Sulfonated Compounds Bind with Prostatic Acid Phosphatase (PAP_{248–286}) to Inhibit the Formation of Amyloid Fibrils

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1. Introduction

Acquired immunodeficiency syndrome (AIDS) is known as a serious infectious disease caused by human immunodeficiency virus (HIV), which greatly threatens the safety of the world and the survival of mankind. Sexual transmission has become the main transmission route of HIV. In 2007, Münch et al. found that semen could significantly promote HIV infection. They described that semen could enhance infection because the endogenous peptide compositions of semen form amyloid fibrils. The most important peptides are 1) prostatic acid phosphatase (PAP) hydrolytic polypeptide PAP_{248–286} and 2) semenogelin 1 hydrolytic polypeptide SEM_{107}. The former assemble themselves into amyloid fibrils termed SEVI (semen-derived enhancer of virus infection), and the latter form SEM1 fibrils.[1–4]

Amyloid fibrils, which are a kind of highly ordered aggregates with rich β-sheet structures, are related to a number of neurological diseases.[5–7] Ramamoorthy et al. recently summarized biophysical and immunological studies on SEVI.[8] SEVI and SEM1 are proposed to promote the effect of HIV infection on account of the strong cationic characteristics of the amyloid fibrils, which possess large amounts of positively charged amino-acid residues such as Arg and Lys.[9,10] There are probably two ways to enhance HIV infection: one way is to ensure close proximity of the negatively charged HIV virus with the target cell by reducing repulsion, and the other way is to capture the HIV virus to assist interaction of the virus with the host cell.[4,9–14] It was reported that PAP_{248–286} could bind to lipid membranes by adopting a helical structure. This membrane fusion process plays an important role in enhancing interactions between the HIV virus and a target cell.[15,16] In addition, the bacterial curli protein was found to catalyze the assembly of PAP_{248–286} to form SEVI as a result of an increase in the rate of aggregation and elongation of PAP_{248–286} cross seeding.[17] Therefore, the amyloid fibrils of semen are an important target to inhibit the transmission of HIV infection. As SEVI is the main component of semen fibrils, three strategies have been developed to inhibit HIV infection: one, inhibition of PAP_{248–286} to form SEVI fibrils at an early stage; two, coat the fibrils to block their infection enhancement properties; three, remodel or degenerate the fibrils.[18] In recent years, some active small mole-

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cules were found to antagonize semen amyloid fibrils. For instance, inhibitors of metal ions, as well as non-natural amino acids, were shown to inhibit the formation of SEVI.\textsuperscript{19,20} Roan et al. found that surfen molecules could effectively interfere during the combination of SEVI and HIV virus particles with target cells, thus weakening the ability of SEVI to promote viral infection.\textsuperscript{21} Furthermore, epigallocatechin-3-gallate (EGCG) was shown to decompose SEVI fibrils under acidic conditions.\textsuperscript{22} EGCG interacts with PAP\textsubscript{248–286} through two steps and binds to the Lys residue of PAP\textsubscript{248–286}. Specifically in the Lys251–Arg257 and Asn269–Ile277 regions.\textsuperscript{23}

Recently, Münch and co-workers also found that CLR01 has anionic phosphate groups.\textsuperscript{18} The compound works as a tweezer to engage the Lys or Arg residues of PAP\textsubscript{248–286} (Figure 1b) to inhibit spontaneous fibrillization of the peptides. Experimental data revealed that the ability of human semen containing the HIV virus to infect cells was diminished greatly.\textsuperscript{18} The properties of CLR01 to antagonize amyloids as well as their antiviral activities prompted us to seek microbicides that could specifically bind with the target peptides. ADS-J1 was the first published anionic sulfonated compound that was shown to inhibit the formation of semen-derived amyloid fibrils and to have high inhibitory activity.\textsuperscript{18,24} Unfortunately, however, ADS-J1 contains an azo group, which has toxic effects. Suramin is also a polysulfonate that has clinical application in the treatment of worm infections and trypanosomiasis.\textsuperscript{25} A lot of commercial derivatives of suramin have been developed, and all of them have anionic sulfonate groups (Figure 1a). As these derivatives possess strong anionic characteristics, we predicted that they could specifically bind with PAP\textsubscript{248–286} and play a role in inhibiting aggregation and fibrillization. Nevertheless, suramin was previously demonstrated to induce severe polyneuropathy, which was related to its dose.\textsuperscript{26,27} In contrast, some suramin derivatives, such as NF110, were reported to inhibit enterovirus A71 infection at lower doses than suramin.\textsuperscript{27} Thus, we will investigate the interactions between PAP\textsubscript{248–286} and the derivatives of suramin in this paper.

However, the answers to many questions still remain obscure. First, it is unclear whether all of the derivatives bind with PAP\textsubscript{248–286}. Second, these sulfonated compounds have various sizes, shapes, and spatial configurations. This diversity may affect the interactions and binding affinities between the peptide and molecules. To discriminate the differences, three molecules (i.e. NF110, NF279, and NF340) were selected as representatives (Figure 1a). Both NF110 and NF340 only have four sulfonate groups, whereas NF279 has six. Moreover, NF110 has an X-shaped configuration, NF279 has an allosteric strategy (linear to V-shaped configuration) upon interaction with the target, and NF340 is not a sulfonated compound. Therefore, the interactions between PAP\textsubscript{248–286} and these derivatives will be investigated in this paper.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure1.png}
\caption{a) Structures of NF110, NF279, and NF340. b) Representative conformations of PAP\textsubscript{248–286} alone, P\textsubscript{248–286}–NF110, P\textsubscript{248–286}–NF279, and P\textsubscript{248–286}–NF340 (from left to right) in the top row, without sulfonated molecules for clarity. The bottom row shows the conformations with sulfonated molecules. The cationic residues of the peptide, such as Lys and Arg, are marked in magenta; sulfonated molecules are displayed in the stick style.}
\end{figure}
and NF340 is smaller and shorter than the previous two molecules. In this manuscript, we explore all of the aforementioned questions by studying the interactions between these molecules (i.e. NF110, NF279, and NF340) and PAP_{248–286} by combining computational and experimental methods. The results provide valuable clues for the development of new molecules that bind with the target peptides specifically to inhibit the aggregation of PAP_{248–286}.

2. Results and Discussion

In this manuscript, we implement molecular dynamics (MD) simulations for four systems to investigate the interactions between PAP_{248–286} and different sulfonated compounds. The four systems include PAP_{248–286}−NF110, PAP_{248–286}−NF279, PAP_{248–286}−NF340, and PAP_{248–286} alone. The root-mean-square deviations (RMSDs) for the four systems are shown in Figure S1 (see the Supporting Information). The plots reveal that all of the systems reach equilibrium within 100 ns simulations and then remain stable. Therefore, the trajectories in the last 50 ns were utilized for analysis.

2.1. PAP_{248–286} Changes Its Conformation Upon Binding to Sulfonated Molecules

We first compared the structures of PAP_{248–286} in the four systems. The timelines for the radius of gyration for PAP_{248–286} are displayed in Figure 2. The results reveal that the peptide presents various properties of compactness upon interaction with the different sulfonated molecules. Even though NF110, NF279, and NF340 are all sulfonated molecules, the radii of gyration for the peptide decrease in the order PAP_{248–286}−NF110, PAP_{248–286}−NF279, PAP_{248–286}−NF340, and PAP_{248–286}. The average radius of gyration for PAP_{248–286}−NF110 system is 1.60 nm. The radii of gyration are 1.42, 1.35, and 1.05 nm in PAP_{248–286}−NF279, PAP_{248–286}−NF340, and PAP_{248–286}, respectively.

The peptide in the PAP_{248–286}−NF340 system becomes tight during the simulation and is more compact than those in PAP_{248–286}−NF110 and PAP_{248–286}−NF279. These changes make PAP_{248–286} in PAP_{248–286}−NF110 more exposed to the sulfonated molecules. Given that PAP_{248–286} has eight positive charged residues (Lys251, Lys253, Lys255, Arg257, Lys272, Arg273, Lys281, and Lys282), open PAP_{248–286} in PAP_{248–286}−NF110 has enough accessible area to interact directly or indirectly with the NF110 molecules (Figure 1b).

We also compared the secondary structures of the peptides in different systems. PAP_{248–286} alone possesses the highest α helix content; the percentage is as high as 38.5%. There are two regions of helical structures: Gly261–Asn269 and Ser279–Met285 (Figure 1b). However, the α helix content decreases in all of the peptides. The α helix content is 15.4% in PAP_{248–286}−NF110 within the Asn265–His270 segment and 25.6% in PAP_{248–286}−NF279 within the corresponding Val262–Met271 fragment, whereas PAP_{248–286}−NF340 loses the helical structure completely and turns into a coil. These changes in the secondary structures of the peptide can also be seen from the contact maps in Figure S2. The contact maps reveal the distances of the heavy atoms from the side chains between two amino acid residues. Distances less than 0.5 nm can be displayed. The decrease in or complete disappearance of the helical confor-

Figure 2. Timelines of the radius of gyration (Rg) for PAP_{248–286} from PAP_{248–286}−NF110, PAP_{248–286}−NF279, PAP_{248–286}−NF340, and PAP_{248–286}.
mation in the peptide structures may be due to the loss of intramolecular interactions within the peptide. The Cα atoms of the residues were calculated by root-mean-square fluctuations (RMSFs) and are shown in Figure S3. These data are consistent with the aforementioned results. The peptide in PAP$_{248–286}$–NF110 has flexible N and C termini. However, the maximum fluctuations of PAP$_{248–286}$–NF340 are present in the middle part, as the peptide loses the α-helix character and becomes a random coil.

2.2. Interactions between PAP$_{248–286}$ and Sulfonated Molecules

PAP$_{248–286}$, an amyloidogenic peptide fragment of prostatic acid phosphatase, tends to assemble to form amyloid fibrils in semen. These characteristic seminal amyloids have the ability to promote the effect of HIV infection.[1,2,4,12] Sulfonated molecules can remodel the peptides and alter their conformations according to previous analysis. These molecules occupy the surrounding sites of PAP$_{248–286}$ by binding with the peptide, and this interferes with interactions between peptides. Therefore, these molecules are very important in inhibiting the aggregation of PAP$_{248–286}$. In this section, we investigate interactions between PAP$_{248–286}$ and different sulfonated molecules and provide detailed information.

Molecules of NF110 bind with PAP$_{248–286}$ through many interactions, such as hydrogen-bonding interactions, hydrophobic interactions, π-π stacking interactions, and electrostatic interactions. The structures of PAP$_{248–286}$ separately bound with NF110, NF279, and NF340 are presented in Figure 1b. Further illustration of PAP$_{248–286}$ interacting with one NF110 molecule is shown in Figure 3a. NF110 is denoted as Mol B, and the other seven NF110 molecules are omitted for clarity. The configuration of NF110 is a distorted X shape owing to the inclusion of several benzene rings, amide bonds, and sulfonate groups. The amide bonds and sulfonate groups of NF110 form hydrogen bonds with the backbones of Gln259 and Gly260 and the side chains of Lys255, Asn269, Lys272, and Arg273 (Figure 3a and Table S1). Moreover, the residues with cationic side chains, such as the ammonium group of Lys and the guanidinium group of Arg, are attracted by the sulfonate groups of NF110 (Figure 3a and Table S1). Besides, Mol B forms hydrophobic and CH–π or NH–π interactions with multiple residues of the peptide (Figure 3a and Table S2). It was reported that π-π interactions affect the fibrillization of amyloid polypeptides.[28,29] EGCG was found to interact with Aβ40 through π-π interactions and to inhibit the formation of amyloid fibrils.[30] Besides hydrophobic interactions, oxidized EGCG could covalently bind to the amyloid fibrils to prevent aggregation.[31] Details about the interactions between PAP$_{248–286}$ and other NF110 molecules can be found in Tables 1, S1, and S2. All of the interactions contribute to strong binding between PAP$_{248–286}$ and NF110. In addition, interactions exist among the NF110 molecules. Figure 3b reveals that one sulfonate group of Mol B coordinates with Mol F through two hydrogen bonds. This not only helps to bind NF110 more tightly but also allows PAP$_{248–286}$ to bind with as many NF110 molecules as possible. NF110 molecules are located on almost the entire surface of PAP$_{248–286}$ and occupy the exposed accessible sites. The whole peptide is buried in a bulky complex. Peptides cannot approach each other, as the NF110 molecules cover the surface of the fibril growth. Furthermore, it is noteworthy that the Gly261–Tyr286 segment was reported to be important for the formation of β-sheet structures in SEVI fibrils,[32] whereas all of these residues interact with the NF110 molecules (Tables 1, S1, and S2). In this way, the NF110 molecules should be able to inhibit the aggregation of PAP$_{248–286}$ through interactions with PAP$_{248–286}$ and among themselves.

NF279, which also comprises a few amide bonds, sulfonate groups, and aromatic rings, has two more sulfonate groups than NF110. Nevertheless, unlike NF110, NF279 has two different configurations: linear and V shaped. Many interactions exist between PAP$_{248–286}$ and NF279. Figure 4a reveals one NF279 molecule binding with the peptide. This NF279 molecule, which is expressed as Mol E, possesses various types of interactions with PAP$_{248–286}$. The side chains of Asn265 and Asn269 from the peptide form hydrogen bonds with one of the sulfonate groups of Mol E (Figure 4a and Table 1). The V-
shaped configuration allows Mol E to work as a clip to pinch the peptide. As a result, the guanidinium group of Arg257 enters into the crack and interacts with the sulfonate group and the oxygen atom of the amide group of Mol E through three hydrogen bonds. In addition, Arg257 and Arg273 are strongly attracted by the nearby anionic groups of Mol E through electrostatic attractions (Figure 4a and Table S1). Furthermore, there are hydrophobic and CH–π or NH–π interactions between PAP248–286 and Mol E (Table S3). Because NF279 adopts an allosteric strategy, this molecule has another linear configuration upon interaction with the peptide. Figure 4b shows the structure of PAP248–286 with two NF279 molecules (Mol E and Mol G). Mol G in the linear conformation is able to contact many residues of the peptide. For instance, Mol G has π–π and CH–π interactions with PAP248–286, from residues Ile249 to Tyr286 (Table S3). The long structure of Mol G makes it more likely to associate with the peptide. As a matter of fact, the linear configuration is predominantly found for NF279 molecules interacting with PAP248–286.

NF340, similar to NF110 and NF279, also contains aromatic rings, amide bonds, and sulfonate groups, but NF340 is smaller and shorter than the previous two molecules. The structures of PAP248–286 interacting with NF340 molecules are observed in Figure 1b and Figure 4c. Detailed information of the interactions can be found in Tables 1, S1, and S4. The results indicate that NF110 is the best compound among the three molecules investigated to interact with PAP248–286. The timeline of the number of hydrogen bonds exhibits that the PAP248–286–NF110 complex forms more hydrogen bonds than the other two peptide–compound systems in Figure S4. The number of other interactions of PAP248–286 with the compounds, including electrostatic and π–π interactions, are shown in Figure 4d.

2.3. Binding of PAP248–286 and Sulfonated Molecules

We know from the aforementioned results that PAP248–286 can interact with NF110, NF279, and NF340 through different kinds of interactions. However, it is still unclear how tightly these compounds bind with PAP248–286. The distance between the nitrogen atom of the ammonium group of Lys or the carbon atom of the guanidinium group of Arg with the nearest sulfur atom of the sulfonated molecules was measured, as the cation–residue interactions of PAP248–286 are attracted by the sulfonate groups of the compounds. Relative to the initial values, most of the distances listed in Table 2 are markedly lower, whereas the distances in the PAP248–286–NF279 and PAP248–286–NF340 complexes are greater than those in the PAP248–286–NF110 complex. It is suggested that the NF110 molecules bind with PAP248–286 more tightly than the other two derivatives. This indication is verified by the binding free energy. Table S5 shows the binding free energy (∆E) of the peptide–compound complexes. The ∆E value of PAP248–286–NF110 is the lowest and is followed by that of PAP248–286–NF279, whereas PAP248–286–NF340 has the highest binding free energy. Consequently, the PAP248–286–NF340 complex has the weakest binding affinity of the three peptide–compound complexes.

We also determined the abilities of these sulfonated compounds to bind PAP248–286 by surface plasmon resonance (SPR). As shown in Figure 5, all three sulfonated compounds bind to PAP248–286 but with different binding affinities. The equilibrium dissociation constant (Kd) of NF110 bound to PAP248–286 is 2.11 × 10⁻⁸ M with an association rate constant (kₐ) of 3.96 × 10⁹ M⁻¹ s⁻¹. Under the same conditions, NF279 bound to PAP248–286 with a Kd of 6.74 × 10⁻⁶ M and kₐ of 59.8 M⁻¹ s⁻¹, whereas NF340 bound to PAP248–286 with a Kd of 4.72 × 10⁻⁵ M. The Kd value of NF110 is 22.4-fold higher than that of NF340. This is in agreement with previous results.

### Table 1. Hydrogen-bonding interactions between sulfonated compounds and PAP248–286

| Entry | NF110 molecule | PAP248–286 residue | NF279 molecule | PAP248–286 residue | NF340 molecule | PAP248–286 residue |
|-------|----------------|-------------------|----------------|-------------------|----------------|-------------------|
| 1     | B              | Lys255            | B              | Arg273            | B              | Gln252            |
| 2     | Gln259         | C                 | Lys281         | D                 | Asn269         |
| 3     | Gly260         | D                 | Gln252         | Met271            |
| 4     | Asn269         | E                 | Arg257         | Lys281            |
| 5     | Lys272         | Arg273            | Arg257         | Arg257            | Gly260         |
| 6     | Arg273         | Asn265            | G              | Arg257            |
| 7     | Arg273         | Arg257            | Asn269         | Gly248            |
| 8     | C              | Arg257            | Arg273         | Arg273            |
| 9     | Arg257         | Arg257            | Lys282         | I                 | Gln276          |
| 10    | Arg273         | Ser256            | H              | Arg273            |
| 11    | Lys251         | Arg257            | Lys282         | Met285            |
| 12    | Ser256         | G                 | Lys282         | Leu283            |
| 13    | Lys272         | I                 | Arg257         | Tyr286            |
| 14    | F              | Arg273            | Ser256         |                  |
| 15    | Ser279         | I                 |                  |                  |
| 16    | H              | Arg273            |                  |                  |
| 17    | Arg273         | Ser256            |                  |                  |
| 18    | Lys281         |                  |                  |                  |
| 19    | Leu283         |                  |                  |                  |
| 20    | Tyr286         |                  |                  |                  |
| 21    | I              | Lys253            |                  |                  |
2.4. Sulfonated Molecules Inhibit the Aggregation of PAP

NF110, NF279, and NF340 were used to examine their abilities to inhibit PAP aggregation. We incubated PAP to detect fibril formation by Congo red staining in the absence and presence of sulfonated molecules. Congo red is a specific dye to detect amyloid, as it can bind amyloid fibrils and induce increased optical absorbance proportional to the level of fibrils. In the absence of sulfonated molecules, PAP aggregated following a lag phase of around 12 h. The Congo red absorbance signal increased gradually until it reached a plateau after 24 h. In the presence of a twofold excess amount of the sulfonated molecules, NF110, NF279, and NF340 all displayed inhibitory effects on PAP fibril formation, as indicated by reduced optical absorbance intensity at 48 h (Figure 5e and Table S6). Relative to the PAP control, NF110 showed a maximum decrease in absorbance intensity and NF340 showed only a slight decrease in the Congo red signals. Therefore, NF110 exhibited the strongest ability to inhibit PAP aggregation and NF279 showed a moderate effect on PAP fibrillization, whereas NF340 weakly inhibited PAP self-assembly.

3. Summary and Conclusions

In this work, we studied the binding of sulfonated compounds with prostatic acid phosphatase (PAP). Three molecules, NF110, NF279, and NF340, were selected from plenty of derivatives of suramin. These sulfonated compounds have different sizes and configurations, although their structures are related, as they all possess amide bonds, sulfonate groups, and aromatic rings. However, many problems remained unsolved, including how these compounds bind with PAP whether
the number of sulfonate groups affects peptide–compound interactions, and how tightly the compounds bind to \( \text{PAP}^{248–286} \). Thus, a computational approach would be useful to investigate protein–ligand interactions and to give us a hand to understand the binding of \( \text{PAP}^{248–286} \) with these sulfonated compounds.

\( \text{NF110} \), having four sulfonate groups, was found to have a distorted X-shaped configuration upon interaction with \( \text{PAP}^{248–286} \). Thus, \( \text{NF110} \) has four flexible arms and each arm has one sulfonate group at the end. As a result, \( \text{NF110} \) can contact the peptide from four different directions. This accounts for good interaction between the peptide and \( \text{NF110} \). \( \text{NF279} \) has two more sulfonate groups than \( \text{NF110} \), three on one tail of the compound and the other three on the other tail. Moreover, \( \text{NF279} \) introduces an allosteric strategy for folding itself into a V shape. V-shaped \( \text{NF279} \) looks like a clip to pinch the peptide. However, \( \text{NF279} \) only has sulfonate groups on both ends, regardless of the configuration it adopts. Finally, the smallest compound, \( \text{NF340} \), similar to \( \text{NF110} \) only has four sulfonate groups. The difference in these two compounds resides in the fact that all of the sulfonate groups of \( \text{NF340} \) are distributed equally on both tails. This is similar to that found for \( \text{NF279} \).

The structural properties of these compounds have a great influence on peptide–compound interactions.

As learned from the results, all of the sulfonated compounds were able to bind with \( \text{PAP}^{248–286} \) through a variety of interactions, including hydrogen-bonding interactions, hydrophobic interactions, \( \pi-\pi \) stacking interactions, and electrostatic interactions. The \( \text{NF110} \) molecules form 21 hydrogen bonds with \( \text{PAP}^{248–286} \) whereas the number of hydrogen bonds is 17 in the \( \text{PAP}^{248–286}–\text{NF279} \) complex and decreases to 13 in the \( \text{PAP}^{248–286}–\text{NF340} \) complex. The number of \( \pi-\pi \) or \( \text{CH}–\pi \) interactions in \( \text{PAP}^{248–286}–\text{NF110} \) is comparable to that in \( \text{PAP}^{248–286}–\text{NF279} \) but is higher than that in \( \text{PAP}^{248–286}–\text{NF340} \). Electrostatic contributions are crucial for binding due to strong attraction between the negatively charged sulfonate groups of the compounds and the positively charged residues of \( \text{PAP}^{248–286} \). From the above, molecules of \( \text{NF110} \) form the most interactions with \( \text{PAP}^{248–286} \) among the three compounds. This validates previous analyses indicating that the structural properties of the compounds were important. On the basis of the data, the interactions between \( \text{PAP}^{248–286} \) and the sulfonated compounds are not determined by the number of sulfonate groups. In contrast, the orientations of the sulfonate groups and the specific configurations of the compounds have a marked effect on binding.

The \( \text{NF110} \), \( \text{NF279} \), and \( \text{NF340} \) compounds occupy binding sites around \( \text{PAP}^{248–286} \). In particular, \( \text{NF110} \) is located on almost the entire exposed surface of \( \text{PAP}^{248–286} \) which leads to an open conformation of the peptide compared to the folded \( \text{PAP}^{248–286} \) system. The average radius of gyration for \( \text{PAP}^{248–286} \) exhibits the compactness of the peptide. In fact, these values increase for all three systems, and \( \text{PAP}^{248–286}–\text{NF110} \) has the largest value of 1.60 nm. Changes in the conformations of \( \text{PAP}^{248–286} \) also support the idea that the \( \text{NF110} \) molecules interact more effectively than the two other compounds with the peptide. In addition, the distances between the cationic residues of \( \text{PAP}^{248–286} \) and the anionic sulfonate groups of the compounds are shortened. The \( \text{PAP}^{248–286}–\text{NF110} \) complex shows the strongest binding affinity of the three peptide–compound complexes due to the fact that it has the most negative binding free energy of all the compounds. The surface plasmon resonance data agree with the results. Moreover, the critical Gly261–Tyr286 fragment of \( \text{PAP}^{248–286} \), interacting with the sulfonated compounds, no longer forms \( \beta \)-sheet structures with other \( \text{PAP}^{248–286} \). It blocks the formation of SEVI (semen-derived enhancer of virus infection) fibrils. A Congo red staining experiment confirmed this analysis.

In consequence, we suggest that \( \text{NF110} \) and its analogues would be promising microbicides to antagonize seminal amyloids. The results will give us useful information to develop novel target selective inhibitors with high efficiency to block HIV transmission. However, more biophysical and biochemical studies are essential to provide in-depth investigation. For example, NMR and HSQC spectra should be included to illustrate the interactions between the compounds and \( \text{PAP}^{248–286} \), and circular dichroism analysis, transmission electron microscopy, and a viral infection assay should be applied to confirm the effects of the compounds on the inhibition of fibril formation. In addition, further in vivo studies in rhesus macaques model to examine the anti-HIV activity would be recommended.

| Residue | Distance [nm] | Representative structure |
|---------|--------------|-------------------------|
| Lys251  | 1.3429 ± 0.0465 | 0.3925 ± 0.0850 |
| Lys253  | 1.0459 ± 0.1169 | 0.3942 ± 0.0691 |
| Lys255  | 0.7601 ± 0.1097 | 0.5684 ± 0.0844 |
| Arg257  | 1.0460 ± 0.0935 | 0.5975 ± 0.0931 |
| Lys272  | 0.4315 ± 0.1184 | 0.3875 ± 0.1130 |
| Arg273  | 1.6453 ± 0.1187 | 0.4186 ± 0.0988 |
| Lys281  | 0.6780 ± 0.1059 | 0.3839 ± 0.1031 |
| Lys282  | 1.2894 ± 0.1076 | 0.5297 ± 0.0927 |
| Lys281  | 2.9181 ± 0.0898 | 0.5450 ± 0.1004 |
| Lys253  | 1.2734 ± 0.0866 | 0.4095 ± 0.1094 |
| Lys255  | 0.8940 ± 0.1082 | 0.5277 ± 0.0150 |
| Arg257  | 0.6510 ± 0.0914 | 0.4558 ± 0.1164 |
| Lys272  | 0.5835 ± 0.0865 | 0.7359 ± 0.0818 |
| Arg273  | 1.0045 ± 0.0156 | 0.6043 ± 0.0841 |
| Lys281  | 1.9025 ± 0.1183 | 0.3734 ± 0.1184 |
| Lys282  | 2.4143 ± 0.0895 | 0.4127 ± 0.1126 |
| Lys251  | 3.5621 ± 0.0407 | 0.3698 ± 0.0913 |
| Lys253  | 2.4326 ± 0.0929 | 0.7635 ± 0.0975 |
| Lys255  | 0.5769 ± 0.0973 | 1.3823 ± 0.0638 |
| Arg257  | 2.0381 ± 0.1154 | 0.6489 ± 0.1146 |
| Lys272  | 0.6953 ± 0.0758 | 0.5733 ± 0.0954 |
| Arg273  | 0.6836 ± 0.1076 | 0.4190 ± 0.0905 |
| Lys281  | 0.7686 ± 0.1143 | 0.3530 ± 0.0723 |
| Lys282  | 2.2401 ± 0.0395 | 0.8556 ± 0.1165 |
Methods

Computational Methods

The structure of PAP<sub>248–286</sub> was gained from the Protein Data Bank (PDB ID 2L3H).<sup>[33]</sup> The NF110, NF279, and NF340 compounds were constructed and optimized by using the Gaussian 09 program package<sup>[34]</sup> without any geometrical constraints.<sup>[35–38]</sup> Molecular docking procedures were implemented by Autodock Vina 1.5.6 software<sup>[39]</sup> to investigate the binding of NF110, NF279, and NF340 to PAP<sub>248–286</sub>. The spacing was set to 1.00 Å. There are eight positively charged amino-acid residues (Lys<sub>251</sub>, Lys<sub>253</sub>, Lys<sub>255</sub>, Arg<sub>257</sub>, Lys<sub>272</sub>, Arg<sub>273</sub>, Lys<sub>281</sub>, and Lys<sub>282</sub>) in PAP<sub>248–286</sub>. Because the Arg and Lys residues of PAP<sub>248–286</sub> can attract the anionic sulfonate groups of NF110, NF279, and NF340, we investigated the following four systems: PAP<sub>248–286</sub> with 8 NF110 molecules, PAP<sub>248–286</sub> with 8 NF279 molecules, PAP<sub>248–286</sub> with 8 NF340 molecules, and PAP<sub>248–286</sub> alone. The side chains of Lys and Arg were set as flexible. On the basis of the interactions and binding energies, the most promising poses were chosen. The structures of the molecules bound to PAP<sub>248–286</sub> were preliminarily energy minimized by YASARA.<sup>[40]</sup> Molecular dynamics (MD) simulations were utilized to investigate the binding of PAP<sub>248–286</sub> and the sulfonated compounds. GROMACS 4.5.3<sup>[41]</sup> with CHARMM27<sup>[42,43]</sup> force field were used for all the systems. The Swissparam server produced the force field parameters of the NF110, NF279, and NF340 molecules.<sup>[44,45]</sup> The peptide–ligand complex was immerged in a cubic box, and the box size was extended 12 Å from the edges of the complex. Then, the box was filled with TIP3P water.<sup>[46]</sup> Sodium and chloride were also added for neutralization and to offer physiological conditions. After that, the system was energy minimized through 3000 steps of steepest descent. The structure produced from the minimization was prepared for MD simulations. The 100 ns MD simulations were implemented with the NPT ensemble, which was used in our previous paper.<sup>[47–49]</sup> The bonds and water were constrained by the LINCS algorithm<sup>[50]</sup> and SETTLE algorithm<sup>[51]</sup> respectively. The Particle-Mesh Ewald was utilized to measure the electrostatic interactions of long range.<sup>[52]</sup> A pressure of 0.1 MPa and a temperature of 300 K were applied. The time step was 2 fs. We utilized the GROMACS tools to analyze the MD trajectories. YASARA<sup>[53,54]</sup> and PyMOL<sup>[55,56]</sup> programs were used in this paper to visualize and prepare the structures. The root-mean-square deviations (RMSDs) evaluate the deviation of the backbone of the peptide. Molecular mechanics/Poisson–Boltzmann surface area (MM/PBSA), which was previously employed in many...
protein–ligand systems, was utilized to study the binding free energy.\textsuperscript{[25–46]}

Experiments

The NF110, NF279, and NF340 compounds were bought from Tocris Bioscience. Scilligent-Peptide (Beijing, China) synthesized PAP\textsuperscript{248–286} (>95 % purity).

SPR Binding Kinetics of PAP\textsuperscript{248–286}—Molecule Interactions

A PlexArray HT system was used to analyze the SPR measurements. The experiment was performed by using the same method as that in our published paper.\textsuperscript{[46]} Details can be found in the Supporting Information.

Congo Red Binding Assay

The fibril formation kinetics of PAP\textsuperscript{248–286} were determined by a Congo red binding assay.\textsuperscript{[46]} For the experimental procedures, see to our previous paper\textsuperscript{[46]} and the Supporting Information.

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Conflict of interest

The authors declare no conflict of interest.

Keywords: amyloid fibrils · HIV · inhibitors · sulfur · viruses

[1] J. Münch, E. Rucker, L. Standker, K. Adermann, C. Goffinet, M. Schindler, S. Wildum, R. Chinnadurai, D. Rajan, A. Specht, G. Gimenez-Gallego, P. C. Sanchez, D. M. Fowler, A. Koulov, J. W. Kelly, W. Mothes, J. C. Grivel, L. Margolis, O. T. Keppler, W. G. Forssmann, F. Kirchhoff, Cell 2007, 131, 1071.
[2] S. M. Usmani, O. Zirafi, J. A. Muller, N. L. Sandi-Monroy, J. K. Yadav, C. Meier, T. Weil, N. R. Roan, W. C. Greene, P. Walther, K. P. Nilsson, P. Harmsstroem, R. Wetzl, C. D. Pilcher, F. Gaggsteiger, M. Fandrich, F. Kirchhoff, J. Münch, Nat. Commun. 2014, 5, 3508.
[3] K. A. Kim, M. Yolamanova, O. Zirafi, N. R. Roan, L. Staendker, W. G. Forssmann, A. Burgener, N. Dejucq-Rainsford, B. H. Hahn, G. M. Shaw, W. C. Greene, F. Kirchhoff, J. Münch, Retrovirology 2010, 7, 55.
[4] N. R. Roan, J. Münch, N. Arhel, W. Mothes, J. Niederman, A. Kobayashi, K. Smith-McCune, F. Kirchhoff, W. C. Greene, J. Virol. 2009, 83, 73–80.
[5] T. Xun, W. Li, J. Chen, F. Yu, W. Xu, Q. Wang, R. Yu, X. Li, X. Zhou, L. Lu, S. Jiang, L. Li, S. Tan, S. Liu, Antimicrob. Agents Chemother. 2015, 59, 5123–5134.
[6] R. Nunes, B. Sarmento, J. das Neves, J. Controlled Release 2014, 194, 278–294.
[7] J. Baeten, C. Celum, Annu. Rev. Med. 2013, 64, 219–232.
[8] Y. H. Lee, A. Ramamoorthy, Protein Sci. 2018, https://doi.org/10.1002/pro.3395.
[36] T. Zhang, M. Ozbil, A. Barman, T. J. Paul, R. P. Bora, R. Prabakar, Acc. Chem. Res. 2015, 48, 192 – 200.
[37] T. T. Zhang, X. X. Zhu, R. Prabakar, Organometallics 2014, 33, 1925 – 1935.
[38] T. Zhang, X. Zhu, R. Prabakar, J. Phys. Chem. B 2014, 118, 4106 – 4114.
[39] O. Trott, A. J. Olson, J. Comput. Chem. 2010, 31, 455 – 461.
[40] E. Krieger, K. Joo, J. Lee, J. Lee, S. Raman, J. Thompson, M. Tyka, D. Baker, K. Karplus, Proteins 2009, 77, 114 – 122.
[41] D. Van Der Spoel, E. Lindahl, B. Hess, G. Groenhof, A. E. Mark, H. J. Berendsen, J. Comput. Chem. 2005, 26, 1701 – 1718.
[42] P. Bjelkmar, P. Larsson, M. A. Cundet, B. Hess, E. Lindahl, J. Chem. Theory Comput. 2010, 6, 459 – 466.
[43] A. D. Mackerell, Jr., M. Feig, C. L. Brooks, J. Comput. Chem. 2004, 25, 1400 – 1415.
[44] D. Bier, R. Rose, K. Bravo-Rodriguez, M. Bartel, J. M. Ramirez-Anguita, S. Dutt, C. Wilch, F. G. Klarner, E. Sanchez-Garcia, T. Schrader, C. Ottmann, Nat. Chem. 2013, 5, 234 – 239.
[45] V. Zoete, M. A. Cundet, A. Grosdidier, O. Michielin, J. Comput. Chem. 2011, 32, 2359 – 2368.
[46] W. L. Jorgensen, J. Chandrasekhar, J. D. Madura, R. W. Impey, M. L. Klein, J. Chem. Phys. 1983, 79, 926 – 935.
[47] T. Zhang, Y. Tian, Z. Li, S. Liu, X. Hu, Z. Yang, X. Ling, S. Liu, J. Zhang, J. Chem. Inf. Model. 2017, 57, 2281 – 2293.
[48] Y. Tian, Y. Yu, Y. Shen, H. Wan, S. Chang, T. Zhang, S. Wan, J. Zhang, J. Chem. Inf. Model. 2017, 57, 977 – 987.
[49] E. S. Song, M. Ozbil, T. Zhang, M. Sheetz, D. Lee, D. Tran, S. Li, R. Prabhakar, L. B. Hersch, D. W. Rodgers, PLoS One 2015, 10, e0133114.
[50] B. Hess, H. Bekker, H. J. C. Berendsen, J. G. E. M. Fraaije, J. Comput. Chem. 1997, 18, 1463 – 1472.
[51] S. Miyamoto, P. A. Kollman, J. Comput. Chem. 1992, 13, 952 – 962.
[52] T. Darden, D. York, L. Pedersen, J. Chem. Phys. 1993, 98, 10089 – 10092.
[53] E. Krieger, G. Koraimann, G. Vriend, Proteins 2002, 47, 393 – 402.
[54] E. Krieger, G. Vriend, Bioinformatics 2002, 18, 315 – 318.
[55] W. L. DeLano, Abstr. Pap. Am. Chem. Soc. 2004, 228, U313 – U314.
[56] W. L. DeLano, J. W. Lann, Abstr. Pap. Am. Chem. Soc. 2005, 230, U1371 – U1372.
[57] N. A. Baker, D. Sept, S. Joseph, M. J. Holst, J. A. McCammon, Proc. Natl. Acad. Sci. USA 2001, 98, 10037 – 10041.
[58] M. Dong, H. Li, D. Hu, W. Zhao, X. Zhu, H. Ai, ACS Chem. Neurosci. 2016, 7, 599 – 614.
[59] M. Dong, W. Zhao, D. Hu, H. Ai, B. Kang, ACS Chem. Neurosci. 2017, 8, 1577 – 1588.
[60] R. Kumari, R. Kumar, Open Source Drug Discovery Consortium, A. Lynn, J. Chem. Inf. Model. 2014, 54, 1951 – 1962.
[61] L. Li, L. He, S. Tan, X. Guo, H. Lu, Z. Qi, C. Pan, X. An, S. Jiang, S. Liu, Antimicrob. Agents Chemother. 2010, 54, 1700 – 1711.
[62] M. R. Nilsson, Methods 2004, 34, 151 – 160.
[63] S. Tan, L. Lu, L. Li, J. Liu, Y. Oksov, H. Lu, S. Jiang, S. Liu, PLoS One 2013, 8, e59777.

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