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Peptide Length and Dopa Determine Iron-Mediated Cohesion of Mussel Foot Proteins

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Mussel adhesion to mineral surfaces is widely attributed to 3,4-dihydroxyphenylalanine (Dopa) functionalities in the mussel foot proteins (mfps). Several mfps, however, show a broad range (30%–100%) of tyrosine (Tyr) to Dopa conversion suggesting that Dopa is not the only desirable outcome for adhesion. Here, a partial recombinant construct of mussel foot protein-1 (rmfp-1) and short decapeptide dimers with and without Dopa are used and both their cohesive and adhesive properties on mica are assessed using a surface forces apparatus. Our results demonstrate that at low pH, both the unmodified and Dopa-containing rmfp-1s show similar energies for adhesion to mica and self–self-interaction. Cohesion between two Dopa-containing rmfp-1 surfaces can be doubled by Fe³⁺ chelation, but remains unchanged with unmodified rmfp-1. At the same low pH, the Dopa-modified short decapeptide dimer did not show any change in cohesive interactions even with Fe³⁺. The results suggest that the most probable intermolecular interactions are those arising from electrostatic (i.e., cation– π) and hydrophobic interactions. It is also shown that Dopa in a peptide sequence does not by itself mediate Fe³⁺ bridging interactions between peptide films: peptide length is a crucial enabling factor.

where Dopa was tethered to a cantilever tip showed Dopa contributes to nano-Newton adhesion on iron oxide, titania, and amine-functionalized surfaces.^[2] Moreover, several studies with Dopa functionalized polymers have demonstrated a strong positive linear correlation between Dopa content and adhesion to different surfaces.^[3] Notwithstanding these trends, much debate persists regarding two critical issues of mfp-mediated adhesion: (1) the actual interfacial chemistry of Dopa side-chains on model surfaces and (2) the contribution of residues other than Dopa to adhesion. The first issue has seen significant progress by the application of resonance Raman microscopy to detect the pH-dependent formation of bidentate binuclear Ti^{IV} coordination complexes between Dopa-containing mfp-1^[4] and mfp-3^[5] on titania surfaces. The second issue, that is, contribution of other residues, is more challenging because the

1. Introduction

Mussels assemble a battery of proteins known as mussel foot proteins (mfps) into a byssus (plaque and the thread) to adhere to solid surfaces in the high-energy intertidal zone. Dopa (3,4-dihydroxyphenylalanine), a post-translational modification from Tyrosine (Tyr), features prominently in mfps, ranging from less than 5 mol% in mfp-4 to 30 mol% in mfp-5.^[1] Single molecule tensile tests using an atomic force microscope

sequences flanking Dopa in most mfps are so variable that selecting a representative or relevant sequence to test is difficult. Mfp-1, a coating protein, is a rare exception in this regard in that it contains more than 70 tandem high fidelity decapeptide consensus repeats, e.g., AKP*SY*P*P*TY*K, where * denotes optional hydroxylation sites. That is to say, peptides can be found in the native protein with none, all, or some combination of hydroxylations present.^[6]

A significant challenge to assessing the adhesive contributions of other amino acids is the complexity of most native mfp sequences, which are polar with high charge density and little to no 2° structure in solution.^[7] The sequences are further complicated by the highly variable post-translational modification by enzymes. In purified native mfp-1, for example, overall Tyr→Dopa and Pro→Hyp conversion can range from 50% to 80%. To reduce sequence complexity, we used a recombinant mfp-1 (rmfp-1) analog that contains 12 tandem repeats of the decapeptide sequence AKPSYPPTYK. This is less than a sixth of the 75 decapeptide repeats in native mfp-1 from *Mytilus edulis*,^[6a] has no post-translational modifications, and limits Tyr to a simple repeating consensus sequence P-T/S-Y-X, where X is P or K. For purposes of the present study, we propose that the decapeptide repeats are not uniformly converted to the fully hydroxylated version because there is some adaptive advantage in not doing so. The present investigation examines the consequences of

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including/excluding hydroxylation on adhesion, cohesion, and Fe^{3+} binding.

In a previous study, the sequence differences of mfp-1 from two related species (*Mc*, *Mytilus californianus*, and *Me*, *Mytilus edulis*) were investigated with regard to Fe-mediated cross-linking of mfp-1 films.^[8] The interaction between Fe^{3+} and mfp-1 using surface sensitive and solution phase techniques showed that the mfp-1 homologs bind Fe^{3+} differently: mfp-1 (*Mc*) Dopa groups interact with Fe^{3+} to form intramolecular complexes, whereas mfp-1 (*Me*) Dopa groups form intermolecular complexes.^[8] Similarly, the adhesive and cohesive contributions of residues other than Dopa in other mfps are the topic of recent studies^[8,9] and will be discussed later.

An important assumption in this study is that an rmfp-1 analog with only 12 tandem repeats of the unmodified decapeptide sequence (the native mfp-1 sequence from *Mytilus edulis*^[6a] has 75 decapeptide repeats) retains some attributes of unmodified decapeptides in native protein. More than 80% of the Tyr in rmfp-1 can be converted to Dopa by tyrosinase,^[10] enabling a separate assessment of contributions by Dopa. Accordingly, rmfp-1 with and without Dopa was tested for adhesion and cohesion on mica using a surface forces apparatus (SFA). We also tested shorter decapeptide dimers (two repeats of the decapeptide sequence, monomer = AKPSYPPTYK) with and without the hydroxylation of Tyr (Y) to Dopa (Y*) and Pro (P) to Hydroxyproline (P*) for cohesion in metal ion (Fe^{3+}) environments to assess the role of peptide length in the formation of metal-protein complexes.

Our results are remarkable in showing that rmfp-1 without Dopa achieves adhesion comparable to Dopa-modified rmfp-1 on mica. Cohesive interactions are also comparable except when Fe^{3+} is added to symmetric surfaces of rmfp-1 with Dopa. However, the cohesive interactions between short decapeptide dimers remained the same regardless of the presence or absence of Dopa, thus stressing the importance of understanding the molecular parameters beyond Dopa that contribute to mussel adhesion.

2. Results and Discussion

2.1. Cohesion (Self-Interaction) between the Protein Films and Interaction with Mica

The cohesive force of interaction between two symmetric rmfp-1 films, Dopa modified and unmodified, was measured in an SFA (Figure 1A) at two different pH values, pH 3.7 and 7.5 (Figure 2). The effect of Fe^{3+} on the cohesive force between the protein films was also investigated (Figure 3). The protein film studies were conducted under low pH environment because it was recently shown that mussels dramatically acidify (pH \approx 2–4) the local environment at the substrate–plaque interface during plaque formation.^[11]

At pH 3.7, similar cohesive interactions were measured for Dopa-containing and unmodified rmfp-1 (no Dopa) when surfaces were kept under compressive contact at $t \geq 10$ min ($W_c = 4.9 \pm 0.6$ mJ m⁻²) (Figure 2A,B). For short contact times, $t_c \approx 2$ min, the Dopa-modified rmfp-1 showed almost 60% higher cohesion ($W_c = 2.40 \pm 0.6$ mJ m⁻²) compared to the unmodified

protein film ($W_c = 1.5 \pm 0.8$ mJ m⁻²). This suggests that Dopa may accelerate the development of cohesion between the protein films; however, given enough interaction time, Dopa adds little to the magnitude of cohesive strength between the protein films at equilibrium. The kinetics of bonding interactions during the contact between the films remains complex and somewhat beyond the reach of experiment; however, we observe that Dopa expedites cohesion between the films at short contact times.

At pH 7.5, the Dopa-containing rmfp-1 ceased to cohere and instead showed long-range steric repulsion (Figure 2D). This is similar to the trend reported for the native mfp-1 and attributed to dopaquinone formation and the conformational consequences of the tautomerization of dopaquinone to Δ -Dopa.^[12] Interestingly, the unmodified rmfp-1 showed significant cohesion ($W_c = 2.0 \pm 0.5$ mJ m⁻²), perhaps because there was no Dopa to oxidize. However, unlike native mfp-1 the range of interaction between the rmfp-1 films was not altered (Figure 1C). The cohesion measured in the unmodified rmfp-1 is contrary to previous observation where the protein did not show cohesion at similar salt concentrations and at a lower pH 5.5.^[13] This could possibly be due to the dimerization of the protein since the authors had observed a thicker hard-wall ($D_H = 20$ –25 nm compared to 3–5 nm in our work) in their experiments and suggested aggregation of the proteins during its synthesis. Recent results suggest that the starting concentration of solutions used for bulk deposition plays a crucial role in determining the adhesive and cohesive properties of a protein film.^[8] Hence, the disparity in the results could also be attributed to the lower protein deposition concentrations (20 $\mu\text{g mL}^{-1}$ compared with 50 $\mu\text{g mL}^{-1}$ in this work) used in the earlier work.

The cohesion between the unmodified rmfp-1 films was completely recovered when the pH of the buffer was switched from 3.7 to 7.5 and back to 3.7 unlike the Dopa-modified rmfp-1 where the protein underwent pH-induced irreversible structural changes and cohesion could not be recovered. At low pH and low salt concentrations, π -cation^[14] and hydrophobic^[15] interactions are strong and these interactions tend to get weaker at higher pH and high salt conditions. Thus, the reversible cohesive behavior of the unmodified rmfp-1 film demonstrates that cohesion in rmfp-1 films could be due to electrostatic (e.g., π -cation),^[16] hydrophobic interactions^[17] and π - π stacking^[18] and that Dopa is not essential for cohesion as has been repeatedly argued in the literature.^[3a,5,19]

Another intriguing finding was related to the adhesion of the unmodified (no Dopa) and the Dopa-modified rmfp-1 film to mica. Both the proteins showed similar time dependence and adhesion energies to mica. Unmodified rmfp-1 adhered to mica with $W_{ad} = 8.0 \pm 0.1$ mJ m⁻², whereas the Dopa-modified rmfp-1 showed similar adhesion energy of $W_{ad} = 9.8 \pm 1.2$ mJ m⁻² at $t_c = 60$ min (Figure S1, Supporting Information). Protein adsorption experiments in a Quartz Crystal Microbalance (QCM) further established that the presence of Dopa in the protein does not change the mass of protein ($m \approx 80$ ng cm⁻²) adsorbed to a TiO₂ surface (Figure S2, Supporting Information). The negligible change in the dissipation of the quartz crystal (Figure S2, Supporting Information) upon the adsorption of the protein at pH 3.7 indicates that rmfp-1, both with and without Dopa, forms a stiff film on TiO₂, and bidentate

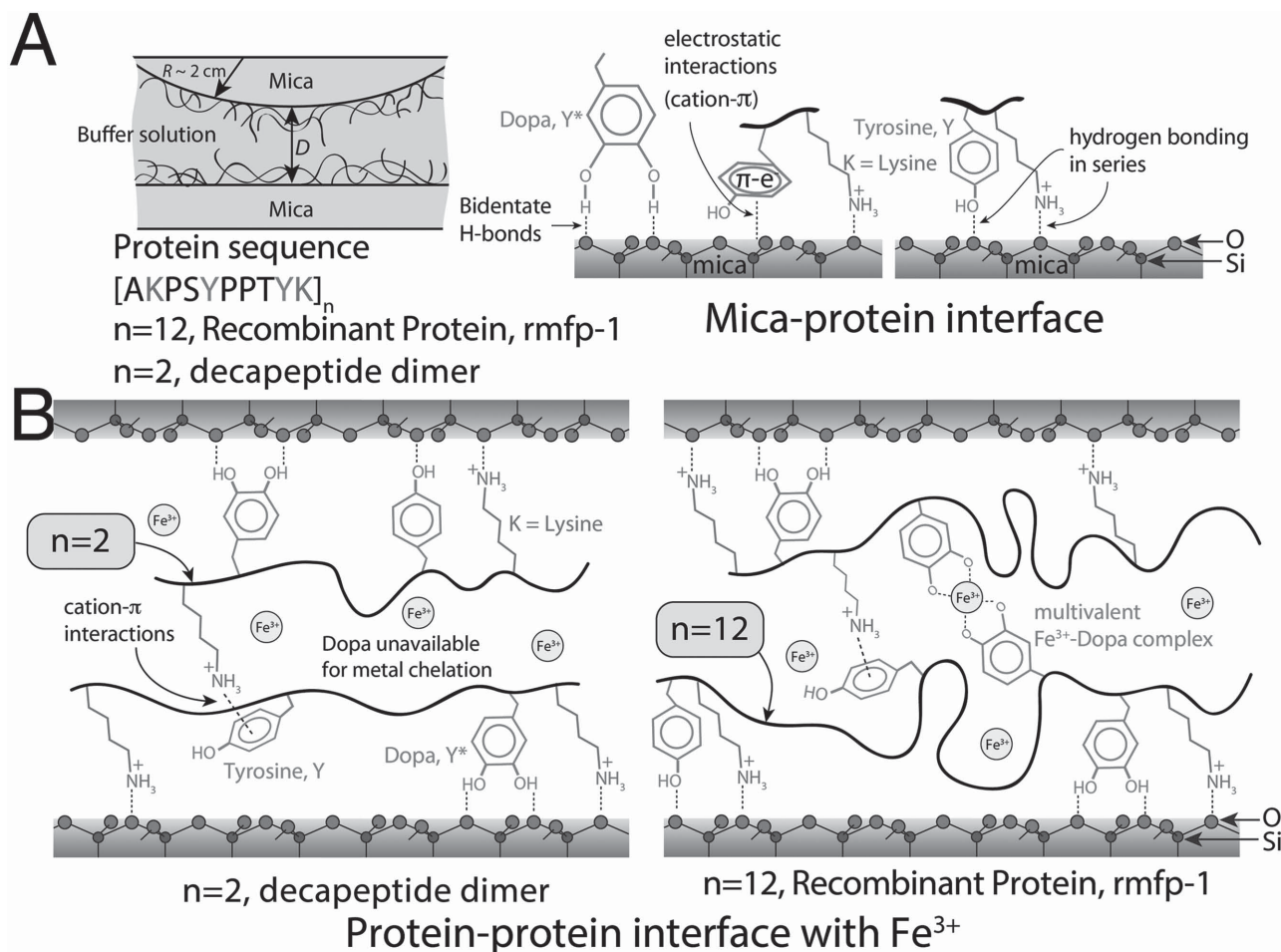


Figure 1. Scheme of the surfaces analyzed by the surface forces apparatus. A) rmfp-1 and short peptides with or without Dopa are adsorbed as thin films onto one or both mica surfaces. Schematics of the bidentate H-bonds, electrostatic, and π -cation interactions between the protein and K⁺ ions adsorbed (not shown for the sake of clarity) to the mica surface (see Figure S1, Supporting Information). Our results suggest that electrostatic, π -cation, and hydrophobic interactions between aromatic residues and mica are more probable than bidentate H-bonding interactions. B) Schematics showing the effect of peptide length on the adhesive interactions between the protein films. Metal-mediated cross-links across the films are possible for proteins containing Dopa residues only when the number of decapeptide monomers is greater than a critical number (n between 2 and 12). For the short decapeptide dimers, most Dopa residues get recruited to the substrate, whereas for the decapeptide 12-mer, free Dopa residues remaining at the protein-solution interface are available to bridge with exposed Dopa on the opposing surface through Fe³⁺-mediated chelation.

coordination bond^[3a,19a] of the Dopa to the crystalline TiO₂ is not the dominant mechanism that binds the protein to the surface at these solution conditions. It was previously demonstrated that hydrophobicity in the mfps mediates dehydration at substrate protein interface to allow force-free adhesion of the protein to a substrate^[20] and that the adsorption of the proteins to a surface depends on the Dopa content for small decapeptide monomers or dimers.^[21] However, present results argue that for a decapeptide 12-mer, the force-free adsorption of the protein (as measured in the quartz crystal microbalance with dissipation (QCM-D)) is surprisingly independent of the presence of the Dopa residue. It should be noted that the thickness of the rmfp-1 film with Dopa was about 4–5 nm compared to 0.7–1.5 nm for the rmfp-1 film without Dopa as measured in the SFA (Figure S1, Supporting Information). The presence of Dopa might affect the structure of the adsorbed rmfp-1 film on the surface; however, both films showed similar adhesive/cohesive properties (SFA studies) and stiffness (QCM-D measurements).

Similar adhesion energies of Dopa modified and unmodified protein to mica also suggest that the primary interaction between the protein film and mica could be due to specific Coulombic interactions between the lysine and negatively charged mica or monodentate hydrogen bonding in series with lysine-mica interactions (Figure 1B). Hydrophobic interactions between the aromatic residues and the hydrophobic domains in the mica crystal^[15] could also cause a strong adhesion between protein and the surface. π -cation interaction between the aromatic residues of the peptides in the protein and the K⁺ in the mica crystal lattice could also possibly cause enhanced interaction between the protein and the surface, and bidentate bonds between Dopa and the polysiloxane lattice of mica might play a minor role in the adhesion. Similar π -cation interactions were previously proposed between lignin and gold^[22] and lipid bilayers and proteins.^[23] The work of adhesion between the mica and rmfp-1 was approximately $W_{ad} = 7.8 \pm 0.6 \text{ mJ m}^{-2}$ for both Dopa modified and unmodified rmfp-1 (bidentate H-bonds not possible) at

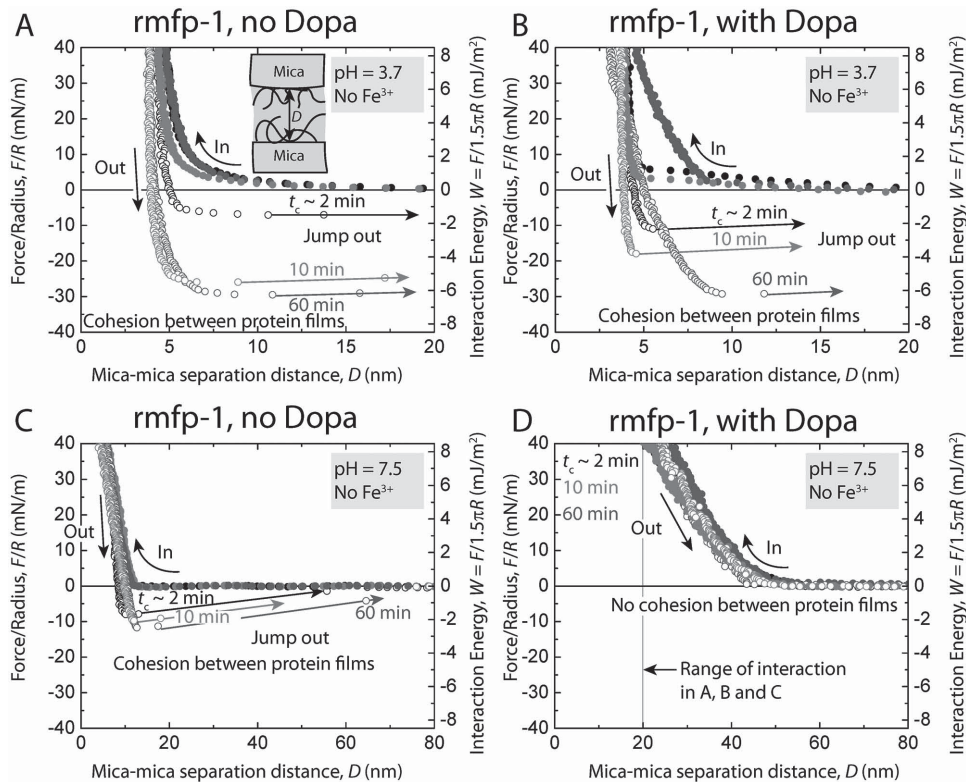


Figure 2. Representative force versus distance plots showing the effect of contact time, t_c , on the cohesion between A,C) two symmetric rmfp-1 films without Dopa and B,D) two Dopa-containing rmfp-1 films at pH 3.7 and pH 7.5, respectively.

short contact times $t_c \approx 2$ min (Figure S1, Supporting Information), which suggests that bidentate Dopa bond to mica cannot be the primary mode of binding to mica surfaces by rmfp-1. It should be noted that the true adhesion energy of the protein to the substrate is likely to be greater than the value measured in the SFA. On preadsorbing the protein to mica, most residues endowed with surface-binding reactivity get recruited to the substrate thus become unavailable to bind the opposing interface. Hence, our measurements show that the binding strength of the decapeptide 12-mer to a mica surface is >7.8 mJ m⁻².

There was no material transfer between the surfaces during the force measurements because the approach force-run

profiles for the very first contact between the surfaces were similar to the successive runs repeated at least six times at the same contact point. The measured cohesive force also did not change significantly (<1%) for the successive force measurements at a given contact point. The failure during the separation of the protein films was determined to be the protein-protein interface and not the mica-protein interface as the adhesion measured between rmfp-1 (unmodified or Dopa-containing rmfp-1) and mica was significantly higher ($W_{ad} = 8.4 \pm 0.8$ mJ m⁻²) than the cohesive energies ($W_c = 3.9 \pm 1.7$ mJ m⁻²) of symmetric rmfp-1 films at $t_c = 2$ to 60 min (Figure 2 and Figure S1, Supporting Information).

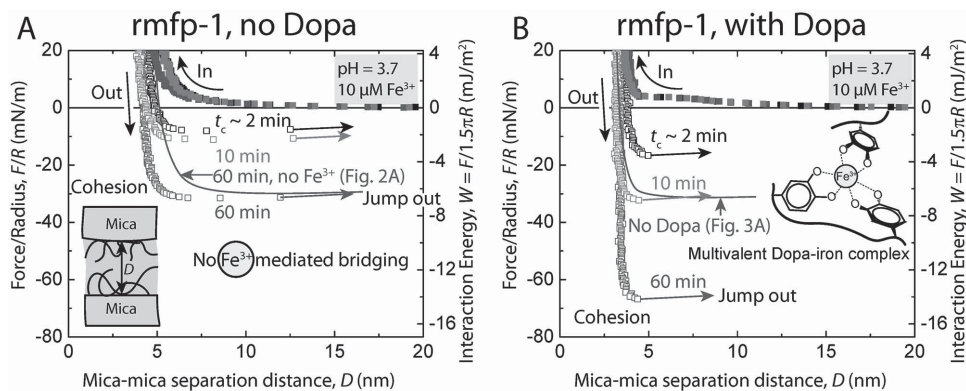


Figure 3. Representative force versus distance plots showing the effect of contact time, t_c , on the cohesion between two symmetric A) unmodified rmfp-1 and B) Dopa-containing rmfp-1 films at pH 3.7 with 10×10^{-6} M Fe^{3+} between the surfaces.

Introduction of 10×10^{-6} M Fe^{3+} into the gap between rmfp-1 surfaces did not change the cohesion between the unmodified rmfp-1 films ($W_c = 5.9 \pm 0.8$ mJ m $^{-2}$ for $t_c = 60$ min with and without Fe^{3+}). However, Fe^{3+} doubled the cohesion energy between the Dopa-containing rmfp-1 after similar contact times (Figure 3) and the forces measured were reversible. Contact time t_c between the surfaces significantly changed the cohesive energy from $W_c = 3.3 \pm 0.4$ mJ m $^{-2}$ for $t = 2$ min to $W_c = 10.0 \pm 2.8$ mJ m $^{-2}$ at 60 min for the Dopa-containing rmfp-1 surfaces apparently due to Fe^{3+} bridging coordination or previously observed Fe^{3+} -mediated covalent cross-linking at low pH.^[24] To determine the mechanism of Fe^{3+} -mediated cohesion between the Dopa-modified rmfp-1 films, the force measurements were repeated several times ($N = 6$; see the Experimental Section) at a given contact point. There was no material transfer between the surfaces during the force measurements because the approach force-run profiles for the very first contact between the surfaces were similar to subsequent force runs and reversible. This observation argues against the covalent cross-linking (irreversible process) of the peptide films by Fe^{3+} in acidic pH and suggests that Fe^{3+} bridging between the Dopa-modified rmfp-1 films is limited to coordination complexes (Figures 1B and 3B).

The temporal increase in the Fe^{3+} -mediated cohesive forces (or energies, W_c increases for contact time, $t_c = 2$ min to 60 min) indicates that it takes time for the Fe^{3+} to recruit two or more Dopa and bridge them across the surfaces. These results also show that Fe^{3+} is involved in chelating only the Dopa moieties in the rmfp-1 films by forming multivalent catecholate-Fe complexes across the surfaces; however, other hard Lewis acid donors such as the -OH of the Tyr or the -NH $_2$ of lysine between rmfp-1 surfaces are not coordinated. The ligand number of the Fe^{3+} -Dopa complex depends on the pH and the ratio of Dopa to Fe^{3+} ,^[24b] and the bridging of rmfp-1 surfaces is by bis- and tris-catecholato- Fe^{3+} complex formation. The local pH within the protein film can be different from the bulk pH^[25] (rmfp-1 has a pI of ≈ 10); hence determining the ratio of bis to tris complexes at an interface is challenging and beyond the scope of this work. The magnitude of Fe^{3+} -mediated cohesion between the Dopa-modified rmfp-1 films measured in this work is comparable with biotin-avidin interfacial bond energy ($W_{ad} \approx 10$ mJ m $^{-2}$),^[26] the strongest known non-covalent

interaction between a protein and a ligand. Two to three Dopa residues of mfp-1 in the cuticle of the marine mussels complex with a single Fe^{3+} ,^[27] thereby creating a stable complex that can, in principle, be translated to cross-link other structural proteins. These iron-protein complexes have a breaking force nearly half that of covalent bonds (as measured in our experiments), but unlike covalent bonds they can form and break reversibly, making them ideal for creating sacrificial cross-links to prevent catastrophic failure of a material.

2.2. Cohesive Interactions between mfp-1 Short Peptide Dimers with Dopa

Cohesive interactions between short decapeptide dimers (Pro-pep, [AKPSYPPTYK] $_2$) of the consensus decapeptide repeat unit of mfp-1 were measured to determine the effect of peptide length on the energy of interaction between the protein films uniformly deposited on mica surfaces. We investigated the effect of Fe^{3+} on the change in cohesive energy between the short peptide films. Another short decapeptide dimer (Hyp-Pep, [AKP*SYP*P*TYK] $_2$, P* = *trans*-4-hydroxyproline) with hydroxyproline modification was also tested for cohesion. Hyp-pep dimer is a closer mimic of the consensus decapeptide repeat unit of mfp-1 which has *trans*-4-hydroxyproline modification at P-3, P-6, and P-7 of the decapeptide (additional *trans*-3 modification occurs at P-6, but was not tested here). We also assessed if hydroxylation of proline has an effect on the cohesive and metal chelating properties between the protein films.

At pH 3.7, the cohesive energy of interaction between unmodified mfp-1 Pro-pep (proline containing dimer) film was $W_c = 8.1 \pm 1.1$ mJ m $^{-2}$ at short contact times, $t_c = 2$ min (Figure 4A), and did not change when the surfaces were kept under compressive contact for $t = 10$ –60 min unlike rmfp-1 (Figure 2A,B). Dopa-modified Pro-pep dimer showed cohesion energy similar to the unmodified dimer. The forces measured between unmodified mfp-1 Pro-pep dimer films on approach were purely repulsive due to steric and hydration forces^[28] (Figure 4A).

The cohesion energy between the mfp-1 peptide films did not change on introducing 10×10^{-6} M Fe^{3+} between the surfaces regardless of the Dopa modification of the decapeptide

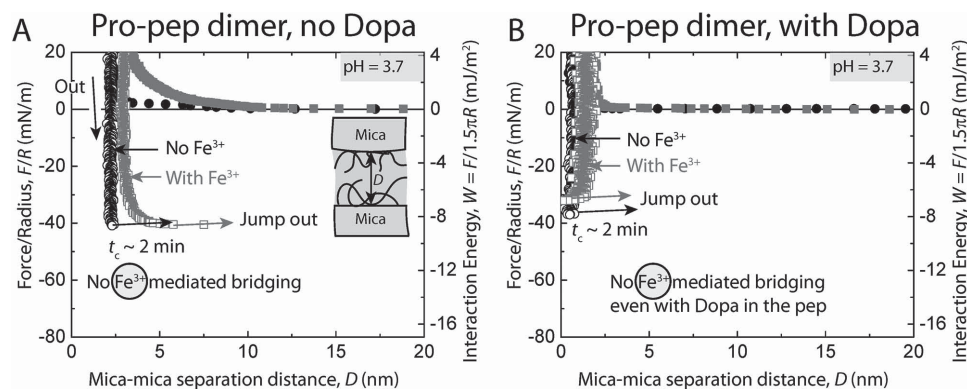


Figure 4. Representative force versus distance plots of cohesion between two symmetric A) unmodified (no Dopa) and B) Dopa-containing mfp-1 peptide dimer (with proline, Pro-pep) films at pH 3.7 with (green points) and without (black points) 10×10^{-6} M Fe^{3+} between the surfaces.

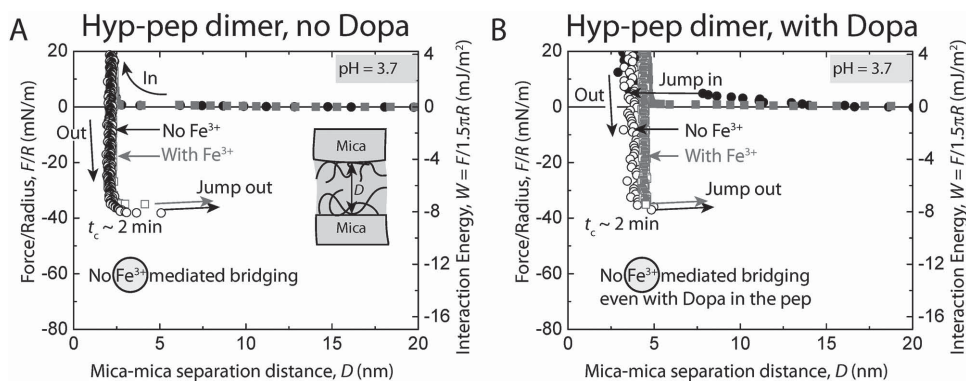


Figure 5. Representative force versus distance plots of cohesion between two symmetric A) unmodified (no Dopa) and B) Dopa-containing mfp-1 peptide dimer (with *trans*-4-hydroxyproline, Hyp-pep) films at pH 3.7 with (black points) and without (green points) 10×10^{-6} M Fe^{3+} between the surfaces.

dimers (Figure 4) for up to $t_c = 60$ min. In a separate experiment, the Dopa-modified decapeptide dimers were given longer times (up to $t_c = 24$ h) to interact cohesively in the presence of Fe^{3+} ; however, the cohesive energy of interaction did not change significantly ($W_c = 7.7 \pm 0.9$ mJ m $^{-2}$, $n_{\text{trials}} = 4$). This is contrary to the commonly observed property of ferric ions to chelate Dopa containing protein films across surfaces as shown in our rmfp-1 films experiments and previously seen in natural mfp films.^[8,13] Perhaps the Dopa needed to coordinate and form Fe^{3+} -mediated bridges between the films is unavailable by virtue of interacting with the mica surface through various interactions as shown in Figure 1B.

Interestingly, the peptide dimers with hydroxyproline (Hyp-pep) showed cohesion energies similar to the Pro-pep dimers ($W_c = 9.4 \pm 1.2$ mJ m $^{-2}$) and Dopa did not have an effect on the interaction energies between the films (Figure 5). Fe^{3+} was also unable to enhance the cohesive interactions between the Hyp-pep films. These results suggest that peptide length is a critical design parameter for Fe^{3+} -mediated cohesive bridging. We showed that there is a critical number for the repeating decapeptide unit of the monomer between 2 and 12 necessary to trigger metal chelation (Figure 1B) between the peptide films and that incorporating Dopa into a peptide sequence does not necessarily guarantee the formation of metal-mediated cross-links between the peptide films.

3. Conclusions

In this work, we demonstrate that bidentate hydrogen bonding by Dopa plays only a minor role in the adhesion of mfp-1 to mica (or adsorption to titania surface). The adhesion of the proteins or peptides to a mica surface is more due to specific Coulombic interactions between lysine and the negative mica surface or monodentate hydrogen bonding in series with lysine–mica interactions. Hydrophobic interaction between the aromatic residues and the hydrophobic domains in the mica crystal lattice or π -cation between the aromatic rings in the protein and the ions adsorbed to the mica interface are possibly responsible for the adhesion.

Since the catechol group did not influence the cohesive strength between the protein films, π - π stacking, hydrophobic and π -cation interactions are more likely to contribute to the

strong cohesion at pH 3.7. Dopa residues tend to accelerate bond formation between the peptide films, however, given enough time, the equilibrium cohesive energy between the films is independent of the Dopa residues in the protein film. The cohesion energy between the protein films was similar for a decapeptide dimer and a 12 mer suggesting that entanglement–entrapment mechanisms^[29] are not responsible for the bonding between the mussel inspired peptide films. Cohesion between Dopa-containing rmfp-1 surfaces can be doubled through Fe^{3+} -mediated chelation resulting in an interfacial energy of $W_c \approx 10$ mJ m $^{-2}$ which is equivalent to biotin–avidin interfacial adhesion energy, the strongest known noncovalent interaction; but unlike the protein and ligand interaction, the iron-mediated cohesive bond can be broken and formed reversibly.^[30] This interaction is absent without Dopa in the protein.

Incorporating Dopa into a peptide sequence does not guarantee the formation of metal-mediated cross-links between peptide films and the length of the peptide is a very crucial parameter that determines the performance of the materials that involve coordination chemistry. Hence, Dopa containing proteins and peptides with appropriate length could be used as tunable systems for applications in strain-resistant coatings, drug delivery, and bio-adhesives.

4. Experimental Section

Modification of rmfp-1: Rmfp-1 used in this work is a shorter synthetic analog of the natural mfp-1 from *Mytilus edulis* with 12 tandem repeat units of the mfp-1 consensus decapeptide AKPSYPPTYK. The protein had an M + H $^+$ of 13 619 Da by MALDI TOF mass spectrometry. Tyr in rmfp-1 was converted to Dopa by mushroom tyrosinase (Sigma-Aldrich) using the borate capture method^[10] and then purified by C-18 reverse phase high performance liquid chromatography (HPLC) column, eluted with a linear gradient of aqueous acetonitrile. Eluent was monitored continuously at 230 and 280 nm, and 0.33 mL fractions containing peptides were pooled and freeze-dried. Sample purity and hydroxylation were assessed by MALDI-TOF. M + H $^+$ was 13 939 Da with >83% conversion efficiency. The short peptide dimers ([AKPSYPPTYK] $_2$ and [AKP*SYP*P*TYK] $_2$, P* = *trans*-4-hydroxyproline) used in these experiments were obtained from GenScript USA Inc. and Tyr was modified to Dopa by similar methods described above.

Measuring the Adhesive/Cohesive Interactions: The SFA (SurForce LLC) was used to measure the normal forces between two mica surfaces in a cross-cylindrical geometry as a function of the separation distance, D , between them and has been described elsewhere.^[28,31] The protein

films were made by adsorbing 50 μL of the protein from a 50 $\mu\text{g mL}^{-1}$ in a buffer solution (10×10^{-3} M sodium acetate buffer, pH 3.7) onto the mica surfaces for 15 min, then rinsing the excess protein with the same buffer. It should be noted that the protein deposition concentration was set at 50 $\mu\text{g mL}^{-1}$ as previously optimized for mfp-1 for achieving maximum cohesive interactions.^[8] During the protein adsorption, the discs were kept in a saturated Petri dish to minimize evaporation of the water from the surfaces. The discs were then mounted in the SFA in one of the two configurations. In a symmetric configuration (Figure 1A), the mussel protein film was deposited on both surfaces in order to measure “cohesion” between the protein films. Cohesion was tested with and without iron. To test the effect of Fe^{3+} , a 10×10^{-6} M FeCl_3 in acetate buffer (as above) was freshly made and added to the reservoir between the symmetrically deposited protein films on mica.

The protein films were always hydrated (i.e., never allowed to dry) and a droplet of the acetate buffer was injected between the surfaces immediately after loading in the SFA. During a typical approach–separation force measurement cycle, the surfaces were first moved toward each other (approach) until reaching a “hardwall” and then separated. The hardwall distance, D_{H} , is the separation distance between the two mica surfaces upon compression that does not change with increased compression. There was no material transfer between the surfaces during the force measurements because the approach force profiles for the initial contact between the surfaces were similar to the successive runs repeated at least six times at the same contact point. All the experiments were repeated three times. The energy of interaction between two crossed-cylinder geometry roughly corresponds to a sphere of radius R approaching a flat surface based on the Derjaguin approximation, $W(D) = F(D)/2\pi R$ where, $W(D)$ is the energy of interaction per unit area between two flat surfaces and $F(D)$ is the measured force of interaction in the SFA.^[29] The measured adhesion (or cohesion) force F_{ad} (or F_{c}) is related to the adhesion (or cohesion) energy per unit area by $W_{\text{ad}} = F_{\text{ad}}/2\pi R$ for rigid surfaces with weak adhesive interactions, and by $W_{\text{ad}} = F_{\text{ad}}/1.5\pi R$ (used in this study) for soft deformable surfaces with strong adhesion or cohesion.^[29,32]

Protein Adsorption Experiments: QCM-D experiments were done with a Q-Sense E4 open module to characterize the adsorption of rmfp-1 (Dopa modified and unmodified) to TiO_2 surfaces independently of the SFA experiments. The QCM crystals were cleaned in 3% sodium dodecyl sulfate (SDS) solution, rinsed in distilled water, cleaned with ethanol, and then treated with UV–Ozone for 10 min. Frequency and dissipation baselines were established in 100 μL of acetate buffer solution on the crystal followed by injection of 25 μL of 50 $\mu\text{g mL}^{-1}$ rmfp-1. The QCM experiments were repeated three times on each surface for each protein.

Supporting Information

Supporting Information is available from the Wiley Online Library or from the author.

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