}$: 0.5-1.76 ppm (3 H, m), 4.05 ppm (0.2 H, s), 7.10 ppm (0.22 H, s) and 7.85 ppm (0.21 H, s). IR: 620 cm⁻¹, 1260 cm⁻¹, 1750 cm⁻¹.



Scheme 2. The synthesis of PAH-His.

Binding constant evaluation from the solution

Optical absorption spectra were recorded using a UV-VIS spectrometer to investigate the complexation of copper ions with PAH-His and to establish equilibrium metal-binding constants for PAH-His in solution. In all experiments, water was used as a reference. Copper ion titration

was performed by adding aliquots of a Cu(NO₃)₂ stock solution (15 mM) to 6 mM of PAH-His solution in water. All calculations and model fittings were carried out using MS Excel software.

Surface modification

Silicon substrates were cut into 2×2 cm² slides by a DISCO dicing machine (DAD 321), cleaned with acetone and isopropanol in ultrasonic bath for 10 min, rinsed with DI water and finally dried with a stream of nitrogen. Subsequently, silicon slides were treated by oxygen plasma (200 W) for 2 min in a triple P plasma processor (Duratek, Taiwan). This cleaning procedure creates a surface rich in hydroxyl groups at the oxide surface to facilitate the subsequent silanization process. The hydroxyl-covered substrates were rinsed with methanol/toluene mixture and toluene, and then immersed in a 3 mM APTMS solution in toluene for 3 h. Subsequently, they were rinsed copiously with toluene, sonicated for 10 min, rinsed again successively with toluene, methanol and water and blown dry with nitrogen and dried under vacuum at 100 °C for 5 h.

Polymeric film fabrication

PIAMA-Me, PAH-His and 1 mM of Cu(NO₃)₂ x $3H_2O$ complexed with PAH-His (referred in this work as PAH-His(Cu)) stock solutions (1 mg/mL) were prepared by dissolving corresponding solids in 0.5 M NaCl. The synthesized PAH-His and PAH-His(Cu) solutions have the pH values of pH = 6.5 and pH = 5.5, respectively. The cleaned silicon wafers were treated with APTMS to impart positive charges onto the surface. Subsequently, substrates were immersed into the PIAMA-Me solution for 10 min, rinsed with DI water for 2 min and dried with flow of nitrogen gas. In the second step, substrates were immersed into PAH-His solution for 10 min, followed by 2 min DI water rinse and blown dry with nitrogen. This cycle was repeated until the desired number of 14 bi-layers was reached. PAH-His was always deposited as the outermost layer. Such fabricated films were cross-linked by heating the silicon substrate at 80 °C for 12 h under vacuum.⁵⁸⁻⁵⁹ The described process of LbL assembly of PIAMA-Me and PAH-His polymers is referred to as non-imprinted LbL_A in this work (Figure 2 and 3).

Cross-linked LbL films with Cu^{2+} ions preloaded at the fabrication stage were obtained by alternating immersions of the silicon slides prepared as described above, but using polymer solutions PIAMA-Me and PAH-His(Cu). 18 bi-layers (PIAMA-Me and PAH-His(Cu)) were

assembled in this way using PAH-His(Cu) as the outermost layer. The process of LbL assembly of PIAMA-Me and PAH-His(Cu) polymers is referred to as imprinted LbL_B in this work (Figure 1).

LbL films - loading and releasing of copper ions

The preparation process of the Cu^{2^+} imprinted LbL_B and non-imprinted LbL_A films on silicon substrates are illustrated in Figure 2 and 3, respectively. Loading and releasing of the copper ions was achieved by the following procedure. Imprinted LbL_B films were first washed using 1 M HCl for 5 min to remove the Cu^{2^+} ions as the Cu^{2^+} were present during the LbL process, and copious amounts of DI water, followed by drying with a stream of nitrogen. Non-imprinted LbL_A and imprinted LbL_B films were immersed in 15 mM solution of $Cu(NO_3)_2$ to absorb the Cu^{2^+} ions for 1h to ensure that equilibrium is reached and the films were washed with copious amounts of water and dried with nitrogen. In the next step, Cu^{2^+} ion-loaded non-imprinted LbL_A and imprinted LbL_B films were rinsed with 1 M HCl for 5 min to extract the Cu^{2^+} ions followed by wash with DI water and drying with nitrogen. The above procedure is repeated for several times to load and release the copper from the films. The non-imprinted LbL_A and imprinted LbL_B films with Cu^{2^+} ions were investigated using XPS to ensure that there were no copper ions left on the surface after washing with HCl.

LbL assembly and metal loading monitored by QCM

In-situ quartz crystal microbalance (QCM) measurements were conducted using a Qsense E4 multichannel instrument equipped with silicon dioxide-coated crystals. The surface of silicon dioxide chips were modified with APTMS similar to the silicon wafers as described above. For in-situ QCM measurements, the as-prepared silane modified QCM chips were fixed in the chamber of the QCM instrument and the crystals were sealed with a silicon rubber O-ring. Then the APTMS films were stabilized in a flow of water followed by 0.5 M NaCl until the frequency reached equilibrium at the flow rate of 250 μ L/min. PIAMA-Me polymer solutions of (1 mg/mL in 0.5 M NaCl) were passed through the chamber at a rate of 250 μ L/min until the frequency reached equilibrium followed by injecting of 0.5 M NaCl through the chamber at the same rate until the frequency became stable to remove any loosely bound molecules. Afterwards PAH-His polymer solutions of (1 mg/mL in 0.5 M NaCl) were pumped at the same flow rate followed by a

NaCl washing step (non-imprinted LbL_A). Similarly, in imprinted LbL_B films, alternate

injections of PIAMA-Me and PAH-His(Cu) polymer solutions in 0.5 M NaCl were delivered at a flow rate of 250 μ L/min until the frequency became stable, followed by 0.5 M NaCl salt injection after every polymer layer was deposited. The procedure was repeated until the required number of bi-layers was reached. Finally, the layers were rinsed with DI water at a flow rate of **Chemical Science Accepted Manuscript**

250 μ L/min to remove any possible excess of salt. Each layer deposition was carried out for 30 min to 1 h to achieve saturated adsorption. The frequency change of the quartz crystals was monitored throughout the adsorption process. The crystals were removed from the QCM instrument and dried with nitrogen. Cross linking of the LbL_A and LbL_B films grown on QCM chips were performed at 80 °C for 12 h under vacuum. Cross linked non-imprinted LbL_A films built on QCM chips were immersed in 15 mM of Cu(NO₃)₂ solutions for 1 h and washed with 1M HCl, similar to silicon wafers. Cross-linked films prepared by the imprinted LbL_B method on QCM were washed with 1 M of HCl for 5 min and dried. For copper loading experiments, non-imprinted LbL_A and imprinted LbL_B films on QCM chips, after washing with HCl as described above, were loaded into the chamber of the QCM instrument. Subsequently, the LbL films were stabilized in a flow of water at rate of 250 μ L/min until the frequency reached equilibrium. Then the sequence of different concentrations of Cu(NO₃)₂ stock solutions (0.1, 0.2, 0.4, 1, 2, 4, 8, 15, 25, 35 and 45 mM) were injected. For each concentration 250 μ L of Cu(NO₃)₂ solutions were injected and the frequency was allowed to reach equilibrium followed by injection of 1 mL of DI water through the chamber at the rate of $250 \,\mu$ L/min. Injection of water was intended to remove any loosely bound Cu²⁺. This procedure was repeated for all concentrations. The frequency change of the quartz crystal was monitored throughout the loading of copper ions. Finally, the crystals were removed from the QCM instrument and dried with nitrogen. Cu2+ ions loaded from non-imprinted LbLA and imprinted LbL_B films on QCM chips were washed with 1 M of HCl for 5 min and copious amounts of DI water to remove copper ions, followed by dry blowing with nitrogen. The above procedure was repeated several times to load and unload the Cu²⁺ ions using the QCM process on non-imprinted LbL_A and imprinted LbL_B films. The mass deposited on the crystal (Δm) was calculated using the Sauerbrey equation.⁶⁰

$$\Delta m = -C \frac{1}{n} \Delta f \tag{1}$$

For a 5, 15, 25, 35, 45 MHz quartz crystals, n = 1, 3, 5, 7; where $C = 17.7 \text{ ng s}^{-1} \text{ cm}^{-2}$, n =overtone number and $\Delta f =$ frequency change.

Stability

Stability and leaching rate of copper from the cross linked non-imprinted LbL_A and imprinted LbL_B films were evaluated by immersing them in artificial sea water solutions. The silicon wafers with 7 bi-layers films from non-imprinted LbL_A and 10 bi-layers from imprinted LbL_B were immersed into artificial seawater (prepared by dissolving 38.5 g/L sea salts in DI water), for up to 90 days. Samples were removed and rinsed by DI water after 1, 7, 15, 45 and 90 days immersions. The surfaces were characterized by XPS after drying with nitrogen.

Larval culture

Amphibalanus amphitrite barnacle larvae were spawned from adults collected from the Kranji mangrove, Singapore. The nauplii larvae were fed with an algal mixture 1:1 v/v of *Tetraselmis suecica* (CSIRO Strain number CS-187) and *Chaetoceros muelleri* (CSIRO Strain number CS-176) at a density of $\sim 5 \times 10^5$ /mL, and reared at 27 °C in 2.7 % salinity, 0.2 µm filtered seawater (FSW). Under these conditions, the nauplii metamorphosed into cyprids in 5 days. The cyprids were aged for 2 days at 4–6 °C prior to use in settlement assays.

Barnacle settlement and toxicity assay

The assay method as described by Wang et al.⁶¹ was adapted in this work. Positive controls consisted of plasma-cleaned silicon wafers. A 200 μ L drop of filtered sea water (FSW) containing 15 - 25 barnacle cyprids were added to each wafer kept in individual petri dish. In addition, 100 μ L of FSW containing 15 - 25 stage 2 barnacle nauplii was introduced to each droplet giving a final volume of 300 μ L. The control, as well as thenon-imprinted LbL_A and imprinted LbL_B films with copper loaded samples were incubated in the dark at 26 - 28 °C and results were scored after 24 hours. For the assessment of barnacle settlement, juvenile barnacles

and metamorphosing cyprids that had attached firmly to the substrate were counted as 'settled'. The number of dead nauplii and cyprids were also enumerated for an assessment of toxicity.

Results and Discussion

To investigate the concept of molecular imprinting of metal receptors in the LbL electrostatic thin film matrix and to employ this material in the fabrication of antifouling polymeric coatings, two custom synthesized polyelectrolytes were developed. The polymer PIAMA-Me was constructed by the opening of polyanhydride units with two nucleophilic agents, namely methanol and hydroxide. The resulting polyanion is bearing negatively charged carboxylic groups and esters functions for covalent cross-linking with amine (Figure 2 and Figure 3).⁵⁸ The results of FTIR analysis confirm the partial conversion of the anhydride polymer (PIAMA) to ester (PIAMA-Me) through the methanolysis reaction as shown in Scheme 1. The double peak (1860 cm⁻¹ and 1780 cm⁻¹) belonging to the C=O of the anhydride polymer (PIAMA) disappears and a new peak at 1720 cm⁻¹, attributed to the C=O of ester bond, becomes visible. In addition, a peak at 1580 cm⁻¹ can be associated to a newly formed COO⁻ (Figure S1A in the supplementary information).⁶² The methanolysis reaction of PIAMA was further verified by comparison of ¹H NMR spectra of PIAMA-Me and purified polyanhydride as shown in Figure S2 (supplementary information). The presence of a peak at 3.66 ppm in the polymer PIAMA-Me spectrum, belonging to the protons of methyl ester, confirms successful conversion. The methyl ester peak (3.66 ppm) integrated area, and the dimethyl group protons of the main chain (peaks between 0.5-1.36 ppm) allow us to estimate degree of substitution with the methyl esters groups to obtain the value of 70% (Scheme 1).

The polycation (PAH-His) was synthesized by grafting of L-histidine side groups on poly allylamine backbone. Following this approach, peptide mimicking polymer bearing histidine moieties for metal ion complexation and primary amine groups used in the LbL assembly was synthesized (Scheme 2). Successful polymer synthesis was confirmed by the appearance of peaks at 620 cm⁻¹ and 1260 cm⁻¹, which are assigned to OCN bending and amide C-N stretching and N-H bending frequencies respectively. The ester peak visible at 1750 cm⁻¹ in pure histidine, and it is absence in PAH-His spectrum (see supplementary information Figure S1B) as well as

presence of the peaks at 1260 cm⁻¹ (amide C-N stretching) and at 1570 cm⁻¹ (amide carbonyl) in the PAH-His polymer spectrum, confirm successful chemical transformation.⁶³ The PAH-His structure and histidine substitution degree were examined and quantitatively evaluated by ¹H-NMR as shown in Figure S3 (supplementary information). Reaction completion is confirmed by disappearance of the methyl ester peak at 3.67 ppm and presence of new peaks at 4.05, 7.10 and 7.85 ppm, which can be associated to the methine and imidazole protons in the His moiety, respectively. The backbone substitution degree is determined to 20 % owing to the clearly visible signal of imidazole protons (7 and 8 ppm) and the aliphatic backbone protons peaks between (1-2 ppm). This allowed us to estimate the indices of repeating units, describing the composition of polymer PAH-His, to 130 and 500 for PAH-substituted and non-substituted groups respectively (Scheme 2).

Complexation of copper ions in aqueous solutions

UV-visible absorption spectroscopy experiments were employed to establish the mechanism of interactions between the synthesized polymer (PAH-His) and Cu^{2+} ions in solution. Titration of PAH-His with $Cu(NO_3)_2$ in water followed by observation of absorption changes allowed us to propose a binding model for these molecules. On the basis of previous reports,^{44, 47} we employed the working hypothesis that two histidine ligands bind to a single copper ion. To simplify the model, repeating histidine units of the polymer were treated as individual, independently binding molecules and polyallylamine interactions with ions were omitted. This is justified since primary amine interactions with Cu²⁺ are reported to be much weaker compare to imidazole complexation.^{18, 36, 64} As shown in Figure 1A, the addition of copper nitrite to the PAH-His solutions results in the appearance of a new peak at 620 nm. The peak intensity is increasing continuously with the addition of copper and reaches the absorption maxima approximately for 1:2 metal to ligand ratio. After the saturation, the intensity of the signal at 620 nm is decreasing with a clear shoulder increase at longer wavelengths. Upon further metal addition a new signal near 820 nm appears which can be assigned to free Cu²⁺ ions.^{44, 65} All UV absorption spectra were fitted by three peaks representing the concentration of three different species present in the solution using asymmetric modification of the Gaussian function.⁶⁶⁻⁶⁷ The peak at 620 nm is attributed to the formation of a 1:2 copper complex with PAH-His, peak around 770 nm corresponds to the formation of a 1:1 copper complex with PAH-His and a peak at 820 nm is 12

linked to free copper ions (Figure 1B and Figure S4).⁴⁴ Examples of fitting procedure are provided in the supplementary information.

The integrated absorption values for the three signals are provided in Figure 1C. The changes observed in the signal intensity variations were subsequently used to establish the binding constants of this system. Titration curves were fitted by assuming that 1:1 and 1:2 equilibriums are present, ⁶⁸

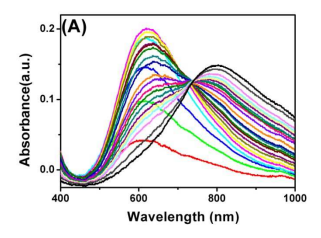
$$M + PAH - His \Leftrightarrow M(PAH - His) \tag{2}$$

$$K_1 = \frac{[M(PAH - His)]}{[M] * [PAH - His]}$$
(3)

$$M(PAH - His) + PAH - His \Leftrightarrow M(PAH - His)_{2}$$
(4)

$$K_2 = \frac{[M(PAH - His)_2]}{[M(PAH - His)]^*[PAH - His]}$$
(5)

The absorbance signals presented in Figure 1C clearly support the co-existence of the 1:1 and 1:2 stoichiometry of Cu^{2+} and PAH-His complex. Fitting of the data provides binding constant values of $K_1 = 8.43 \times 10^3 \text{ M}^{-1}$ (log $K_1 = 3.93$) and $K_2 = 2.63 \times 10^4 \text{ M}^{-1}$ (log $K_2 = 4.42$). Based on the above observations it appears that the 1:2 metal / PAH-His coordination dominates the 1:1 interactions. The results are consistent with a large stability constant observed for other histidine grafted polymers.^{44, 47}



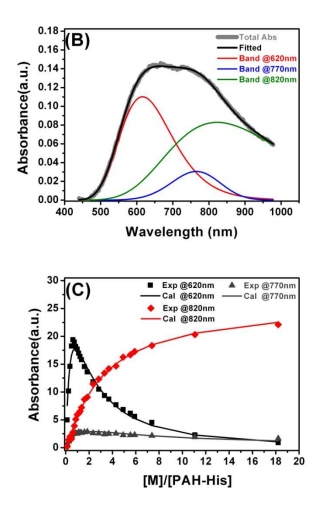


Figure 1. (A) UV-VIS absorption spectra of PAH-His coordinated with Cu, (B) Example of absorption spectra fitted with three different signals for 2.3 mL of PAH-His (6 mM) added to the 2 mL of $Cu(NO_3)_2$ (15 mM) (C) integration of the signal intensity at 620 nm, 770 nm and 820 nm, fitting data to the binding model.

LbL film fabrication

Synthesized polymers were used for fabrication of thin film structures using alternating deposition of polyelectrolytes in a typical electrostatic LbL protocol,¹⁰⁻¹³ followed by the interlayer film cross-linking performed through amide bond formation.⁵⁸⁻⁵⁹ Two different designs were compared to investigate the effect of metal imprinting on the affinity of the polymeric matrix to bind Cu^{2+} ions. Imprinting of ion receptors was achieved by immobilization of

histidine moieties around preorganized metal atoms in the covalent, interlayer cross-linking process (Figure 3).

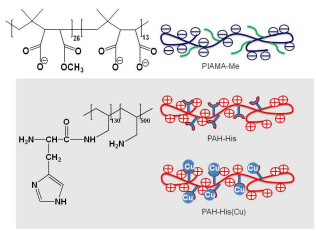


Figure 2. Schematic illustration of PIAMA-Me, PAH-His and PAH-His(Cu).

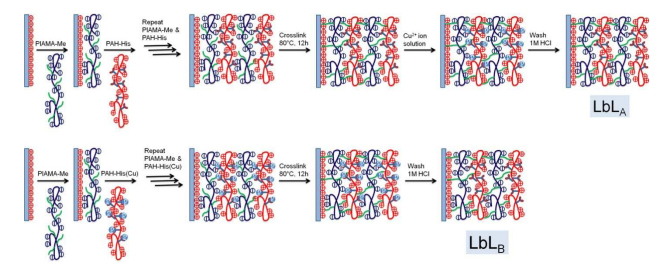


Figure 3. LbL film formation by different processing methods, (non-imprinted LbL_A refers to assembly of PIAMA-Me and PAH-His, and imprinted LbL_B refers to assembly of PIAMA-Me and PAH-His(Cu)). Cross linking of the films as employed followed by loading and releasing of copper within the LbL films. Schematic representation of the polymer PIAMA-Me, PAH-His and PAH-His(Cu) are shown in Figure 2.

The two methods used essentially differ in the cross linking step. Non-imprinted LbL_A films are cross-linked without the metal template present and as such, the mobility of the receptors is reduced, maintaining random arrangement not influenced by the presence of copper. Contrary, in the imprinted LBL_B system, the film is loaded with Cu^{2+} ions at the stage of fabrication, so the 15

subsequent cross-linking process is intended to "freeze" histidine receptors in the given conformation around the guest molecules (Figure 3). Using these approaches two types of structures with similar chemical composition but different arrangement of metal receptors can be obtained. Film cross linking evidence was acquired using FTIR and XPS methods. Disappearance of the ester peak (1720 cm⁻¹) in the non-imprinted LbL_A film and the evolution of a new peak at 1590 cm⁻¹ (amide C-N stretch) indicate amide bond formation (Figure 4) and successful covalent attachment between the layers.⁶²

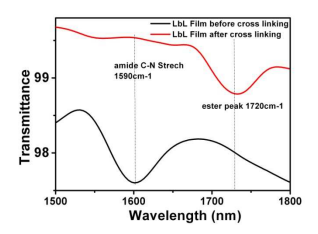


Figure 4. FTIR spectra of non-imprinted LbL_A film before and after cross linking.

XPS spectra (Figure S5 in supplementary information) further confirmed the cross-linking of the LbL films in addition to the FTIR analysis. S5A and S5B are the N1s core-level spectra for noncross linked and cross-linked LbL_A films, respectively, which are fitted into primary amine, amide, imide and protonated amine components with binding energy (B.E.) values of 398.9 eV, 399.9 eV, 400.8 and 401.8 eV.⁶⁹ In the spectra of cross-linked non-imprinted LbL_A films, the peak component for the free and protonated amines are small. The increase of the amide peak intensity demonstrates that free amine is reacting with ester. A similar trend is also observed for the imprinted LbL_B films (Figure S5C and S5D). The thickness of the cross linked LbL films were measured by ellipsometry and by the AFM scratch method as shown in Figure 5. For the non-imprinted LbL_A, the thickness growth of the films is linear up to 8 bi-layers and exponential afterwards, following trends reported in the literature (Figure 5A).⁷⁰ Similarly, for the imprinted LbL_B we observed linear growth until 14 bi-layers and exponential increase afterwards. The 16 thickness of the non-imprinted LbL_A and imprinted LbL_B films were also measured independently using AFM scratch method. The obtained thickness values correlate well with the ellipsometry measurements as shown in the figure 5A and 5B. To achieve a good comparison among the structures studied, films of similar thicknesses and comparable masses of the polymers deposited were used. This corresponds to 7 bi-layers and 10 bi-layers for nonimprinted LbL_A and imprinted LBL_B, respectively.

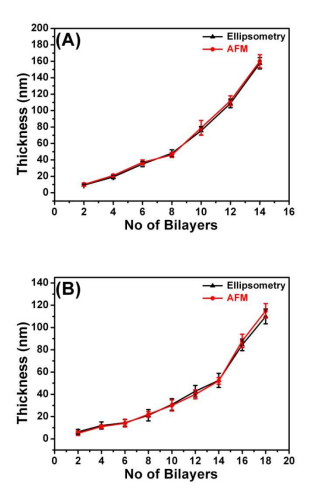


Figure 5. Thickness data of non-imprinted LbL_A (Figure 5A) and imprinted LbL_B (Figure 5B) films from ellipsometry and AFM scratch test.

The morphology of the LbL films was studied using AFM (Figure. S6 in supplementary information). The AFM images in Figure S6 show clear variations of the film surface depending on the fabrication method applied. Figure S6A shows cluster/coil-like features on the non-cross

linked LbL_A film, and no significant changes in morphology upon application of the crosslinking procedure (Figure S6B). Rather different morphology is observed for LbL_B films. The LbL_B film is fairly smooth (Figure S6D) when compared to LbL_A (Figure S6B). The surface roughness R_q (the quadratic mean value of roughness, obtained over a 5 x 5 μ m² surface area) for an LbL_A film without cross-linking exhibited a value of Rq = 16±0.5 nm (Figure. S6A) compared to Rq = 13±0.5 nm (Figure. S6B) for cross-linked film. Similarly, for the imprinted LbL_B films, the roughness values of the non-cross linked film (Rq = 10±1 nm from Figure S6C) decreased to 8±1 nm (Figure. S6D) after cross-linking. The results are consistent with previous reports showing that the surface roughness values are reduced with an increased cross-linking density.^{58, 70}

LbL growth and metal affinity assessed by QCM

To compare LbL_A and LbL_B film affinity to ions by establishing binding constants for each film and finally to answer the question if the proposed molecular imprinting method allows us to enhance the metal binding capacity of LbL structures, QCM experiments were performed. The buildup of the macromolecules during fabrication of LbL structures on the APTES modified chips, was monitored following frequency changes of QCM signals (Figure 6). The alternating addition of polycation and polyanion leads to a stepwise rise of the mass for the deposited layers in both non-imprinted and imprinted films (Figure 6), demonstrating successful layer by layer assembly. The observed increase in the mass of the LbL_A (Figure 6A)is higher than the LbL_B (Figure 6B). This is most likely related to the the pH of the polymer solution. This parameter is lowered by the presence of Cu salt in the PAH-His(Cu), and is responsible for thinner layers of LbL_B.⁷¹

Successful LbL deposition is known to be facilitated by the overcompensation of charges, which in turn promotes the binding of the subsequent polyelectrolytes.¹⁰ Earlier studies suggest that for electrostatically driven LbL deposition both the substrate and the polyelectrolyte species must be sufficiently charged. If the charge is insufficient, the adsorbed layers may be partially removed upon adsorption on the next polyelectrolyte layer.⁷² Similar behavior is observed here when the LbL films were assembled by LBL_A with PIAMA-Me and PAH-His polymers. It is also suggested that reason for such behavior is the formation of free polyelectrolyte complexes in the

solution, which are entropically favored in this case compared to a multilayer on the surface.⁷³⁻⁷⁴ Similar complexes may occur when PAH-H is interacting with PIAMA-Me.

Thickness values found from QCM data for the cross-linked 7 bi-layer non-imprinted LbL_A and the 10 bi-layer imprinted LbL_B films are 35 ± 5 nm. The total mass of the PAH-His layers in the non-imprinted LbL_A and imprinted LbL_B films are 15.9, and 9.3, μ g/cm², respectively.

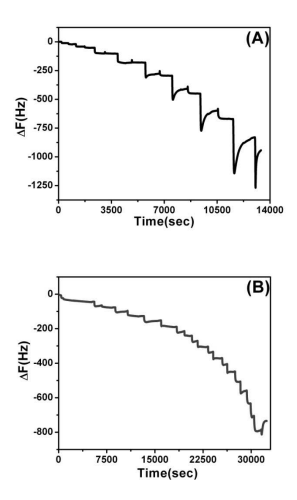


Figure 6. Frequency changes monitored by QCM-D for (A) non-imprinted LbL_A (7 bi-layers) and (B) imprinted LbL_B (10 bi-layers) films.

Film - metal affinity experiments were carried out by passing Cu^{2+} solutions of different concentrations over the thin film until equilibrium state is reached (Figure S7 in the supplementary information). The adsorption isotherms obtained in this way display a typical profile of saturation with increased concentration of guest molecules (Figure 7A). The amount of

absorbed mass was calculated from the Sauerbrey equation as stated in the experimental section. The equilibrium adsorbed amount of metal for imprinted LbL_B films is 11.25 μ g/cm², whereas for non-imprinted LbL_A only 4.95 μ g/cm² could be immobilized. These observations clearly demonstrate the higher adsorption capacity of the LbL_B compare to the non-imprinted LbL_A and support the hypothesis of enhanced affinity of the polymeric film to the metal upon imprinting of receptor structure during the fabrication process.

QCM equilibrium adsorption experiments were conducted to investigate the binding behavior and binding constants for both LbL_A and LbL_B (Table 1). The saturation isotherms of these films are displayed in Figure 7A. With the concentration of the copper solution increased, the adsorption capacity of both films first increases sharply then the increase slowes down until saturation is reached.

A typical method for assessing guest affinity for thin films is based on the application of the simple Langmuir adsorption isotherm model (Figure 7B). Using this approach we assume a homogeneous distribution of binding sites with equal energy across the film and that the guest diffusion through the thin film is fast and is not limiting step.^{36, 41, 75} Learning from initial experiments in solution we also know that both films are interacting with metal ions utilizing predominantly receptors composed of two histidine groups per binding site. In case of the film affinity this simplifies binding to 1:1 type of interaction but with stoichiometry of two histidine moieties per Cu²⁺. Data obtained in this way (Table 1) display substantially higher binding affinity of Cu²⁺ for imprinted LbL_B film (logK=3.9) compare to non-imprinted LbL_A (logK=3.0).

Process	Method	Binding constant	log K
PAH-His (solution)	UV-Visible	$K_1 = 8.43 \times 10^3$	$K_1 = 3.93$
		$K_2 = 2.63 \times 10^4$	$K_2 = 4.42$
Non-imprinted - LbL _A	QCM (from Langmuir isotherm)	$K_1 = 1.04 \text{ x } 10^3$	$K_1 = 3.02$
Imprinted - LbL_B	QCM (from Langmuir isotherm)	$K_1 = 7.24 \text{ x } 10^3$	$K_1 = 3.86$

 Table 1. Equilibrium constants for PAH-His and imprinted and non-imprinted LbL films from different methods

 Process
 Nothed

 Process
 log K

The Cu^{2+} template-defined organization of PAH-His receptors was protected in the cross-linking process of the LbL_B film fabrication. After the template extraction, imprinted LbL_B structures remained in place to furnish a spatial configuration of histidine ligands for the Cu^{2+} coordination. Contrary, enhanced metal binding cavities derived from spatial arrangements of functional groups, and corresponding to Cu^{2+} , were absent in the LbL_A film due to the lack of metal template in the preparation process. As such LbL_A displays randomly immobilized histidine receptors with lower affinity to copper ions compare to LbL_B.

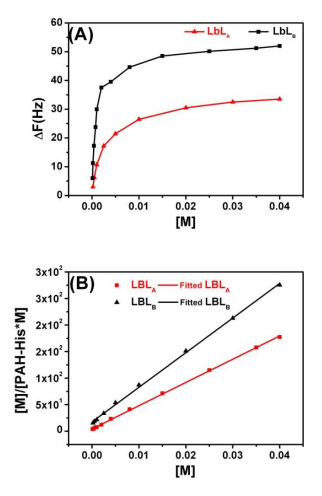


Figure 7. (A) Saturation isotherms monitored by QCM-D for Cu^{2+} loading in non-imprinted LbL_A and imprinted LbL_B films. The surface proved fully saturable and followed normal Langmuir-like adsorption behavior. ΔF represents the frequency change (Hz). (B) Adsorption isotherms of non-imprinted LbL_A and imprinted LbL_B films for Cu²⁺ ions fitted by the Langmuir model. [M] denotes the molar concentration of Cu²⁺.

Rebinding capacities of the LbL_A and LbL_B films were calculated using QCM collected data after washing the films with 1M HCl. Both films showed Cu rebinding levels of 90% and 95% respectively (see the saturation isotherms in the supplementary information Figure S8). Comparison of LbL_A and LbL_B for affinity toward Cu²⁺ ion was also investigated using XPS (Figure 8). The intensities of the Cu2p_{3/2} peak at 933 eV and the Cu2p_{1/2} peak at 953 eV were used for monitoring the trapped copper inside the LbL films.¹⁵ Copper can be detected for both types of imprinted LbL_A and non-imprinted LbL_B films, and it can be removed by washing with 1 M of HCl for 5 min. The cycle can be repeated for several times and these results are indicating that both films possess stable affinity toward copper ions.

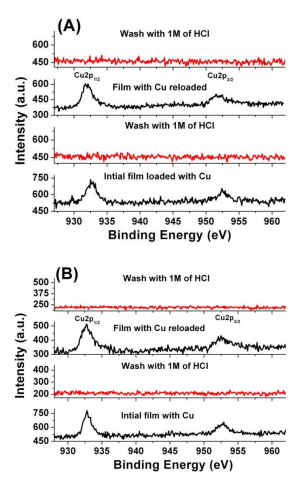


Figure 8. Cu 2p XPS spectra of the LbL film before and after loading of copper in the (A) non-imprinted LbL_A films and (B) imprinted LbL_B films.

The study of the leaching behavior for the metal-bearing films is an important way to obtain valuable information about the chemical properties of the metals in stabilized products and their potential environmental risks in antifouling applications. A better understanding of the leaching rate of the metals trapped inside the LbL films exposed to different environmental conditions is helpful for practical applications. To compare the copper stabilization effect the non-imprinted LbL_A and imprinted LbL_B films were subjected to prolonged 90 days leaching experiments (Figure 9) in artificial sea water environment.

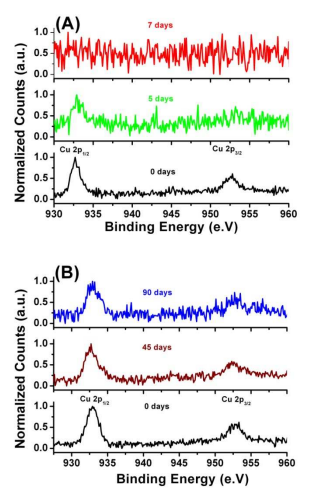


Figure 9. Cu 2p XPS spectra of the non-imprinted LbL_A (Figure 9A) and imprinted LbL_B films (Figure 9B) against sea salts.

XPS results provide another confirmation of the enhanced affinity of the imprinted LbL_B film toward Cu^{2+} ions. Substantially lower amounts of copper ions leach out of the LbL_B after six

weeks of exposure to artificial sea water as compared with the LbL_A. The leaching rate of copper from non-imprinted LbL_A is thus obviously faster and all the copper is leached out from the film within a week as shown in the XPS spectra (Figure 9A). In addition, we have also monitored the N1s XPS spectra of both types of the films. Corresponding results (Figure S9A and S9B) showed similar spectra indicating that the films are stable in the sea water environment for a prolonged period of time. Thickness values for both films showed the same values even after 45 days in sea salts solution with minor 5-10% loss after 90 days (measured by ellipsometry). From the above results we can state that the cross-linking between the ester groups from the polyanion and amine groups from the polycation, plays a vital role in improving chemical stability of the films under seawater conditions. Moreover, the presented method can be also used to design surfaces with customized leaching rate of metal ions in addition to enhancement of copper binding within the imprinted film.

Antifouling properties:

Barnacles are common, cosmopolitan macrofouling organisms and can rapidly colonize unprotected surfaces submerged in the marine environment. Barnacle larvae (cyprids) are widely used in the lab scale antifouling tests, since cyprids larvae of the barnacles settle readily in static water assays.⁷⁶ In this study a barnacle cyprid bioassay was conducted to evaluate the settlement and toxicity effects of imprinted LbL_A and non-imprinted LbL_B films with copper.

Slightly elevated mortality was observed for naupli incubated for 24 hours on surfaces coated with copper loaded LbL films fabricated with both discussed methods compare to the control sample. There was no settlement of the cyprids on non-imprinted LbL_A and imprinted LbL_B films with copper. (Figure 10) These results indicate that Cu^{2+} released from the surface is reaching required fouling preventive concentrations. As the assay is conducted in a static droplet, the concentration of copper present in the seawater droplet is the cumulative value achieved over the 24 hour leaching period.

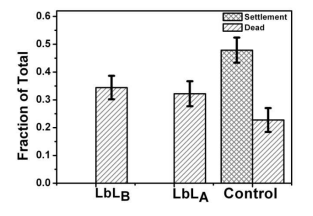


Figure 10. Cyprid settlement and toxicity of imprinted and non-imprinted LbL films with copper loading. Each value is the mean of 8 replicate measurements. The error bars here are standard deviations.

Statistical comparisons were performed using GraphPad Prism 5 (GraphPad Software Inc.). Cyprid mortality and settlement test data were analyzed using a Kruskal-Wallis test and Dunn's post test to evaluate the performance of the LbL films with copper loading. There is no significant difference between the cyprid mortality on imprinted LbL_A and non-imprinted LbL_B films and control samples (Kruskal-Wallis test, p = 0.1133 and Dunn's post test, p > 0.05).⁷⁷ Therefore we can conclude that there is no toxicity difference among those surfaces. However, there was no cyprid settlement for non-imprinted LbL_A and imprinted LbL_B films. Lethal concentration (LC₅₀) reported for barnacles naupli and barnacle cyprids is 0.71 and 63 µg/mL respectively.⁷⁸⁻⁷⁹ Leached Cu concentration in the model experiment, when LbL_A sample was immersed in 300 µL of water, was established on 60 µg/mL. This value is considerably lower in the real biological experiments when volume of the solution is much higher. The results suggest that the amount of copper released is below the toxicity threshold to kill barnacle cyprids but sufficient to prevent the settlement.

Conclusions:

A peptide mimicking polymer, bearing histidine side groups, was used to construct thin LbL architectures with high affinity to bind Cu^{2+} ions. Enhanced binding ability was achieved through a metal imprinting process involving covalent immobilization of receptors within the film 25

structure in presence of metal ions. Polycation bearing imidiazole-type histidine ligands for metal binding, and polyanion with high density of methyl esters for covalent cross-linking were custom synthesized for this task. A comparison of metal imprinted LbL_B and non-imprinted LbL_A films allowed us to conclude that by following the established approach it is possible to enhance the affinity of thin films to bind copper up to a seven-fold. This number is likely to be increased when further process optimization at the fabrication stage is considered. Physical and chemical characterization techniques showed that the prepared imprinted LbL_B films display excellent stability under harsh seawater conditions and very slow leaching rate of metals compared to non-imprinted LbL_A films.

Finally, the films are promising antifouling materials showing noticeable performance against barnacles. We believe that the presented approach may lead to a generic platform for marine antifouling materials with very low net release rate of biocides. Particularly it may help to reduce the amount of copper currently used commercial antifouling products. By further improvement of metal ion binding one can imagine that this technology can be directly used to obtain paints with very high metal affinity. For example, absorbing copper in places with high concentration like harbors or polluted river mouths and then slowly releasing it in an open sea could be envisaged. The fabrication approach presented can also benefit metal extraction processes used for example by the water purification or natural resources recovery industries. The phenomenon described here for copper ions could be further explored in heavy metal separations, antifungal layers, protein binding, catalysis and fluorescence sensing.

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Notes and References

Electronic Supplementary Information available: FTIR, NMR spectra of synthesized polymers, XPS spectra and AFM images of non-cross linked and cross linked LBL_A and LBL_B films, UV-Visible absorption spectra of copper complexation with PAH-His, QCM data of LBL_A and LBL_B films and stability of the films are provided in the electronic supplementary information.

- ¹ T. Farhat and P. T. Hammond, *Adv Funct Mater.*, 2005, **15**, 945.
- ² T. Farhat and P. T. Hammond, *Adv Funct Mater.*, 2006, **16**, 433.
- ³ J. E. Gu, S. Lee, C. M. Stafford, J. S. Lee, W. Choi, B. Y. Kim, K. Y. Baek, E. P. Chan, J. Y. Chung, J.
- Bang, and J. H. Lee, *Adv Mater.*, 2013, **25**, 4778.
- ⁴ M. C. Petty, *Stud. Interf. Sci.*, 2002, **16**, 317.
- ⁵ I. Welterlich and B. Tieke, *Macromolecules*, 2011, **44**, 4194.
- ⁶ S. A. Makohliso, L. Giovangrandi, D. Leonard, H. J. Mathieu, M. Ilegems, and P. Aebischer, *Biosens. Bioelectron.*, 1998, **13**, 1227.
- ⁷ A. Shukla, J. C. Fang, S. Puranam, and P. T. Hammond, *J. Control. Release*, 2012, **157**, 64.
- ⁸ J. Song, D. Jańczewski, Y. Guo, J. Xu, and G. J. Vancso, *Nanoscale*, 2013, **5**, 11692.
- ⁹ J. Song, D. Jańczewski, Y. Ma, M. Hempenius, J. Xu, and G. J. Vancso, *J. Colloid. and Interf. Sci.*, 2013, **405**, 256.
- ¹⁰ G. Decher, *Science*, 1997, **277**, 1232.
- ¹¹ Y. Yan, M. Björnmalm, and F. Caruso, *Chem. Mater.*, 2014, **26**, 452.
- ¹² A. L. Becker, A. P. Johnston, and F. Caruso, *Small*, 2010, **6**, 1836.
- ¹³ G. Decher and J. B. Schlenoff, 'Multilayer Thin Films: Sequential Assembly of Nanocomposite Materials', ed. G. Decher and J. B. Schlenoff, Wiley-VCH, 2003.
- ¹⁴ D. Jańczewski, J. Song, and G. J. Vancso, *Eur. Polym. J.*, 2014, **54**, 87.
- ¹⁵ S. R. Puniredd and M. P. Srinivasan, *Ind. Eng. Chem. Res.*, 2005, **46**, 464.
- ¹⁶ A. Mentbayeva, A. Ospanova, Z. Tashmuhambetova, V. Sokolova, and S. Sukhishvili, *Langmuir*, 2012, **28**, 11957.
- ¹⁷ M. A. Rahim, M. S. Islam, T. S. Bae, W. S. Choi, Y.-Y. Noh, and H.-J. Lee, *Langmuir*, 2012, **28**, 8486.
- ¹⁸ S. Bhattacharjee, J. Dong, Y. Ma, S. Hovde, J. H. Geiger, and G. L. Baker, *Langmuir*, 2012, **28**, 6885.
- ¹⁹ R. G. de Paiva, M. A. de Moraes, F. C. de Godoi, and M. M. Beppu, J. Appl. Polym. Sci., 2012, **126**, E17.
- ²⁰ D. Jańczewski, D. N. Reinhoudt, W. Verboom, C. Hill, C. Allignol, and M. Duchesne, *New J. Chem.*, 2008, **32**, 490.
- ²¹ A. H. Chen, S. C. Liu, C. Y. Chen, and C. Y. Chen, J. Hazard. Mater., 2008, **154**, 184.
- ²² G. P. Rao, C. Lu, and F. Su, *Sep. Purif. Technol.*, 2007, **58**, 224.
- ²³ K. Haupt and K. Mosbach, *Chem. Rev.*, 2000, **100**, 2495.
- ²⁴ W. Li and S. J. Li, 'Oligomers Polymer Composties Molecular Imprinting', Springer, 2006.
- ²⁵ R. Y. Liu, G. J. Guan, S. H. Wang, and Z. P. Zhang, *Analyst* 2011, **136**, 184.
- ²⁶ P. Turkewitsch, B. Wandelt, G. D. Darling, and W. S. Powell, *Anal. Chem.*, 1998, **70**, 2025
- ²⁷ G. Wulff, *Chem. Rev.*, 2002, **102**, 1.
- ²⁸ Y. Zhan, X. Luo, S. Nie, Y. Huang, X. Tu, and S. Luo, *Ind. Eng. Chem. Res.*, 2011, **50**, 6355.
- ²⁹ X. Y. Bi, R. J. Lau, and K. L. Yang, *Langmuir*, 2007, **23**, 8079.
- ³⁰ B. J. Gao, F. Q. An, and Y. Zhu, *Polymer*, 2007, **48**, 2288.
- ³¹ Buhani, Narsito, Nuryono, and E. S. Kunarti, *Desalination*, 2010, **251**, 83.
- ³² Buhani, Suharso, and Sumadi., *Desalination*, 2010, **259**, 140.
- ³³ G. Z. Fang, J. Tan, and X. P. Yan, *Anal. Chem*, 2005, **77**, 1734.
- ³⁴ X. Huang, A. B. Schubert, J. D. Chrisman, and N. S. Zacharia, *Langmuir*, 2013, **29**, 12959.
- ³⁵ K. C. Krogman, K. F. Lyon, and P. T. Hammond, *J. Phys. Chem. B*, 2008, **112**, 14453.
- ³⁶ S. Wijeratne, M. L. Bruening, and G. L. Baker, *Langmuir*, 2013, **29**, 12720.
- ³⁷ H. Chen, G. Zeng, Z. Wang, and X. Zhang, *Macromolecules*, 2007, **40**, 653.
- ³⁸ P. Schuetz and F. Caruso, *Adv. Funct. Mater.*, 2003, **13**, 929.
- ³⁹ A. M. Balachandra, J. Dai, and M. L. Bruening, *Macromolecules*, 2002, **35**, 3171.
- 27

- ⁴⁰ T. Li, S. Chen, H. Li, Q. Li, and L. Wu, *Langmuir*, 2011, **27**, 6753.
- ⁴¹ H. Deng, L. Gao, S. Zhang, and J. Yuan, *Ind. Eng. Chem. Res.*, 2012, **51**, 14018.
- ⁴² D. Sanna, G. Micera, C. Kállay, V. Rigóc, and I. Sóvágóc, *Dalton Trans.*, 2004, 2702.
- ⁴³ I. Sóvágó and K. Ősz, *Dalton Trans.*, 2006, 3841.
- ⁴⁴ X. Bi and K.-L. Yang, *Langmuir*, 2007, **23**, 11067. ⁴⁵ B. Daschamps, P. P. Kulkarni, M. Gautam Pasak, a
- P. Deschamps, P. P. Kulkarni, M. Gautam-Basak, and B. Sarkar, *Coord. Chem. Rev.*, 2005, 249, 895.
- ⁴⁶ H. Sigel and R. B. Martin, *Chem. Rev.*, 1982, **82**, 385.
- ⁴⁷ D. E. Fullenkamp, L. He, D. G. Barrett, W. R. Burghardt, and P. B. Messersmith, *Macromolecules*, 2013, **46**, 1167.
- ⁴⁸ Y. Kobayashi, Y. Takashima, A. Hashidzume, H. Yamaguchi, and A. Harada, *Sci. Rep.*, 2012, **3**, 1243.
- ⁴⁹ Y.-C. Chung, W.-Y. Hsieh, and T.-H. Young, *Biomaterials*, 2010, **31**, 4194.
- ⁵⁰ S. Dobretsov, R. M. Abed, and M. Teplitski, *Biofouling*, 2013, **29**, 423.
- ⁵¹ M. Tasso, S. L. Conlon, A. S. Clare, and C. Werner, *Adv. Funct. Mater.*, 2012, **22**, 39.
- ⁵² D. M. Yebra, S. Kill, and K. Dam-Johansen, *Prog. Org. Coat.*, 2004, **50**, 75.
- ⁵³ E. Almeidaa, T. C. Diamantinoa, and O. de Sousab, *Prog. Org. Coat.*, 2007, **59**, 2.
- ⁵⁴ A. P. Negri and A. J. Heyward, *Mar. Environ. Res.*, 2001, **51**, 17.
- ⁵⁵ A. Sousa, L. Genio, S. Mendo, and C. Barrosoi, *Appl. Organomet. Chem.*, 2005, **19**, 324.
- ⁵⁶ I. Omae, *Chem. Rev.*, 2003, **103**, 3431.
- ⁵⁷ G. Damiens, C. Mouneyrac, F. Quiniou, E. His, M. Gnassia-Barelli, and M. Romeo, *Environ. Pollut.*, 2006, **140**, 492.
- ⁵⁸ X. Zhu, D. Jańczewski, S. S. C. Lee, S. L.-M. Teo, and G. J. Vancso, *ACS Appl. Mater. Interfaces*, 2013, **5**, 5961.
- ⁵⁹ X. Zhu, S. Guo, D. Jańczewski, F. J. P. Velandia, S. L.-M. Teo, and G. J. Vancso, *Langmuir*, 2014, **30**, 288.
- ⁶⁰ K. A. Marx, *Biomacromolecules*, 2003, **4**, 1099.
- ⁶¹ Y. Wang, D. E. Betts, J. A. Finlay, L. Brewer, M. E. Callow, J. A. Callow, D. E. Wendt, and J. M. DeSimone, *Macromolecules*, 2011, **44**, 878.
- ⁶² E. Pretsch, P. Bnhlmann, and C. Affolter, 'Structure Determination of Organic Compounds', Springer, 2009.
- ⁶³ J. G. Mesu, T. Visser, F. Soulimani, and B. M. Weckhuysen, *Vib. Spectrosc.*, 2005, **39**, 114.
- ⁶⁴ R. J. Sundberg and R. B. Martin, *Vib. Spectrosc.*, 1974, **74**, 471.
- ⁶⁵ H. Yang, M. Pritzker, S. Y. Fung, Y. Sheng, W. Wang, and P. Chen, *Langmuir*, 2006, **22**, 8553.
- ⁶⁶ J. Song, D. Jańczewski, Y. Ma, L. V. Ingen, C. E. Sim, Q. Goh, J. Xu, and G. J. Vancso, *Eur. Polym. J.*, 2013, **49**, 2477.
- ⁶⁷ A. L. Stancik and E. B. Brauns, *Vib. Spectrosc.*, 2008, **47**, 66.
- ⁶⁸ A. E. Hargrove, Z. Zhong, J. L. Sessler, and E. V. Anslyn, *New J. Chem.*, 2010, **34**, 348.
- ⁶⁹ S. R. Puniredd and M. P. Srinivasan, *Langmuir*, 2005, **21**, 7812.
- ⁷⁰ I. Choi, R. Suntivich, F. A. Plamper, C. V. Synatschke, A. H. E. Muller, and V. V. Tsukruk, *J. Am. Chem. Soc.*, 2011, **133**, 9592.
- ⁷¹ K. Glinel, C. Déjugnat, M. Prevot, B. Schöler, M. Schönhoff, and R. V. Klitzing, *Colloids Surf.*, A, 2007, 303, 3.
- ⁷² B. Schoeler, G. Kumaraswamy, and F. Caruso, *Macromolecules*, 2002, **35**, 889.
- ⁷³ S. T. Dubas and J. B. Schlenoff, *Macromolecules*, 2001, **34**, 3736.
- ⁷⁴ D. P. Go, A. Hung, S. L. Gras, and A. J. O'Connr, *J. Phys. Chem. B*, 2012, **116**, 1120.
- ⁷⁵ Z. Pei, H. Anderson, T. Aastrup, and O. Ramstrom, *Biosens. Bioelectron.*, 2005, **21**, 60.
- ⁷⁶ E. R. Holm, *Integr. Comp. Biol.*, 2012, **52**, 348.
- ⁷⁷ D. Rittschof, B. Orihuela, T. Harder, S. Stafslien, B. Chisholm, and G. H. Dickinson, *PLoS One*, 2011, **6**, e16487.
- ⁷⁸ C. Bressy, C. Hellio, J. P. Maréchal, B. Tanguy, and A. Margaillan, *Biofouling*, 2010, **26**, 769.
- ⁷⁹ V. W. W. Bao, K. M. Y. Leung, J.-W. Qiu, and M. H. W. Lam, *Mar. Pollut. Bull.*, 2011, **62**, 1147.