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Shinde, S., Incel, A., Mansour, M., Olsson, G., Nicholls, I., Urraca, J., & Sellergren, B. (2020). Urea-Based Imprinted Polymer Hosts with Switchable Anion Preference. Journal of the American Chemical Society (JACS), 142(26), 11404-11416. https://doi.org/10.1021/jacs.0c00707

Published in:
Journal of the American Chemical Society (JACS)

Document Version:
Publisher's PDF, also known as Version of record

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Urea-Based Imprinted Polymer Hosts with Switchable Anion Preference

Sudhirkumar Shinde, Anil Incel, Mona Mansour, Gustaf D. Olsson, Ian A. Nicholls, Cem Esen, Javier Urraca, and Börje Sellergren*  

ABSTRACT: The design of artificial oxyanion receptors with switchable ion preference is a challenging goal in host–guest chemistry. We here report on molecularly imprinted polymers (MIPs) with an external phospho-sulpho switch driven by small molecule modifiers. The polymers were prepared by hydrogen bond-mediated imprinting of the mono- or dianions of phenyl phosphonic acid (PPA), phenyl sulfonic acid (PSA), and benzoic acid (BA) using N-3,5-bis-(trifluoromethyl)-phenyl-N-4-vinyl-phenyl urea (1) as the functional host monomer. The interaction mode between the functional monomer and the monoanions was elucidated by 1H NMR titrations and 1H−1H NMR NOESY supported by molecular dynamic simulation, which confirmed the presence of high-order complexes. PPA imprinted polymers bound PPA with an equilibrium constant $K_{eq} = 1.8 \times 10^5 \text{M}^{-1}$ in acetonitrile (0.1% 1,2,2,6,6-pentamethylpiperidine) and inorganic HPO$_4^{2−}$ and SO$_4^{2−}$ with $K_{eq} = 2.9 \times 10^3 \text{M}^{-1}$ and $4.5 \times 10^3 \text{M}^{-1}$, respectively, in aqueous buffer. Moreover, the chromatographic retentivity of phosphonate versus sulfonate was shown to be completely switched on this polymer when changing from a basic to an acidic modifier. Mechanistic insights into this system were obtained from kinetic investigations and DSC, MALDI-TOF-MS, 1H NMR-studies of linear polymers prepared in the presence of template. The results suggest the formation of template induced 1−1 diad repeats in the polymer main chain shedding unique light on the relative contributions of configurational and conformational imprinting.

INTRODUCTION

The molecular constituents of a living cell are predominantly water-soluble molecules carrying a net negative charge, e.g., cofactors, enzyme substrates, nucleic acids, the lipid double layer, and the glyocalix. As a consequence, evolution has resulted in highly refined receptors capable of recognizing anions in water. Some of the most impressive examples are the proteins designed to selectively bind and/or transport either sulfate or phosphate ions. These two pyramidal anions resulted in highly reactivity especially with respect to the relative preference for phosphate/sulfate. Moreover, these anion receptors are seldomly capable of switching the anion preference or of releasing the guest upon external stimuli. Inspired by this remarkable performance, much effort has been devoted to the development of neutral sulfate/phosphate binding hosts. In contrast to the more common approach based on charged receptors, neutral hosts can achieve higher selectivity but are challenging to construct since H-bonding is considerably weaker in polar solvents. Nevertheless, both macrocyclic e.g. pyrrole, saphyrine, peptide and acyclic podand hosts e.g. ureas, have been reported showing high sulfate affinity. However, these hosts rarely overturn the Hofmeister solvation governed series of anion affinity especially with respect to the relative preference for phosphate/sulfate. Moreover, these anion receptors are seldomly capable of switching the anion preference or of binding and releasing the guest upon external stimuli.

Received: January 23, 2020  
Published: May 19, 2020
This highlights a need of neutral hosts with tunable microenvironment capable of alternating phosphate/sulfate recognition. This can in principle be achieved by embedding designed low molecular weight hosts in a polymer scaffold imparting a lower dielectric microenvironment. Alternatively, a macromolecular host can be designed from simpler building blocks by molecular imprinting. Monomers are here chosen or designed to complement functional groups of a template molecule. Polymerizing the monomer complexes into a cross-linked polymer matrix followed by removal of the template, leaves behind sites capable of recognizing the template. This approach benefits from spontaneous self-assembly guiding the binding groups to their positions in the receptor site. Thus, the structure of the final binding site is a priori unknown.

Notable examples of MIPs for anion recognition in water have been reported using both charged, e.g., imidazolium, amidinium, and neutral host monomers, e.g., urea, thiourea, squaramide, and others. Some time ago we introduced urea based host monomers acting as potent H-bond donors for stoichiometric imprinting of oxyanions or other H-bond acceptors. Receptors for β-lactam antibiotics, phospho-peptides, peptide biomarkers, phospholipids, sugar acids, and sulfated peptides were demonstrated.

To gain insight into the nature of these imprinted receptors at a molecular level we here report an in-depth investigation of MIPs for simple oxyanions (Scheme 1).

### Scheme 1. Plausible Structural Formulas of H-Bonded Complex between Oxyanions and 1,3-Diaryl Urea Monomer 1 (left) and Structures of Oxyacids Used in the Study (Right)\(^a\)

\(\text{PO}_4^{3-}, \text{SO}_4^{2-}, \text{PO}_4^{3-}\)

\(\text{PPA} = \text{phenylphosphonic acid}; \text{PSA} = \text{phenylsulfonic acid}; \text{BA} = \text{benzoic acid}; \text{PPPrA} = \text{phenylphosphoric acid.}\)

With this we aimed to answer the following questions: (1) What level of oxyanion selectivity and affinity can be achieved with MIPs in organic and aqueous media? (2) Can MIPs recognize inorganic anions in water? (3) Can the Hofmeister preference be overturned? (4) Can we switch oxyanion preference using the same MIP? (5) What are the mechanisms underlying the imprinting process?

On the basis of combined physical characterization, modeling, spectroscopic studies of prepolymerization complexes, and studies of the polymerization kinetics we here offer answers to the aforementioned questions.

### RESULTS AND DISCUSSION

#### Monomer/Template Complex Formation.

1,3-Disubstituted ureas have long been exploited as neutral hosts (H) for complexing oxyanion guests (G), \(^3\), \(^38\), \(^39\). They provide a 2-fold H-bond donor which with the acceptor (e.g, carboxylate, phosphate, sulfate) results in a cyclic H-bonded complex. Also, several structural and compositional parameters can be readily tuned to enhance HG association strengths (Scheme 2).

#### Scheme 2. Structural and Compositional Parameters for Tuning Host-Guest Association Strength\(^a\)

| Parameter                              | Calculated Values |
|----------------------------------------|-------------------|
| pH (6)                                  | –498 kJ/mol       |
| pH (6)                                  | –1130 kJ/mol      |

\(^a\)EWG = electron withdrawing group.

The affinity for the guest increases with the acidity of the urea protons (donor capacity, \(R_3, R_4, R_5 = \text{electron withdrawing groups (EWGs)}\)) and the basicity of the oxyanion (acceptor capacity), but this can be offset by the ability of the host to self-associate and host deprotonation.\(^40\) The latter leads to nondirectional electrostatic interactions and is promoted by increased solvent polarity.\(^41\) This is an undesired effect since it undermines the structural integrity of the hydrogen bond connected HG complex—hence a compromise between host acidity and guest basicity has to be found that maximizes the complex strength. In this context, no deprotonation has so far been observed for the host monomer, 1, used in the current study (Scheme 1). The nature of the countercation also influences the HG complex stability. Complexation can be easily induced by proton transfer to amine bases but this causes undesirable competition between the urea host and the protonated amine for binding the guest. Stronger complexation is therefore induced by the use of bulky tertiary amines (e.g., pentamethylypiperidine, PMP) or quaternary ammonium ions like tetrabutylammonium (TBA), which have been shown to produce pronounced counterion memory effects.\(^29\),\(^42\) Another factor is the level of host preorganization, which impacts on the size of the entropic penalty upon guest complexation due to the need to freeze internal rotors. For monourrea host monomers this can in principle be enhanced by introducing two polymerizable groups (cross-linkers) in the host thereby

### Table 1. Physical Properties of Sulfate and Phosphate Anions

| Anion   | \(pK_a, 1^{\circ}\) | \(pK_a, 2^{\circ}\) | \(pK_a, 3^{\circ}\) | Molecular Volume | Anion Hydration Energy |
|---------|-------------------|-------------------|-------------------|----------------|-----------------------|
| \(\text{PO}_4^{3-}\) | 2.1               | 7.2               | 10.9              | 56             | –498 |
| \(\text{SO}_4^{2-}\) | –3                | 1.9               |                   | 51             | –1130 |

\(^{1}\)Calculated values assuming pH = 6 and an average charge of –2 for sulfate and –1.4 for phosphate.\(^3\)

\(^{2}\)Calculated values assuming pH = 6 and an average charge of –1.4 for phosphate.\(^3\)

\(^{3}\)Calculated values assuming pH = 6 and an average charge of –2 for sulfate and –1.4 for phosphate.\(^3\)

\(^{4}\)Calculated values assuming pH = 6 and an average charge of –1.4 for phosphate.\(^3\)

\(^{5}\)Calculated values assuming pH = 6 and an average charge of –2 for sulfate and –1.4 for phosphate.\(^3\)

With this work we aimed to answer the following questions: (1) What level of oxyanion selectivity and affinity can be achieved with MIPs in organic and aqueous media? (2) Can MIPs recognize inorganic anions in water? (3) Can the Hofmeister preference be overturned? (4) Can we switch oxyanion preference using the same MIP? (5) What are the mechanisms underlying the imprinting process?

On the basis of combined physical characterization, modeling, spectroscopic studies of prepolymerization complexes, and studies of the polymerization kinetics we here offer answers to the aforementioned questions.
increasing host stiffness (R = vinyl in Scheme 2). For ternary HG complexes formed from phosphate dianions and urea hosts (2:1 complex in Scheme 3), host preorganization has been enhanced by introducing the cleft-like host monomer 2 (1:1 complex in Scheme 3).

Scheme 3. Anticipated Ternary or Binary Monomer—Template Complexes Formed Using Mono- (Left) Or Di- (Right) Functional Urea Host Monomers

Considering oxyanion—oxourea complexation, the stability has been shown to increase in the order of increasing acceptor basicity, i.e., PhSO3− < PhP(OH)O2− < PhP(OH)O2− ≤ PhCOO− < PhPO3− ≤ PhPO3−. To verify this trend we decided to investigate 1 with respect to its complex stability with model anions in the form of TBA-salts of PPA, PPrA, BA, and PSA. Interaction strength and complex stoichiometries were investigated through 1H NMR titration or isothermal titration calorimetry (ITC) (see the Supporting Information (SI)). Titrating TBA in DMSO-d6 and PPA-2PMP in THF-d8 resulted in strong downfield shifts of urea protons Ha and Hb and more moderate shifts of the aromatic protons Hc-f (Figure S1). The moderate downfield shift of Hf are in agreement with previous NMR titration study of 1,3-bis(4-nitrophenyl)urea with acetate. 1H NMR studies using Jobs method of continuous variation confirmed a preferred 1:1 stoichiometry. Complex stability constants were calculated using the induced downfield shifts of the urea protons of 1 (Figure S1). With respect to monoanions, 1 formed more stable complexes with benzoate anion BA (Keq = 8820 M−1) followed by PPA (Keq = 7005 M−1) and PPrA (Keq = 676 M−1), whereas the complex with the sulfonate PSA was too weak to be detected (Table S1). This agrees with the trend determined by Kelly et al. Comparing with monoanions, the dianions of PPA and PPrA interact more strongly with 1. The relative affinity of the two dianions is reflected in the downfield shifts of Ha and Hb (Figures 1 and 2) where PPrA (Figure 1c) display larger shifts than PPA (Figure 1b), again in line with the literature results.

To gain more insight we performed molecular dynamics (MD) simulations of the prepolymerization mixtures. MD simulations offer unique insights into the nature of the interactions between polymerization mixture components providing diagnostic/prognostic correlation to polymer performance. Compositions of simulated and evaluated systems and chemical structures are presented in Table 2 and Figure S15.

Full system all-atom simulations were performed using the same stoichiometries employed for polymer synthesis. The results overall confirmed the presence of stable interactions between the neutral, mono-, and di-valent anions of PPA and 1 (Tables 3 and S8–S10 and Figure S15) with the dianion of PPA (PP2) (PPrA not investigated using MD) displaying a higher degree of H-bonding than the monoanion (PP1) or neutral PPA template.

However, the monoanionic PSA template (PS1) performs as well, even slightly better than PP2 considering complex formation with 1 (Tables S8–S10). A rank order of affinity for 1 was observed where anionic neutral PSA < neutral PPA < anionic PP1 < dianionic PP2 < PS1 (Tables 3 and S8–S10). This obviously disagrees with the NMR results where no interactions between PS1 and 1 could be detected. The systems differ though with respect to solvent polarity, the base, presence of cross-linker, and the concentration of all species which we believe can account for these contrasting results (vide infra). 1H NMR chemical shift changes of urea host monomer 1 complexation with PPA-2PMP or PPrA-2PMP. ∆δ = δ (in the presence of anion) − δ (in the absence of anions), induced by addition of 1/2 equiv of different anions to receptor 1.

To gain further insight into the nature of the urea complexes between 1 and the two anions we performed 2D-NOESY NMR experiments and molecular modeling as reported by others. As shown in Figure 3a the free 1 shows strong cross-peaks with the residual water signal at ca. 2.9 ppm indicating water protons close in space to the urea protons. In addition, weaker cross-peaks are observed between the two-urea protons Ha and Hb resulting from the Z,Z-urea conformer with the two protons juxtaposed (Figure 3e). The addition of half an equivalent of PPA-2PMP (Figure 3b) and PPrA-2PMP (Figure 3c) shows extensive cross-peaks with the residual water signal at ca. 2.9 ppm indicating cross-links of water protons close in space to the urea protons and proton exchange reactions. The cross-peaks between the two-urea protons Ha and Hb in the presence of 1/2 equiv of anionic PSA template (PS1) at 5 ppm is a direct consequence of cross-linker formation where the water protons close in space to the urea protons exchange with the residual water signal at ca. 2.9 ppm but interestingly the cross-peaks between the two-urea protons Ha and Hb in the presence of 1/2 equiv of dianionic PP2 template (PP2) is broader at 10 ppm suggesting the ura complexes between 1 and anionic 2PP2 are less stable. The presence of cross-links of water protons close in space to the urea protons exchange with the residual water signal at ca. 2.9 ppm but interestingly the cross-peaks between the two-urea protons Ha and Hb in the presence of 1/2 equiv of dianionic PP2 template (PP2) is broader at 10 ppm suggesting the ura complexes between 1 and anionic 2PP2 are less stable.
ammonium phosphonate salt was poorly soluble in this solvent, the solution became completely clear after addition of 2 equiv of 1. This provides unequivocal visual proof for the presence of prepolymerization complexes between 1 and PPA-2PMP.

Polymer preparation and characterization. Imprinted and nonimprinted polymers were thereafter prepared and characterized (see SI and Tables S2–S4) using the urea host monomer 1 in a 2:1 and 1:1 stoichiometric ratio to the templates: PPA, PSA, or BA in their monoanionic or dianionic forms (Scheme 4). Nonimprinted polymers \( P_{N} \) were prepared identically to the imprinted polymers though in the absence of template. Figure S2 shows the 25–36 \( \mu \)m particle fraction obtained after crushing and sieving of the polymer monoliths and template removal by solvent extraction. These were subjected to physical characterization as follows: C, H, and N elemental analysis of imprinted and nonimprinted polymers after removal of template matched calculated values after correction for water uptake. This is reflected in the agreement between calculated and found N/C ratios (Table S3). Thermogravimetric analysis (TGA) of the polymers produced weight loss curves typical for highly cross-linked networks with onset above 200 °C (Figure S3) and complete weight loss obtained at 450 °C. Meanwhile, the transmission FTIR spectra (Figure S4) showed all characteristic bands with

| Table 2. Composition of Systems Simulated and Evaluated\( ^{ac} \) |
|-----------------|--------------------|-----------------|--------------------|--------------------|--------------------|
| \( T^{b} \)     | P1N                | P1C             | P2N                | P2C                | P3N                |
| PPA             | 2PP1               | PPA             | PP2                | PSA                | PS1                |
| PMP\( ^{b} \)   | 20                 | 40              | 40                 | 20                 | 40                 |
| PMPH\( ^{b} \)  |                    |                 |                    |                    |                    |
| THF             | 2760               | 2760            | 2760               | 2760               | 2760               |

\( ^{a} \)Numbers of virtual molecular models of each molecule included in performed simulations of systems. See Figure S15 for molecular representations and defined abbreviations. \( ^{b} T = \) template (20 equiv), PP1 = PPA monoanion, PP2 = PPA dianion, PS1 = PSA monoanion, PMP = 1,2,6,6-pentamethylpiperidine, EGDMA = ethyleneglycoldimethacrylate. \( ^{c} \)PN/C, Polymer system X, neutral (N) or charged (C).

| Table 3. H-Bond Analysis Results\( ^{a} \) |
|-----------------|-----------------|-----------------|-----------------|-----------------|
| \( T^{b} \)     | PMP             | PMPH\( ^{b} \)  | 1               | EGDMA          |
| P1C             | 0.00            | 0.00            | 0.11            | 0.00            |
| PMP\( ^{b} \)   | 64.54           | 0.00            | 0.00            | 0.00            |
| P1N             | 24.26           | 0.00            | 0.11            | 18.27           |
| P2C             | 77.27           | 0.00            | 0.04            | 46.69           |
| P2N             | 13.42           | 0.00            | 0.10            | 25.93           |
| P3C             | 33.57           | 0.00            | 0.09            | 77.59           |
| P3N             | 74.37           | 0.00            | 0.27            | 15.11           |
| P4C             | 38.96           | 0.00            | 0.00            | 14.66           |
| P4N             | 47.39           | 0.00            | 0.50            | 48.62           |
| P1N             | 1.00            | 0.00            | 0.00            | 11.84           |
| P2N             | 1.06            | 0.00            | 0.19            | 29.18           |
| P3C             | 1.50            | 0.00            | 0.05            | 24.26           |
| P4C             | 1.50            | 0.00            | 0.36            | 15.11           |

\( ^{a} \)Summarized average interactions, as percent of total simulation time, between all atom pairs of indicated molecular species. See Figure S15 for molecular representations and defined abbreviations. All analyses involving templates were averaged against the number of templates (20 in each system). All interactions including the urea monomer I but not the template was averaged against the number of urea monomer molecules in each system (see Table 2). Cross-linker interactions with solvent and bases were averaged against the number of cross-linkers (800 in all systems) and, finally, base self-interaction and solvent interaction were averaged against the number of base molecules in each system (20 in P1, P3 N/C systems and P4C, 40 in P2 N/C and P4N systems). \( ^{b} \)Interactions with the template in the analyzed system.

The former reflects stabilization of the Z,Z-conformer, whereas the latter a close contact between I and PMP, this providing an indirect support for the existence of the anticipated ternary complex between I, PMP and the dianions (Scheme 4) This is in agreement with previous reports on urea- and squaramide-based sulfate receptors. Results from MD simulations also support the argument for higher order complexes observed in cameo images from the production run (Figures S16 and S17) and in the radial distribution function (RDF) analyses (Table 4).

As expected, the degree of complexation increases with the template charge, e.g., P2C (\( T = \) PPA dianion) display clusters involving any part of 6 molecules of I within 5 Å from a template molecule, whereas only half of this number is seen in P1C. Generally, PMPH\( ^{+} \) was found in the simulations to be in close proximity to the templates and engaged in extensive H-bonding with charged templates (Table 3). These results support the observed solubilization of the template PPA-2PMP in the presence of I in THF (Scheme 4). Whereas the ammonium phosphonate salt was poorly soluble in this solvent, the solution became completely clear after addition of 2 equiv of I. This provides unequivocal visual proof for the presence of prepolymerization complexes between 1 and PPA-2PMP.

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no apparent difference between imprinted and nonimprinted polymers all in all, indicating a stoichiometric monomer incorporation and successful template removal from the imprinted polymers.

**Figure 3.** $^1$H−$^1$H NMR NOESY spectra of 1: (a) and (b) (close-up), (c) 2:1 ratio of 1:PPA·2PMP and (d) 2:1 ratio of 1:PPrA·2PMP in THF-$_d_8$ and (e) anion induced conformational stabilization detected by an increase $^1$H−$^1$H NMR NOESY between Ha and Hb. [1] = 30 mM. The arrows indicate the urea−water (a) and urea−Ha,Hb cross peaks (b−d).

**Scheme 4.** Solubilization of the Bis-PMP-Salt of PPA by the Addition of 2 Equiv of Urea Host Monomer 1 in THF and Procedure for Imprinting

**Table 4.** Numbers of Molecules within a 5.05 Å Cutoff from Each Template

|        | EGDMA | PMPH$^*_1$ | PMP | T  | THF | 1  |
|--------|-------|------------|-----|----|-----|----|
| T-P1C  | 8     | 13         | 0   | 15 | 3   |    |
| T-P1N  | 12    | 0          | 5   | 17 | 1   |    |
| T-P2C  | 5     | 33         | 0   | 7  | 6   |    |
| T-P2N  | 13    | 0          | 2   | 17 | 3   |    |
| T-P3C  | 8     | 14         | 1   | 14 | 4   |    |
| T-P3N  | 12    | 0          | 0   | 21 | 1   |    |
| T-P4C  | 8     | 11         | 0   | 14 | 6   |    |
| T-P4N  | 12    | 1          | 0   | 20 | 1   |    |

$^*_1$Values presented are obtained through integration of RDFs, assuming the template as the solute molecule with an average number of molecules within the cutoff for an average template.
This contrasted with the results from nitrogen sorption analysis and swelling tests, both reflecting the porous and structural properties of the polymers (Figure S5, Table S4). All polymers exhibited mesoporous morphology with surface areas exceeding 200 m²·g⁻¹ and average pore diameters of roughly 4 nm. However, in agreement with our previous report, the imprinted polymers showed a lower surface area and a higher swelling factor than the nonimprinted polymers. We tentatively attributed this template-induced difference in morphology to two effects of the divalent anion, (1) its action as a physical cross-linking agent and (2) its action as a good solvent for the growing chains leading to delayed phase separation and a more gel-like morphology. Below we will invoke a third alternative explanation to the observed differences based on the dianions ability to coordinate two monomer units of I in a geometry favoring cyclopolymerization (vide infra).

Template Binding Studies. To evaluate the anion recognition properties of the materials we first assessed their ability to rebind their corresponding templates under static conditions. The polymers were incubated in acetonitrile in presence or absence of base (PMP) with the anion templates (Figure 4) followed by quantification of unbound template by HPLC.

Figure 4. Absorbed amount in percent of PPA (a), PSA (b), and BA (c) by anion imprinted and nonimprinted polymers in the absence and presence of 0.1% PMP modifier.

Figure 4 shows the degree of template binding to the polymers in presence or absence of PMP. MIPs generally bound more of the templates than nonimprinted polymers (NIPs) with the largest uptake shown by the polymers prepared from a 2:1 monomer/template ratio and with a significant increase observed upon PMP initiated deprotonation. These results are all in line with the anticipated binding model and both observations correlate with the MD simulation results, whereby increasing the amount of functional monomer in the prepolymerization mixtures increases the extent of template H-bonding (Table 3). Furthermore, template complexation with PPA increases with the degree of deprotonation (neutral < monoanion < dianion). Increasing the amount of PMP will increase the degree of template deprotonation, which increases the degree of interaction with I. Ranking the polymers in order of decreasing uptake showed that only PPA, and to a lesser extent BA, led to polymers exhibiting these effects, whereas the PSA imprinted polymers lacked significant affinity for the template. This agrees with the weak basicity of the corresponding anions and weak interactions observed for this anion in aprotic solvents. The poor performance of the BA-MIPs P5 and P6 however, disagrees with the basicity trend. To understand the reason for this, the differences in anion size and functionality should be considered. PPA monoanion is larger and features an additional H-bond donor in PhP(OH)O₂⁻ which can interact with the polymer. This is lacking in PhCO₂⁻. In view of the observed trends, we chose to characterize the PPA imprinted polymers in more detail.

Adsorption Isotherms and Binding Parameters. The binding-energy distribution of the polymers was obtained from single-component adsorption isotherms determined by a batch equilibrium binding experiment. The binding curves of PPA on P1/Pₙ₁ and P2/Pₙ₂ in acetonitrile buffered with PMP are seen in Figure 5a and 5b. Comparing different binding equilibria of unbound template by HPLC.

Figure 5. Equilibrium binding isotherms of PPA adsorbed to P1 (blue circles) and Pₙ₁ (red squares) (a) and P2 (blue circles) and Pₙ₂ (red squares) (b) in acetonitrile (0.1% PMP). In (c) the binding to P1 (red squares) and P2 (blue circles) corrected for binding to the corresponding nonimprinted polymers is shown. The isotherms were fitted to a Langmuir binary-site (a, b) or monosite (c) models resulting in the binding parameters listed in Table 5. In c, the data points above X = 0.2 have been excluded from the fitting of the data for P2. (d) shows the binding parameters obtained from Figure 5c. Equations (Figure S6) showed that the data were best fitted to the Langmuir binary site model. The associated binding parameters are given in Table 5. The curves exhibited saturation behavior with P₂/P₂,₂ taking up approximately two times more PPA at a given free concentration. This is in line with the nominal capacity of the materials, which is two times higher for P₂/P₂,₂ than P₁/Pₙ₁ and correlates with results derived from MD simulations. The NIPs displayed consistently less binding than the MIPs with the corresponding binding curves rapidly diverging in the low concentration interval of 0–0.5 mM. This appears clearly in the plots of the differential uptake in Figure 5c reflecting MIP binding corrected for binding to the NIP (assuming the latter to reflect the nonspecific binding contribution). Both of these curves fitted with a Langmuir monosite model resulted in nearly identical B_max values reflecting a similar imprinting efficiency of ca. 30% (given identical template concentration in the prepolymerization mixture) and with P2 showing a markedly higher affinity than P1 (Figure 5d). The latter is reflected in the higher equilibrium constant of P₂ (K_eq = 1.9 × 10⁸ M⁻¹) versus P₁ (K_eq = 3.1 × 10⁷ M⁻¹).

This difference can be attributed to the state of template complexation prior to polymerization and is supported by the results obtained from MD simulations. Again, both H-bond
Plausible explanations for this disagreement will be discussed. Accounting for the entropic gain, the 1:1 complex will yield a site with a 2-fold H-bond donor. PPA to monoanionic PPA. This suggests a complexation model that indicates the formation of ternary complexes are dominating. Turning to the RDF-results, the calculated number of molecules within the given cutoff varies with the cutoff and order of the selected solvent/solute pairs. However, in the neutrally templated PPA system (P3), there is one molecule of 1 within 5 Å of any average template. This number increases to three when either increasing the amount of 1 or deprotonating the template PPA to monoanionic PPA. This offers support for the suggested complexation model that indicates the formation of ternary complexes. Increasing the amount of base further leads to dianionic PPA for which RDF results (Table 4) show six molecules of 1 in close proximity to any template. The results obtained moving from 1:1, through 3:1 to 6:1 stoichiometries of 1 and template complexes, in association with the base driven in increase in degree of deprotonation, add support for the formation of ternary complexes.

Assuming that the structures of the prepolimerization complexes are carried into the polymer scaffold, the 1:2 complex will result in a cleft-like receptor featuring four H-bond donors, whereas the 1:1 complex will yield a site with a 2-fold H-bond donor. Accounting for the entropic gain, the resulting binding constant can exceed that of the 1:1 site by 2–3 orders of magnitude, which is clearly not the case here. Plausible explanations for this disagreement will be discussed below.

**Switchable Oxyanion Selectivity.** The crushed polymer monoliths of P1/P1 and P2/P2 were sieved and packed in columns for chromatographic characterization of their retenive properties for the model oxoacids (Scheme 1). The acids were injected in an acetonitrile rich mobile phase as such or buffered with either triethylamine (TEA) or trifluoroacetic acid (TFA). Basic conditions will promote deprotonation leaving the oxyanions to interact with the MIP via H-bonding.

The binding parameters were obtained by fitting of the binding data in Figure 5 to a Langmuir monosite (P1corr and P2corr) or binary site binding model. Imprinting efficiency IE = 100 × B_{max}/B_{max}^a, where B_{max}^a = 125 μmol/g = nominal capacity assuming quantitative template removal and reoccupancy. Fitting parameters for isotherms corrected for nonspecific binding to nonimprinted polymer (e.g., ΔΔEB = BTP/B_{PN1}H). The MIP exhibited strong affinity for its template (PPA) in this mobile phase (Figures 6 and Figures S7) with ca. 75% of injected analyte remaining stuck on the column. Meanwhile, with the exception of PPrA, none of the other acids were retained whereas P2 exhibited no affinity for any of the analytes. This contrasts with the retention results in the TFA-buffered mobile phase. Here we note a complete switch of anion preference with PSA being the only retained anlyte. In absence of modifier both PPA and PSA are retained by P1 and P2 with no retention observed by nonimprinted P1 and P2.

A detailed look at the ionization states of the acids may offer clues to the origin of the turn on/off effect. Table S5 lists the pK_a values and anticipated charges in the presence of the two modifiers of the acids in the study. In presence of base, PSA and PPrA carry a net charge of −2, whereas the monovalent acids are negatively charged −1. In this state both PPA and PPrA can bind to the P2 site while benefiting from 4 complementary H-bond donors. The PPA dianion is the most strongly retained anion agreeing with the fact that this anion was used as template and hence fits best into the binding sites. The weak retention of the monovalent anion of PSA we ascribe to its weak basicity and its hydrophilicity, whereas planar BA lacks steric complementarity with the PPA templated site. In presence of TFA, however, (pK_a = 0.5), PPrA (pK_a = 0.9), and PSA (pK_a = 1.8) with pK_a values in vicinity are partially or fully protonated weakening their interactions with the ura of the stationary phase. PSA (pK_a = 2.8) should be fully ionized in water but less so in 95% acetonitrile in the presence of TFA. As a consequence, PSA is less hydrated under these conditions and can more easily partition into the stationary phase. With only one of the three oxygen acceptors blocked by protonation PSA, as a crude mimic of the PPA dianion, is effectively recognized by the urea binding site.

Some insight is provided by the MD simulations. Regarding template rebinding in absence of modifier, it is clear from the H-bond analysis results (Table 3) that with respect to neutral templates, 1 will interact more strongly with PPA than with PSA. This agrees with the observed retention order in pure acetonitrile (Figure 6a). These experiments are based on investigations of polymer system P2, prepared using higher levels of I and PMP (Table 2, Figure S15). Interestingly, the H-bond analysis results (Table 3) revealed that increasing the amount of I promotes template interaction while barely

### Table 5. Equilibrium Constants (K_{eq}) and Binding Capacities (B_{max}) of PPA Imprinted Polymers^a

| polymer | site class | K_{eq} (× 10^3 M⁻¹) | B_{max} (μmol/g) | R² | IE^b (%) |
|---------|------------|---------------------|-----------------|----|---------|
| P1      | HI         | 265 ± 107           | 23 ± 3          | 0.99 | 18      |
|         | LO         | 4.4 ± 0.8            | 63 ± 2          | 0.99 | 50      |
| P2      | HI         | 24 ± 22              | 14 ± 8          | 0.99 |         |
|         | LO         | 1.1 ± 0.6            | 56 ± 2          | 0.99 |         |
| P3      | HI         | 110 ± 52             | 70 ± 14         | 0.97 | 56      |
|         | LO         | 0.4 ± 1.2            | 326 ± 804       | 0.97 | 260     |
| P4      | HI         | 53 ± 30              | 24 ± 8          | 0.99 |         |
|         | LO         | 2 ± 0.5              | 115 ± 5         | 0.99 |         |
| P1 corr | HI         | 31 ± 2               | 36 ± 0.4        | 0.96 | 29      |
|         | LO         | 194 ± 16             | 38 ± 0.3        | 0.96 | 31      |

^a The binding parameters were obtained by fitting of the binding data in Figure 5 to a Langmuir monosite (P1corr and P2corr) or binary site binding model. Imprinting efficiency IE = 100 × B_{max}/B_{max}^a, where B_{max}^a = 125 μmol/g = nominal capacity assuming quantitative template removal and reoccupancy. Fitting parameters for isotherms corrected for nonspecific binding to nonimprinted polymer (e.g., ΔΔEB = BTP/B_{PN1}H).

Hence, increasing the extent of deprotonation of PPA strongly influences template complexation with both I and PMPH. The MIP exhibited strong affinity for its template (PPA) in this mobile phase (Figures 6 and Figures S7) with ca. 75% of
affecting involvement of PMPH⁺. Further, increasing the degree of deprotonation of template reduces template–template interactions, which is reasonable given the increased charge repulsion. Strong support for the formation of higher order complexes is offered by the RDF analysis. The degree of aggregation is noticeable when inspecting the simulated systems visually (Figures 7, S16, and S17). There is a markedly higher presence of PMPH⁺ (2-fold) in the vicinity of PPA in Figure 8 to a Langmuir monosite binding model.

Binding of Inorganic Phosphate and Sulfate in Water. Given the strong and switchable ion binding shown by the MIP in water poor media, our next goal was to investigate whether the polymers would cross-react with inorganic anions in buffer. Several synthetic receptors have been developed for this purpose to address various water processing or sensing applications, but so far only few receptors have shown effective anion recognition in water and even more rare are those functioning in high ionic strength media. A key problem is that both phosphate and sulfate are strongly hydrated in water (Table 1) which has a destabilizing effect on the interactions with the imprinted site.

To increase particle wettability, we prepared a new set of polymers as P2/P3,2 but using pentaerythritoltriacrylate (PETA) as a hydrophilic crosslinker. Ion binding was measured by conductometry in buffer pH = 9.0 (c) Binding isotherms of P7 corrected for binding to the nonimprinted polymer P N7. (d) Equilibrium constants (Keq, black bars) and binding capacities (Bmax, gray bars) for the indicated anions interacting with P7 in buffer pH 9.0. The binding parameters were obtained by fitting of the corrected binding data in (c) to a Langmuir monosite binding model.

Table 6. Equilibrium Constants (Keq) and Binding Capacities (Bmax) of PPA Imprinted Polymers

| polymer | anion    | Keq (× 10⁶ M⁻¹) | Bmax (µmol/g) | R² |
|---------|----------|-----------------|---------------|----|
| P7      | NaHPO₄   | 1.5 ± 0.3       | 53 ± 5        | 0.971 |
|         | Na₂SO₄   | 1.1 ± 0.2       | 25 ± 3        | 0.976 |
|         | PPA      | 2.2 ± 0.6       | 34 ± 3        | 0.966 |
|         | PSA      | 1.0 ± 0.3       | 24 ± 3        | 0.959 |
| P₇corr  | NaHPO₄   | 0.5 ± 0.2       | 51 ± 18       | 0.940 |
|         | Na₂SO₄   | 0.4 ± 0.2       | 30 ± 8        | 0.970 |
|         | PPA      | 1.5 ± 0.5       | 18 ± 2        | 0.950 |
|         | PSA      | 0.3 ± 0.2       | 27 ± 11       | 0.940 |
| P7      | NaHPO₄   | 3.9 ± 0.5       | 19 ± 0.6      | 0.990 |
|         | Na₂SO₄   | 4.6 ± 1.0       | 58 ± 0.3      | 0.973 |
|         | PSA      | 3.1 ± 0.1       | 18 ± 0.1      | 0.999 |
|         | PSA      | 3.4 ± 0.5       | 61 ± 0.3      | 0.990 |

The binding parameters were obtained by fitting of the binding data in Figure 8 to a Langmuir monosite binding model.

![Figure 7. Example of an observed ternary complex in the P2C system, here involving three molecules of I (gray) engaged in H-bonding interactions with the template (PP2) (purple/magenta) and two non-H-bonding base molecules (PMPH⁺) (yellow). H-bonds are indicated by dashed lines.](image)

![Figure 8. Equilibrium binding isotherms of NaHPO₄ (black curve, triangles), PPA (brown curve, circles), PSA (red curve, diamonds), and Na₂SO₄ (blue curve, squares) on polymer P7 (a) and P N7 (b) in 0.1 M sodium bicarbonate buffer pH = 9.0 (c) Binding isotherms of P7 corrected for binding to the nonimprinted polymer P N7. (d) Equilibrium constants (Keq, black bars) and binding capacities (Bmax, gray bars) for the indicated anions interacting with P7 in buffer pH 9.0. The binding parameters were obtained by fitting of the corrected binding data in (c) to a Langmuir monosite binding model.](image)
preferred sequence or stereoregularity in the polymer main chain that may in turn influence its molecular recognition properties?

We raised this question in view of the ternary monomer−template complexes anticipated to be present during polymerization, the presence of which finds strong support in results from the MD simulations (Tables 3 and 4 and Figures 7, S16, and S17). Such an arrangement may place the reactive double bonds of two monomers in close vicinity (see the left two molecules of 1 in Figure 7). One consequence of this arrangement may be a template induced incorporation of two urea groups juxtaposed in the same chain (Figure 9, route A) akin to the established template polymerization mechanism.55−58 The role of the template is here to bring two or more polymerizable groups close in space to achieve rate enhancement combined with stereo- and sequence-controlled monomer incorporation. The template can be covalently or noncovalently attached to the monomers and is cleaved off after polymerization to free up pendent functional groups.56 This results in high molecular weight copolymers with controlled tacticity and regularity.

To investigate whether monomer reactivity in our system is template controlled or the monomers are randomly incorporated, we used differential scanning calorimetry (DSC), NMR, and MALDI-TOF mass spectrometry to study the initial stages of polymerization. Cross-linked polymers P1 and P2 were compared with linear copolymers of 1 and methyl methacrylate (MMA). All polymers were prepared in absence or presence of template, the latter in the form of mono- or bis-TBA salts of PPA. First, we studied the curing process by DSC monitoring the heat generation upon double bond conversion. The onset and peak maxima temperatures of the curing exotherm were registered (Figures 10a, S9, and S10), and the double bond conversion was estimated from the total heat generated per unsaturation divided by the literature value for the MMA double bond enthalpy (ΔH = 13.1 kcal/mol) (Table S6). All copolymer curing systems experienced a lowering of the peak maxima temperatures upon added template, the effect being more pronounced for PPA-2TBA than for PPA-TBA (Figure 10a). This contrasted with the effect of template addition on the curing of MMA alone (Figure S10), where a slight increase in the onset temperature as well as a significantly lower conversion was observed. This indicates that PPA acts as an inhibitor in absence of the host monomer. Overall, the template had no significant effect on the monomer conversion, which was 40−50% in all cases. We cautiously interpret these results as being due to a template assisted monomer incorporation with PPA-2TBA prearranging two molecules of 1 leading to a catalyzed propagation. The molecular weight distribution of p-(1-co-MMA) determined by MALDI-TOF-MS supports this explanation (Figures 10b and S11). This technique provides detailed information on the molar mass distribution, end groups, and repeat units of linear polymers and in the tandem mode, confirmation of their structural identity.59

The polymers were isolated from the crude reaction mixtures by precipitation in methanol and deposited together
Table 7. Properties of Linear Copolymers Prepared in the Presence of the Templates Indicated

| polymer | template | \(\Delta T_{\text{max}}^\text{o} (^\circ \text{C})\) | \(M_n^b\) (g/mol) | \(M_w^b\) (g/mol) | PDI\(^b\) | MMA/1\(^c\) |
|---------|----------|---------------------------------|-----------------|-----------------|--------|----------|
| p-MMA   | PPA-2TBA | 0                               | 1114            | 1025            | 1.09   | n/a      |
| p-(1-co-MMA) | none    | n/a                             | 1493            | 1174            | 1.27   | 1.74     |
| p-(1-co-MMA) | PPA-TBA | -1.0                            | 1642            | 1319            | 1.24   | 2.36     |
| p-(1-co-MMA) | PPA-2TBA | -2.8                            | 2700            | 1926            | 1.40   | 1.57     |

\(^a\)Change in exotherm peak maxima calculated from the data in Figures S9, S10, and S10a. \(^b\)Number (\(M_n\)) and weight (\(M_w\)) average molecular weights and polydispersity index (PDI) calculated as described in the SI. \(^c\)Calculated from the integrals of the signals corresponding to the \(-\text{OCH}_3\) protons of MMA (\(\delta = 3.6\) ppm) and the aromatic protons Hc of 1 (\(\delta = 8.2\) ppm).

with the MALDI matrix on the target plate followed by recording of the spectra. As seen in Figures 10b and S11 the oligomer distributions were found in the ranges \(m/z\) 100–1500 p-MMA; 500–4000 p-(1-co-MMA); 700–5000 p-(1-co-MMA) (template: PPA-TBA); 800–7000 p-(1-co-MMA) (template: PPA-2TBA), with the signal mass separations corresponding to the expected repeat units i.e. 100.1 g/mol. The corresponding number (\(M_n\)) and weight (\(M_w\)) average molecular weights and polydispersity index (PDI) are listed in Table 7.

From these results we conclude that polymerization in the presence of the doubly charged template leads to a pronounced increase in the average molecular weight of the oligomers, and a significant reduction in polydispersity, all in agreement with the anticipated mechanism.

Finally, we analyzed all polymers by \(^1\text{H}\) NMR spectroscopy. This technique has a broad information value in polymer chemistry.\(^{29}\) From integral ratios of incorporated monomers, the copolymer composition can be estimated, and the technique can be used to estimate the number-average molecular weight,\(^{6,61}\) and polymer stereo- and sequence-regularity.\(^{25}\) Of particular relevance in our case is whether the signals from pendent urea groups are visible and if yes, whether their affinity vis-à-vis the template can be determined.

Figure S12a shows a representative solution spectrum (DMSO-\(d_6\)) of p-(1-co-MMA) prepared in absence of template. We gratefully noted that protons characteristic for both monomers were clearly visible, i.e., the urea protons Ha and Hb and the aromatic protons Hc of 1 and the \(-\text{OCH}_3\) protons of MMA. The integrals of the Hc and \(-\text{OCH}_3\) signals were used to estimate the stoichiometry of incorporated monomers. This is expressed as the ratio MMA/1 and the values are listed in Table 7. For all polymers the ratios are lower than 5/1 indicating an enhanced incorporation of monomer 1 in the chains. More interesting, however, is the low ratio observed for the template induced polymers, most notably for the polymer prepared in the presence of the doubly charged template. This agrees with the anticipated catalytic action of PPA-TBA enhancing the reactivity of 1. We thereafter monitored the complexation induced shifts of the urea protons upon addition of PPA-TBA. As seen in Figure S12, titration was accompanied by pronounced downfield shifts and signal broadening which confirm the expected template-polymer interactions. The CIS plots for all polymers are seen in Figure S13 with the binding parameters listed in Table S7. The increase of the average equilibrium constants from \(K_{\text{eq}} = 139 \text{ M}^{-1}\) for the nontemplated polymer to \(K_{\text{eq}} = 198 \text{ M}^{-1}\) for the templated polymer we ascribe to an enhanced contribution of cross-linking to the polymer a

main chain alone. So, what is the role of the cross-linking agent? Judging from the equilibrium constants for adsorption, the affinity of the cross-linked P2 for PPA (\(K_{\text{eq}} = 1.9 \times 10^5 \text{ M}^{-1}\)) exceeds by far that of the linear polymer. The contribution of cross-linking to the polymer affinity for the template is hence obvious. The importance of the cross-linking monomer is further highlighted in the results obtained from MD simulations. Looking at the neutral templated systems (PXN), cross-linker and 1 form H-bonds between 30 and 50% of the total simulation time (Table 3). Even in the PPA templated systems, extensive H-bonding between the template and the cross-linking monomer is observed, 60–80%. There is a high presence of cross-linker around templates and competition for access to functional groups on templates. The interactions are also stable, often more so than interactions between template and 1 or PMP/PMPH+. H-bond interactions with template are lower in PSA systems than in PPA systems, even if the presence of cross-linker is similar. As observed in several other systems,\(^{445–446}\) the presence of cross-linker in close vicinity to the template is observed in prepolymerization mixtures and is observed to influence template-polymer rebinding.

\section{FINAL DISCUSSION}

Charge neutral receptors interacting with ligands via H-bonding have for long been associated with poor compatibility with aqueous media. The presence of water and other protic solvents effectively disrupts the HG interactions in these systems.\(^1\) Some exceptions to this rule have recently been reported, although neutral receptors displaying affinity for phosphate and sulfate in pure water or buffer are still rare. Inspiration from naturally occurring receptors can be used to move forward. These display high affinity and selectivity in physiological conditions by shielding the anion from the surrounding solvent molecules and thereby freeing it to interact by H-bonding in a nest like site.\(^{1,5}\) To mimic this principle, we have designed oxoanion imprinted polymers incorporating charge neutral urea groups in a hydrophobic scaffold. The polymers exhibit an unprecedented switchable anion binding behavior as well as strong ion affinity in buffered media comprising both phosphate and the strongly hydrated sulfate anion. This performance has a practical relevance in view of the broad need for robust anion binders. As we previously showed, this behavior may be exploited in bioseparations for fractionation of phosphorylated and sulfated peptides or saccharides.\(^{29}\) Another classical problem concerns sulfate separation from nitrate-rich radioactive mixtures which has been a long-term goal in nuclear waste remediation.\(^6\) Several synthetic receptors have been developed for this purpose but so far only few receptors offer effective anion recognition in water and even more rare are those functioning in high ionic strength media. Through further improvements of
the imprinted receptors, such as incorporation of dual receptors or other scaffolds, we hope to sufficiently boost affinity and selectivity for the MIPs to offer a viable alternative host for these purposes.

We also provide detailed molecular level insight into the anion recognitive sites. Comparing stoichiometric imprinting based on 1:1 or 1:2 phosphate anion–urea monomer ratios shows the superiority of the ternary complex imprinting to generate high fidelity binding sites. In spite of the stoichiometric complexion as proven by 1H NMR 2D-NOSY and CIS plots, the imprinting efficiency did not exceed 30% when counting the high affinity sites. This is likely related to the amorphous nature of this class of MIPs supported by the heterogeneity of observed complexes in these mixtures (Figures S16 and S17). MIPs prepared by free radical polymerization belong to the thermostet class of materials characterized by extensive and irreversible cross-linking and insoluble end products. The heterogeneity of these materials ranges from the molecular level to the micro scale which precludes molecular level insights into the structural and dynamic features of the binding sites. This stands in contrast to the range of available tools for characterizing biomacromolecules. Nevertheless, reports focusing on well characterized monomer template systems employing advanced techniques such as solid state NMR with isotopically enriched templates, infrared or fluorescence spectroscopy have increased our knowledge in this regard. As discussed above, the propagation rate versus the on/off rate of the template-monomer interaction is an indicator for the templates ability to induce repeat units in the main chain. Very few studies on how the template influences the initial stages of the polymerization have been reported. This warrants further studies in this regard. As we have shown in this report, a solution binding constant for the host monomer template interaction exceeding 7000 M−1 is sufficient for achieving this. This contrasts with situations where the rate of propagation is slow with respect to the on/off rate. Here the template will have time to dissociate before the next monomer addition. In this case, imprinting follows the mechanism outlined in Figure 9b where preformed polymer chains are conformationally stabilized by the template via multidentate interactions. Although this is likely the dominating mechanism in most noncovalent imprinting systems, studies of corresponding linear polymers can provide further important insights into the imprinting process.

**ASSOCIATED CONTENT**

**Supporting Information**

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacs.0c00707.

Experimental section outlining synthetic procedures, characterization techniques and data evaluation and dynamic simulations, and supporting tables and graphics (PDF)

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**Funding**

Marie Skłodowska-Curie Actions PITN-GA-2010-264699 and H2020-MSCA-ITN-2016, 722171; Swedish Vetenskapsrådet (VR) under grant number 2014–3794; Sweden Knowledge-Foundation (KKS) for research under project number 20150086. Deutsche Forschungsgemeinschaft (DFG) (Se 777/9-1).

**Notes**

The authors declare no competing financial interest.

**ACKNOWLEDGMENTS**

We gratefully acknowledge financial support from the Marie Skłodowska-Curie Actions (PITN-GA-2010-264699 and H2020-MSCA-ITN-2016, 722171), Vetenskapsrådet (VR) under grant number 2014–3794, the Deutsche Forschungsgemeinschaft (DFG) under grant number Se 777/9-1 and the Sweden and the Sweden Knowledge-Foundation (KKS) for research under project number 20150086. S.S. gratefully acknowledges a Matching Funds fellowship from the Deutsche Akademische Austauschdienst (DAAD). C.E. is grateful for the postdoctoral research fellowship from the Council of Higher Education in Turkey (YÖK) and for the visiting academic funding from the German Academic Exchange Service (DAAD).

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