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EXINTERFACES

Spray Layer-by-Layer Assembled Clay Composite Thin Films as Selective Layers in Reverse Osmosis Membranes

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S Supporting Information

[AB](#page-6-0)STRACT: [Spray layer-b](#page-6-0)y-layer assembled thin films containing laponite (LAP) clay exhibit effective salt barrier and water permeability properties when applied as selective layers in reverse osmosis (RO) membranes. Negatively charged LAP platelets were layered with poly(diallyldimethylammonium) (PDAC), poly(allylamine) (PAH), and poly(acrylic acid) (PAA) in bilayer and tetralayer film architectures to generate uniform films on the order of 100 nm thick that bridge a porous poly(ether sulfone) support to form novel RO membranes. Nanostructures were formed of clay layers intercalated in a polymeric matrix that introduced size-exclusion transport mechanisms into the selective layer. Thermal crosslinking of the polymeric matrix was used to increase the mechanical stability of the films and improve salt rejection by constraining swelling during operation. Maximum salt rejection of 89% was observed for the tetralayer film architecture, with an order of magnitude increase in water permeability compared to commercially available TFC-HR membranes. These clay

composite thin films could serve as a high-flux alternative to current polymeric RO membranes for wastewater and brackish water treatment as well as potentially for forward osmosis applications. In general, we illustrate that by investigating the composite systems accessed using alternating layer-by-layer assembly in conjunction with complementary covalent cross-linking, it is possible to design thin film membranes with tunable transport properties for water purification applications.

KEYWORDS: layer-by-layer, spray layer-by-layer, laponite clay, composite thin film, reverse osmosis membrane, nanofiltration membrane, water permeation, salt permeation

NO INTRODUCTION

The Earth's surface is dominated by oceans, seas, and glaciers, but geological surveys indicate only 0.8% of these water resources meet the minimum standard for freshwater adequate for human consumption.¹ Additionally, agriculture and poor irrigation practices,^{2,3} mining operations,^{4−6} industrial production, $7,8$ [an](#page-6-0)d other human activity⁹ can produce significant amounts of waste[wat](#page-6-0)er that must be tr[ea](#page-6-0)t[ed](#page-6-0) before it can be reus[ed](#page-6-0) or discharged to the enviro[nm](#page-6-0)ent without a significant ecological impact. Efficient water desalination is already vital and will continue to be to sustain the quality of life for societies living without sufficient access to freshwater.

Membrane processes, the most commonly implemented being reverse osmosis (RO),¹⁰ are playing a larger role in desalination because the energy costs per volume of freshwater produced are an order of [mag](#page-6-0)nitude lower than the costs associated with thermal desalting processes. $11,12$ In RO, water is desalted via pressurized flow past a salt-selective membrane which produces two product streams: a [des](#page-7-0)alted permeate stream and a retentate stream of concentrated brine. Although RO has been proven to be a robust process, challenges remain to fully optimize RO membranes to increase throughput and lower power consumption. These challenges may be addressed by manipulating the structure and composition of the RO membrane.^{10,13} Currently, state-of-the-art thin film composite RO membranes are comprised of two or more layers: (1) a dense poly[a](#page-6-0)[mid](#page-7-0)e selective layer deposited through interfacial

polymerization that accomplishes the separation, (2) a polysulfone or poly(ether sulfone) support layer to provide a mechanically robust, porous support for the selective layer, and optionally (3) polyester or other backing materials for mounting the RO membrane into an appropriate module.¹⁴

While a number of transport mechanisms can partially describe the flux of solvated ions and water through [t](#page-7-0)he selective layer, the exact mechanism is the subject of debate. It is generally accepted that solvated ions diffuse at a slower rate than water molecules through the polymer matrix.¹⁵ The rate of diffusion through a dense polymer layer is a function of several factors, including the hydrodynamic radius of t[he](#page-7-0) solute, the charge or lack thereof on the solute, and environmental factors such as temperature and applied pressure that affect the activity and chemical potential of the solutes.¹⁶ Note the hydrodynamic radius of a diffusing ion is not simply the radius of the ion or the polyatomic species, but also in[clu](#page-7-0)des the shell of closely hydrogen-bonded water molecules around the ion, termed a solvated ion. The size of the solvation shell is a function of the attractive electrostatic forces between the ion and the oppositely charged dipole of the water molecules and the repulsive forces between the nuclei.^{17,18} These solvation shells increase the effective size of the solute to the order of a single

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Figure 1. Spray-LbL assembly process and film architectures of (a) polymer and composite bilayer films, and (b) nanocomposite tetralayer films. Process is repeated to deposit desired number of bilayers or tetralayers.

nanometer for common monatomic ions like sodium and chloride to several nanometers for larger polyatomic ions such as sulfates, with lighter monatomic ions being relatively larger than heavier monatomic ions due to the increased charge density.19[−]²¹

There is an inverse relationship between salt rejection and water fl[ux](#page-7-0), [an](#page-7-0)d manipulating membrane properties such as the porosity of either the selective or support layer to maximize one desired attribute will reduce the other. To address this trade-off, researchers have investigated the use of nanostructured materials that enable less-hindered water transport while still providing mechanisms for salt rejection in the membrane.²² The literature on nanostructured materials for RO membranes contains experimental and theoretical work on selective lay[ers](#page-7-0) using carbon nanotubes,²³ graphene,²⁴ metal oxide nanoparticles, $25,26$ and zeolites²⁷ with varying degrees of success. Underlying these investi[ga](#page-7-0)tions is t[he](#page-7-0) hypothesis that the incorpor[ation](#page-7-0) of imperm[eab](#page-7-0)le nanomaterials into a selective layer introduces effective nanosized channels inhibiting the flux of large solutes. Particularly, the focus is on developing channels or pores with a length scale on the order of 1 nm, which enables solute rejection via size exclusion, 28 as this number is between the hydrodynamic radius of a water molecule and small solvated ions such as Na⁺ and [Cl](#page-7-0)⁻¹⁵ For this reason, we are interested in the use of laponite clay, a highaspect ratio nanoplatelet that may be layered into films t[o f](#page-7-0)orm highly tortuous diffusive pathways for solvated ions. Claycontaining ceramic membranes have been used with some success in water microfiltration applications²⁹ and oil−water separations, 30 while recent research into the use of similar composite thin films as selective layers in R[O](#page-7-0) membranes has primarily f[ocu](#page-7-0)sed on the incorporation of carbon nanotubes and zeolites.²² More generally, polymer−clay composites have unique properties that have been utilized in a number of applications [in](#page-7-0)cluding gas barriers,^{31–34} medical devices and biocompatible materials,^{35–37} and the release of chemical agents.³

Many of the assem[bly te](#page-7-0)chniques commonly used to incorp[ora](#page-7-0)te these nanomaterials into membranes, such as the inclusion of nanoparticles to the polymerization step of the selective layer or the phase inversion processing of the support layer, generate composites with a disordered internal structure that do not form uniform barrier layers. A potential improvement to this approach would be to build a more uniform selective layer through a sequential assembly process, enabling finer control over the internal structure of the selective layer. The layer-by-layer (LbL) assembly process, in which thin films are assembled via the sequential deposition of film components with complementary functionality such as electrostatic interactions, 39 is one such method to accomplish this control while enabling the construction of highly conformal thin films atop a r[ang](#page-7-0)e of porous supports. The composition of LbL films can be varied through the manipulation of process conditions such as deposition times, concentration of the adsorbing solutions, and ionic strength of the film components.^{40,41} Aerosolizing the film components with a forcing gas and spraying onto the substrate in what is termed Spray Layerby-La[yer](#page-7-0) (Spray-LbL) assembly can decrease the processing time for depositing large films by an order of magnitude⁴² and further enhance the lateral order of nanomaterials incorporated in the film. Previous research has examined the use of Lb[L](#page-7-0) films containing polyelectrolytes^{43–49} and metal-ion complexed polymers⁵⁰ to serve as selective layers in osmotic membranes. The flexibility of the Lb[L](#page-7-0) [ass](#page-7-0)embly process enables the incorpor[atio](#page-7-0)n of clay platelets into LbL-assembled films,⁵¹⁻⁵³ but there is a gap in the scientific literature concerning the use of LbL to generate nanostructured clay composite sel[ect](#page-7-0)i[ve](#page-7-0) layers for RO membranes.

In this work, we investigate several spray-LbL assembled film architectures that consist of laponite clay layers (LAP) and polyelectrolyte layers. Analogous to what is observed in models for composite polymer−clay membranes used in gas permeation applications, 54 we hypothesize alternating layers of clay intercalated with polyelectrolytes will increase the tortuosity of the diffusive path [len](#page-7-0)gth for solutes and introduce size-exclusive transport mechanisms that impact the transport of solvated ions to a greater degree than for smaller water molecules. We ground this hypothesis in three primary observations: first, prior work has shown laponite clay-containing thin films inhibit the in-plane mobility of $Li⁺$ ions by up to 2 orders of magnitude compared to the cross-plane mobility, thus serving as a barrier to ion mobility.⁵⁵ Second, incorporation of clay platelets into polymer composites was found to significantly increase selectivity in g[as](#page-7-0) barrier and separation membranes, showing that LbL-assembled clay composites serve as an effective small molecule barrier.31−³⁴ Third, recent studies on RO membranes

Figure 2. Growth curves of spray layer-by-layer assembled (a) polymer bilayer (PAH/PAA) and composite (PDAC/LAP) thin films, and (b) composite tetralayer (PAH/PAA/PAH/LAP) thin films assembled at different pH values.

containing inorganic nanoscale materials suggest the possibility of preferential water flow channels through the composite membrane.¹⁵ Through tuning of the LbL assembly conditions, films can be assembled with a relatively high degree of incorporati[on](#page-7-0) of nanomaterial into the polymer matrix, on the order of 50% of the weight fraction of the film,⁵⁶ compared to weight fractions under 10% observed in traditional polymer composites.57,58 We believe an increased qu[ant](#page-8-0)ity of nanostructured materials can further enhance selectivity in RO membranes[.](#page-8-0)

■ RESULTS AND DISCUSSION

In this work, the spray-LbL assembly technique is used to deposit composite thin films containing LAP on poly(ether sulfone) ultrafiltration membranes to generate novel RO membranes. LAP clay platelets were selected for the composite film assembly for three reasons. First, the cation-exchanged platelets have a negative surface charge and thus can be incorporated into LbL film architectures. Second, the platelets have a flat disc-like geometry with an aspect ratio of approximately $25:1^{59}$ to $30:1,^{60}$ and can be layered into films aligned with the membrane surface as shown in Figure 1. Three, LAP clay h[as](#page-8-0) a small[er p](#page-8-0)article size than other silicate clays such as montmorillonite, and thus is more compatib[le](#page-1-0) with the aerosolization technologies used in spray-LbL deposition. These features make LAP clay platelets ideal for the spray-LbL deposition of these types of clay composite thin films.

We examine two composite film architectures, the first containing the strong polycation poly(diallyldimethylammonium) (PDAC) and laponite (LAP) clay, shown in Figure 1a. For this film architecture, we control the composition of clay in the film by adjusting the spray times of the t[wo](#page-1-0) film components. For the first film, the PDAC spray time was held constant at 3 s and the LAP clay spray time was held at 3 s (3s:3s); for the second bilayer composite film, the LAP clay spray time was increased to 9 s $(3s:9s)$.

The second is a cross-linked tetralayer architecture of poly(allylamine) (PAH) and poly(acrylic acid) (PAA) bilayers between clay-containing bilayers of PAH and LAP, shown in Figure 1b. As both PAH and PAA are weak polyelectrolytes their degree of ionization depend on assembly pH. We examin[ed](#page-1-0) films assembled at pH 5 and pH 6, which yield different film properties. Both the pH 5 and pH 6 films swell under aqueous conditions, as is expected for most LbL polyelectrolyte film systems, so thermal cross-linking⁶¹ was used to form covalent bonds in the polymer matrix to lower film swelling. This additional step maintains the average [spa](#page-8-0)cing between layers of clay platelets to maintain an effective barrier

layer, while toughening the polymer matrix and increasing the overall hardness and elastic modulus of the film, a phenomenon observed in prior research on polymer−clay nanocomposite films.⁶² Finally, a cross-linked (PAH/PAA) film assembled at pH 5.0 without clay was used as a control system.

Fi[lm](#page-8-0) Thickness and Composition. The dry thicknesses of the assembled thin films were measured by profilometry. All of the examined film architectures appear to grow linearly with respect to the number of bilayers or tetralayers deposited (Figure 2a and 2b) up to a maximum of 60 repeating units. The thickness per bilayer for the (PDAC/LAP) 3s:3s composite bilayer films was approximately 2.6 nm per bilayer, and for (PDAC/LAP) 3s:9s system 5.2 nm per bilayer, approximately 84% thicker. This corresponds to an increase in the clay content of the films from 52% at 3s:3s to 83% at 3s:9s (Table 1), indicating the increase in thickness is correlated with the

Table 1. Film Clay Content As a Function of Assembly Conditions

additional incorporation of LAP into the film. The increase in clay content with longer spray times indicates that for these LAP containing film architectures, the spray-LbL deposition does not achieve full equilibrium at the film interface per layer deposition. Rather, more clay is deposited at longer exposures, which suggests kinetic control of LbL film content over these spray conditions, in contrast to the equilibrium-controlled mass content usually observed with dip-LbL with appropriately long dipping times.

The tetralayer films were assembled with spray times of 3 s maintained for each film component but at three assembly pH values. At pH 6.0, the PAH and PAA are both highly ionized when deposited at the film interface based on their respective pK_a values (PAA pK_a = 6.0−6.5; PAH pK_a = 8.0−9.0⁶³), which results in very thin, dense polymer layers being deposited. On average, the dry thickness for (PAH/PAA/PAH/LA[P\)](#page-8-0) pH 6.0 films was 2.6 nm per tetralayer, similar to what was observed for the (PDAC/LAP) 3s:3s assembly conditions (Figure 2b). The clay content of these two films were also statistically indistinguishable (Table 1), indicating the polymeric matrices of the (PDAC/LAP) 3s:3s and (PAH/PAA/PAH/LAP) pH 6.0 films were of similar dry film thickness.

Figure 3. ATR-FTIR spectrograph showing the untreated films (solid line) and the films cross-linked at 175 °C for 5 h (dashed line) for (a) (PAH/ PAA/PAH/LAP) pH 5.0 films, and for (b) (PAH/PAA/PAH/LAP) pH 6.0 films.

The (PAH/PAA/PAH/LAP) pH 5.0 and pH 5.5 films assembled were significantly thicker than the pH 6.0 films of the same architecture, with a thickness of approximately 5.1 and 4.8 nm per tetralayer deposited respectively (Figure 2b). At pH 5.0 assembly, the PAA chains are less ionized, resulting in the deposition of thicker polyelectrolyte layers betwe[en](#page-2-0) the clay layers, decreasing the overall clay content of the film to approximately 38% (Table 1). This is a similar trend to what is observed in the literature for dipped (PAH/PAA) bilayer films over the same pH range.⁴⁰ Within the margin of error, the thickness of the tetralayer [p](#page-2-0)H 5.0 film appears to be a linear sum of the LAP-containin[g b](#page-7-0)ilayer and the (PAH/PAA) pH 5.0 bilayer, implying the intercalating clay layers do not have a significant impact on the dynamics on the polyelectrolyte deposition, unlike what has been observed for other LbL polymer-clay nanocomposite films.⁶⁴ There are two contributing factors that explain this observation. First, the smaller LAP clay platelets form more uniform [su](#page-8-0)rface layers during each deposition cycle. Second, the short deposition times employed by the spray-LbL assembly process tend to generate linearly growing films, in contrast to the exponential growth patterns observed with weak polyelectrolytes assembled via dip-LbL.

Thermal Cross-Linking. The tetralayer composite films were cross-linked to form amide bonds via uncatalyzed heating in a 175 °C oven at atmospheric pressure for 5 h. The resulting covalent bond formation was characterized by ATR-FTIR spectroscopy (Figure 3). Qualitatively, the 1550−1540 cm⁻¹ peak corresponds to the $N-H$ bending in the amine and is significantly reduced after cross-linking, indicating the reaction of primary amines to form secondary amides. Through the application of ORIGIN software for linear baseline adjustment, peak deconvolution, and peak integration, an estimate of roughly 20.6% of the amine groups were cross-linked for the pH 5.0 films (Figure 3a), compared to 12.8% for pH 6.0 films (Figure 3b). The increase in the degree of cross-linking for the pH 5.0 films is attributable to the increased number of free, unbounded acid functional groups present in the pH 5.0 films that are available at the lower pH of assembly. An additional side reaction possible with these free carboxylic acid groups is the formation of acid anhydrides. The formation of acid anhydride bonds was not observed; the key peaks for the identification of these bonds are in the 1830−1800 cm[−]¹ and $1775-1740$ cm⁻¹ range, neither of which were appreciable in the spectra.

To show the impact of the cross-linking under aqueous conditions, spectroscopic ellipsometry was used on 20 bilayer and tetralayer samples to evaluate the degree of swelling in both in the untreated and cross-linked state (Table 2). The uncrosslinked (PAH/PAA) pH 5.0 films were observed to significantly

Table 2. Swelling in Untreated and Cross-Linked Films

swell as the films undergo a significant pH shift from the assembly conditions as well as have more free amine and carboxylic acid groups present to form hydrogen bonds with water molecules. The composite bilayer of (PAH/LAP) swelled approximately 62% upon exposure to DI water, significantly less than the polyelectrolyte (PAH/PAA) pH 5.0 films. Crosslinking reduced the observed film swelling to roughly 10−15% regardless of assembly pH. This suggests that the covalent bonds formed during the cross-linking process both reduce the number of free amine and carboxylic acid groups that can interact with water and physically constrain the swelling of the film.

Imaging of Composite Films. The uncoated substrate, a poly(ether sulfone) (PES) ultrafiltration membrane, has a regular distribution of surface pores of approximately 30 nm diameter (Figure 4a). Deposited film components, such as $(PAH/PAA/PAH/LAP)_{40}$ film architecture shown in Figure 4b, bridge the und[er](#page-4-0)lying pore structure to form thin, relatively smooth surface films. These films do not appear to penetrate [in](#page-4-0)to the underlying porous structure of the PES membrane, as observed by comparing the cross-sectional SEM micrographs for the uncoated membranes in Figure 4c and a (PAH/PAA/ $PAH/LAP)_{40}$ -coated membrane in Figure 4d. Clay platelets with intercalating polymer regions we[re](#page-4-0) observed by TEM imaging of the cross sections of these me[mb](#page-4-0)ranes, shown in Figure 4e. Through ImageJ software analysis, the approximate size of the clay platelets observed in the cross-sectional SEM was on [th](#page-4-0)e order of 50 nm, indicating the clay platelets are not aggregated within the film and are intercalated within the polymer matrix. There was no statistically significant difference apparent in the average platelet spacing for the uncross-linked pH 5.0 films $(2.41 \pm 0.23 \text{ nm})$ and the cross-linked films $(2.57$ $± 0.39$ nm).

Micrographs were also taken of the membranes as assembled and after water and salt permeation experiments were conducted (Figure 5). Significant defects formed in (PDAC/ LAP)₄₀ and $(PDAC/LAP)_{60}$ films following the permeation trials with applied p[re](#page-4-0)ssures in excess of 100 psig (Figure 5a,b). Qualitatively, it would appear that matrix of the PDAC and LAP film formed during the LbL deposition process do[es](#page-4-0) not have the mechanical cohesiveness and integrity to withstand the

Figure 4. (a) SEM micrograph of uncoated PES ultrafiltration membrane with 30 nm pores; (b) SEM micrograph of $(PAH/PAA/PAH/LAP)_{40}$ pH 5.0 composite tetralayer film on PES membrane; (c) cross-sectional SEM micrograph of uncoated PES membrane; (d) cross-sectional SEM micrograph of (PAH/PAA/PAH/LAP)40 pH 5.0 composite tetralayer film on PES membrane; and (e) TEM micrograph of (PAH/PAA/PAH/ $LAP)_{40}$ pH 5.0 composite tetralayer film deposited on Si wafer.

Figure 5. (a) SEM micrograph of $(PDAC/LAP)_{40}$ 3s:3s film as prepared before permeation experiments; (b) SEM micrograph of $(PDAC/LAP)_{40}$ 3s:3s film after water and salt permeation experiments; (c) SEM micrograph of $(PAH/PAA/PAH/LAP)_{40}$ pH 5.0 film as prepared before permeation experiments; and (d) SEM micrograph of (PAH/PAA/PAH/LAP)₄₀ pH 5.0 after water and salt permeation experiments.

in situ RO conditions. The cross-linked (PAH/PAA/PAH/ LAP)₄₀ and (PAH/PAA/PAH/LAP)₆₀ films, however, do not exhibit such defects after permeation trials (Figure 5c,d), indicating the covalently bonded LbL film is more mechanically robust under RO conditions. Salt deposits in the form of micron-sized salt crystals were observed on the surfaces of LbL films following the permeation trials, however these crystals appear independently of the fractures in the (PDAC/LAP) films and are the result of the washing and drying of the films in the sample preparation for the SEM.

Permeation Properties of LbL Films. Water flux and salt rejection data on the thin films were collected from dead-end permeation cell measurements over a pressure range of 50 psig to 250 psig with 10 000 ppm of NaCl solution (Figure 6a,b, respectively). A localized permeability coefficient was calculated through a regression of the Spiegler−Kedem model to yi[el](#page-5-0)d a single value to compare films where several different transport mechanisms are at work. The water permeability of the commercial TFC-HR polyamide active layer under these operating conditions is 7.6×10^{-15} m²/Pa s with 98% salt

rejection. All the examined LbL films exhibited significantly greater water permeability, on the order of one to 2 orders of magnitude.

Due to the formation of defects in the (PDAC/LAP) selective layers, the salt rejection was low: 28% for the 3s:3s film and 10% for the 3s:9s film. The water permeability for the 3s:3s film was 2.27×10^{-13} m²/Pa s and for the 3s:9s film 8.96 \times 10⁻¹³ m²/Pa s, roughly four times greater. It is likely that the lower salt rejection and increased water flux correspond to an increase in the cracks formed in the (PDAC/LAP) films during operation (Figure 5). However, this phenomenon appears to have been eliminated in the tetralayer composite films, which exhibit similar water permeability but significantly increased salt rejection.

It is notable that for the composite tetralayer series from pH 6.0 to 5.0, the water permeability dropped by approximately a factor of 2 from 4.60×10^{-13} m²/Pa s to 2.82×10^{-13} m²/Pa s. However, the salt rejection increased from an average of 46% to 89%. We suspect this occurs for two reasons: (1) the increased degree of cross-linking in pH 5.0 films reduces the effective averaged channel width between clay layers after swelling under in situ RO conditions, and (2) additional free carboxylic acid groups present in the pH 5.0 film slow the diffusion of solvated ions due to charge effects. The ultimate impact of these two phenomena is an increase in tortuosity of the selective layer, resulting in longer diffusive path lengths for solvated ions.

It is also notable that the overall clay content of the film does not directly correlate to its performance in the salt rejection trials. The best performing film, the (PAH/PAA/PAH/LAP) pH 5.0 film, had an average clay content of 38% and a salt rejection of 89%. Extremely high clay content films assembled via the (PDAC/LAP) method perform similarly with regards to water permeation, but were more brittle due to the high clay content, and thus more susceptible to critical defect formation during RO, with a 83% clay membrane only rejecting 10% of salt ions. This demonstrates the importance of the cross-linked polymeric matrix that reduces the brittleness of the film; these cross-links fortify the polymeric matrix encompassing the clay platelets, and prevent the formation of cracking and critical faults under in situ RO conditions. The film architecture without clay, the (PAH/PAA) bilayer architecture, only exhibited 53% salt rejection. This is indicative of a swollen polymeric matrix that does not serve as an effective barrier to

layer-by-layer thin films against commercially available Koch TFC-HR RO membrane, and (c) prior data plotted on two-axis graph of water permeability against salt rejection.

solvated ion diffusion due to the lack of size exclusion-driven rejection through the nanochannels present in the clay composite films. To better illustrate the trade-off between increased water permeability and salt rejection, the salt and water permeation data is plotted against each other (Figure 6c).

■ CONCLUSIONS

In this report, we have demonstrated the effectiveness of using LAP clay in spray-LbL assembled selective layers for RO membranes in both a strong polyelectrolyte film architecture, (PDAC/LAP), and a weak polyelectrolyte film architecture, (PAH/PAA/PAH/LAP). The physical and permeability properties of these two architectures were characterized and compared. The calculated water permeability through the Spiegler−Kedem model for all LbL-assembled composite thin films was between one and 2 orders of magnitude greater than what was observed for commercially available thin film composite RO membranes. The salt rejection measured for uncross-linked bilayer (PDAC/LAP) films was between 10% and 28% and for the cross-linked tetralayer (PAH/PAA/PAH/ LAP) films was between 46% and 89%, with the greatest salt rejection observed for films assembled at pH 5.0.

The (PDAC/LAP) composite bilayer film architectures were too brittle under in situ RO conditions and formed critical defects during operation. These defects were effectively eliminated by the introduction of a cross-linkable polymer matrix of PAH and PAA that reduced brittleness and film swelling under aqueous conditions, which made the composite films more mechanically tough and maintained the nanochannels between platelet layers. We hypothesize that the selective salt rejection and high water permeability is the result of a combination of two transport mechanisms: a size-exclusion

transport mechanism that hinders the flow of solvated ions between clay layers to a greater degree than individual water molecules, and charge interactions between the polyelectrolytes and the solvated ions in the selective layer. We attribute the efficacy of the pH 5.0 tetralayer architecture over the pH 6.0 to the increased degree of cross-linking via free carboxylic acid groups which strengthens the polymeric matrix, as well as the enhanced presence of free carboxylic acid groups that may act as hydrated regions to further enhance water permeability while retarding ion transport.

Interestingly, although the (PDAC/LAP) films have higher clay content than the tetralayer (PAH/PAA/PAH/LAP) films and naively one would predict a greater degree of tortuosity for these films leading to an increase in salt selectivity, a more complicated trend was observed. This was due to the high rate of defect formation observed for the (PDAC/LAP) bilayer films under RO conditions, which was reduced through increasing the polymer content of the film and forming a strong polymeric matrix through thermal cross-linking. However, given the polymer (PAH/PAA) bilayer films exhibited 53% salt rejection, there is necessarily an optimal value for maximizing salt rejection as a function of clay content. Of the film architectures investigated, the best performing film was tetralayer pH 5.0 film with an average clay content of 38%, a degree of incorporation that is difficult to achieve with other assembly methods but is attainable through spray-LbL assembly.

These findings represent an opportunity for further development of LbL-assembled clay composite formulations that can be adapted to water filtration applications. Furthermore, if the fouling properties were characterized or dedicated antifouling layers were introduced, then these high-flux clay composite

EXPERIMENTAL SECTION

Materials. Laponite clay was provided by Southern Clay Products; clay dispersions were prepared at a concentration of 1.0 wt % clay and the balance reagent-grade water. Poly(diallyldimethylammonium chloride) (MW: 200−350 kDa) was obtained from Sigma-Aldrich, and both poly(allylamine hydrochloride) (MW: 60 kDa) and poly(acrylic acid) (MW: 20 kDa) were obtained from Polysciences, Inc. Polyelectrolyte solutions were prepared at 10 mM concentration and were adjusted to the assembly pH by using a Φ340 pH/Temp Meter and concentrated HCl or NaOH solution as appropriate. Millipore PES ultrafiltration membranes with 30 nm pores were purchased and used as a substrate for deposition. PES membranes were plasma-cleaned in a Harrick Plasma Cleaner/Sterilizer PDC-32G at 18 W for 30 s and soaked in PDAC or PAH solutions before spray-LbL film assembly.

Spray Layer-by-Layer (Spray-LbL) Deposition. Films were constructed using a custom-built spraying apparatus. Solutions and clay dispersions were aerosolized with N_2 gas at 20 psi and are sprayed onto the substrate rotated at 10 rpm. The basic program for each layer involved spraying the film component for 3 s, pausing for a 5 s drain period, rinsing for 10 s with pH-adjusted water, and then and then a final 5 s drain period. The sequence is repeated for each film component listed to assemble a bilayer or tetralayer. Films assembled at different component spray times are identified by the expression ns:ms, where n refers to the spray time of PDAC, and m refers to the spray time of LAP.

Characterization. A Dektak 150 profilometer was used to determine the film thickness. Profilometry samples were deposited on glass slides plasma-cleaned using the above equipment for 5 min; otherwise, the standard protocol above was used. Both a JEOL JSM-6060 and a JSM-6010LA Scanning Electron Microscopes (SEM) were used to image both film surfaces and cross sections. Cross-sectional SEM samples were prepared via the cryo-fracture method by submerging the sample in liquid N2 and then physically separated. A J.A. Woollam Co., Inc. M-2000D spectroscopic ellipsometer was used to determine swelling through the measured change in film thickness between the dry state and being immersed in DI water for films deposited on Si wafers. The spectra were fitted with a (1) Si model, (2) SiO₂ model, and (3) Cauchy model corresponding to (1) the bulk of the Si wafer, (2) the oxidized surface of the Si wafer, and (3) the LbL-deposited thin film. A TA Instruments Discovery Series Thermogravimetric Analyzer was used to determine the film composition of LbL films (150−200 bilayer/tetralayer depositions on polystyrene chips) with the following program: temperature equilibration step at 65 °C for 5 min, followed by a ramp up to 800 °C at the rate of 10 °C/min, followed by a final temperature equilibration step at 800 °C for 5 min. The percentage clay content was calculated by taking the final mass of the remaining film after thermal decomposition and dividing by the mass taken after the temperature equilibriation step at 65 °C. A Sterlitech HP4750 dead-end permeation cell was used to determine both water permeability and salt rejection. The cell was operated with an applied pressure between 50 and 300 psi and on a Benchmark H4000−S-E Magnetic Stirrer at speed "8" for films assembled on ultrafiltration membranes. A Spiegler−Kedem model was applied to determine the local water permeability constants. The chloride ion concentration of the collected permeate was measured with an Oakton Ion 700 conductivity meter and Thermo-Scientific Orion 9617BNWP IonPlus Probe.

■ ASSOCIATED CONTENT

6 Supporting Information

The method for calculating water permeability via the Spiegler−Kedem method, a pore size distribution analysis on the PES-30 nm UF membranes, and nanoindentation testing on a selection of films from this report.The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsami.5b01879.

[■](http://pubs.acs.org) AUTHOR I[NFORMATION](http://pubs.acs.org/doi/abs/10.1021/acsami.5b01879)

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Notes

The authors declare no competing financial interest.

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■ ABBREVIATIONS

LAP, laponite clay LbL, layer-by-layer PAA, poly(acrylic acid) PAH, poly(allylamine) PDAC, poly(diallyldimethylammonium chloride) PES, poly(ethersulfone) RO, reverse osmosis spray-LbL, spray layer-by-layer

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