**N-Methylation as a Strategy for Enhancing the Affinity and Selectivity of RNA-binding Peptides: Application to the HIV-1 Frameshift-Stimulating RNA**

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**ABSTRACT:** Human Immunodeficiency Virus (HIV) type 1 uses a −1 programmed ribosomal frameshift (−1 PRF) event to translate its enzymes from the same transcript used to encode the virus’ structural proteins. The frequency of this event is highly regulated, and significant deviation from the normal 5–10% frequency has been demonstrated to decrease viral infectivity. Frameshifting is primarily regulated by the Frameshift Stimulatory Signal RNA (FSS-RNA), a thermodynamically stable, highly conserved stem-loop that has been proposed as a therapeutic target. We describe the design, synthesis, and testing of a series of N-methyl peptides able to bind the HIV-1 FSS RNA stem loop with low nanomolar affinity and high selectivity. Surface plasmon resonance (SPR) data indicates increased affinity is a reflection of a substantially enhanced on rate. Compounds readily penetrate cell membranes and inhibit HIV infectivity in a pseudotyped virus assay. Viral infectivity inhibition correlates with compound-dependent changes in the ratios of Gag and Gag-Pol in virus particles. As the first compounds with both single digit nanomolar affinities for the FSS RNA and an ability to inhibit HIV in cells, these studies support the use of N-methylation for enhancing the affinity, selectivity, and bioactivity of RNA-binding peptides.

**H**uman Immunodeficiency Virus type 1 (HIV-1), like many retroviruses, encodes some of its proteins in an overlapping open reading frame (ORF) that does not contain its own ribosomal initiation site. This overlapping ORF can only be translated by a programmed ribosomal frameshifting event. In HIV-1, the ORF encoding the polyprotein for viral enzymes (Pol) is in a −1 reading frame relative to that encoding the structural proteins (Gag). Pol is only translated with a frequency of 5–10% of Gag, reflecting how often frameshifting occurs. Deviation from this narrow range causes defects in the virus’ ability to infect new cells. This was robustly demonstrated by Shehu-Xhilaga et al. in 2001. Using a reverse transcriptase activity assay as a marker for infectivity in 293T cells, viral fitness was found to degrade by approximately 100-fold as the Gag/Gag-Pol ratio changed from 20:1 to 20:21. These viral particles were also unable to correctly dimerize their genomic RNA, although recent work suggests that the frameshift site is not directly involved in packaging when the proper ratios of Gag/Gag-Pol are provided in trans. Reduction in frameshift efficiency likewise interferes with viral replication. Control of frameshifting in HIV-1 occurs via a frameshift stimulatory signal (FSS), a structured piece of the viral mRNA. Extensive analysis of this element, including NMR solution structures and chemical probing using SHAPE, suggests that a key structural feature of the HIV-1 FSS is a thermodynamically stable stem loop (Figure 1, boxed upper region). This is preceded by a heptameric, uridine-rich “slippery sequence,” where the frameshift actually occurs, and other dynamically regulated regions of secondary structure. The stem loop is highly conserved in HIV-1, with over 85% of all isolates containing the rare ACA tetraloop motif, and all subtypes of group M are predicted to fold into identical structures.

**Supporting Information**

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secondary structures with a nearly 100% conservation of base pairs. Due to the virus’ stringent control of the frequency of frameshifting, it has been suggested that small molecules able to bind the FSS-RNA and modulate this process could point the way to future HIV-1 therapeutics, as well as serving as prototypes for small molecules able to interfere with recoding events in other viruses. Efforts to identify such compounds have been complicated by the general challenge posed by sequence-selective RNA recognition. While a bisbenzamide, doxorubicin, and guanidinoneomycin have been examined as FSS RNA binders, their relatively low affinities and lack of specificity have limited their applicability as probes or drug leads. Building on the discovery of an HIV-1 FSS RNA binder identified via Resin-Bound Dynamic Combinatorial Chemistry, we recently reported the first synthetic compound able to bind the HIV-1 FSS RNA with a submicromolar dissociation constant and inhibit pseudotyped HIV infectivity. Compound 1Z was found to bind FSS RNA with a $K_D$ of approximately 70 nM, was cell permeable, and strongly inhibited the infectivity of pseudotyped HIV in a dose-dependent manner. While successful, the selectivity of the 1Z—FSS RNA binding interaction was modest (5:1 vs total yeast tRNA). In order to access compounds with enhanced anti-HIV activity and selectivity, we sought a simple modification that would be compatible with our synthetic scheme.

Table 1. Fluorescence Titrations Confirm N-Methyl Analogs Have Higher Affinity and Selectivity for HIV-1 FSS RNA

| compound | FSS-RNA $K_D$ (nM) | FSS-DNA $K_D$ (nM) | yeast tRNA $K_D$ (nM) | FSS RNA: FSS DNA selectivity | FSS RNA: yeast tRNA selectivity |
|----------|---------------------|---------------------|------------------------|-----------------------------|--------------------------------|
| 1E       | 100 ± 30            | 230 ± 20            | 220 ± 30               | 2.3:1                       | 2.2:1                          |
| 1Z       | 66 ± 30             | 120 ± 20            | 330 ± 30               | 1.8:1                       | 1:1                            |
| 2E       | 2.7 ± 1.0           | 81 ± 30             | 120 ± 40               | 30:1                        | 44:1                           |
| 2Z       | 2.7 ± 0.7           | 65 ± 10             | 180 ± 20               | 24:1                        | 67:1                           |
| 3E       | 3.0 ± 0.1           | 79 ± 10             | 90 ± 10                | 26:1                        | 30:1                           |
| 3Z       | 1.1 ± 0.5           | 60 ± 8.0            | 67 ± 10                | 55:1                        | 61:1                           |
| 4E       | 5.0 ± 2.0           | 34 ± 10             | 91 ± 20                | 6.8:1                       | 18:1                           |

Dissociation constants ($K_D$) are reported in nanomolarity. Titrations were carried out in 20 mM HEPES, 150 mM NaCl, and 5 mM MgCl$_2$. Each measurement was taken in triplicate to assure equilibrium was reached, and each titration was repeated twice. Data were fit to a single-site binding model. Error reported is the distance between values from complete titrations.
mediated degradation.\textsuperscript{27} As such, we hypothesized that introduction of N-methyl groups could bias the conformational ensemble of FSS-binding compounds favorably, resulting in higher affinity and enhanced frameshift-stimulating activity.

To test this hypothesis, a series of three different symmetrical methylation patterns were targeted. Combined with the potential to generate $E$ and $Z$ olefin isomers during synthesis, this would yield six new compounds. To that end, compounds $2\text{–}4E$ and $-Z$ were synthesized on a solid phase, via a scheme incorporating N-methylation methodology reported by Miller and Scanlan.\textsuperscript{28} Five of the six targeted compounds were obtained in quantity; cross-metathesis to produce the tetra N-methyl compound did not yield a sufficient quantity of the $Z$ isomer (4Z) to permit characterization.

\section*{RESULTS}

\textbf{N-Methylation Increases Binding Affinity and Selectivity for FSS-RNA.} The benzo[G]quinoline moiety of compounds 1–4 is fluorescent and quenches on binding DNA or RNA.\textsuperscript{29} As such, fluorescence titration provides a simple, direct means for determining compound affinity. Dissociation constants ($K_D$) measured for compounds 1Z–4E are reported in Table 1; titration curves are provided in the Supporting Information. In order to evaluate sequence selectivity, binding constants were also measured for each compound to total yeast tRNA (a standard selectivity measure)\textsuperscript{30} and to a DNA sequence homologous to the FSS RNA. The concentration-independent melting profile of the latter suggests that it is also a hairpin or hairpin-like structure (Supporting Information). These results indicate that N-methylation enhanced compound affinity for the FSS-RNA relative to 1E and 1Z by as much as 100-fold and improved the selectivity relative to tRNA by 6- to 65-fold.

To confirm the high affinity binding and to determine if this resulted from changes in binding kinetics, compounds 3Z and 4E were further investigated by surface plasmon resonance. Compared to the previously reported compound 1Z, N-methyl compounds 3Z and 4E both show an increase in $k_1$ with little to no change in $k_2$ (Table 2 and Supporting Information). This result is consistent with the hypothesis that N-methylation alters the conformational ensemble such that conformers close to the bound form are favored ($k_2$) but that the mode of binding is the same (no change in $k_1$). The small difference in $K_D$ values measured between the two techniques is not unexpected, as SPR utilizes a surface immobilized RNA as opposed to the fully solution-phase fluorescence titration.

\textbf{N-Methylation Increases Cell Permeability.} We used flow cytometry to assay the effect of N-methylation on the ability of HIV FSS-targeted compounds to cross cell membranes. Dose-dependent cell toxicity was first measured using a standard WST-1 proliferation assay\textsuperscript{31} (Supporting Information S20) in HEK293T cells. These experiments established LD$_{50}$ values for compounds 1Z–4E in the range of 20–30 μM. For flow cytometry experiments, HEK293 cells were incubated with either 5 or 15 μM of each compound. Cells treated with N-methyl compounds show an increased amount of fluorescence at an equivalent concentration to unmethylated compound 1Z, suggesting that N-methylation enhances the ability of HIV RNA-targeted compounds to cross the cell membrane (Figure 2).\textsuperscript{32,33} The rank order of compound penetration (Figure 2) correlated strongly with differences in cLogP, with less-polar compounds displaying better permeability. Differences in permeability are also readily observable via fluorescence microscopy (Supporting Information).

\textbf{N-Methylation Increases Antiviral Potency.} To assess the effect of compounds 2E–4E on viral fitness, an infectivity assay was performed with pseudotyped HIV.\textsuperscript{34,23} HEK293T producer cells were transfected with a pro-viral plasmid for HIV in which the nef gene is replaced with green fluorescent protein and the env gene is deleted. A plasmid coding for VSV-G was cotransfected to allow for production of single cycle infectious virions. Producer cells were incubated with the compound for 24 h, after which the virions were harvested and normalized for viral load based on p24 ELISA. Equal loads of these viral particles were then incubated with TZM-bl reporter cells that express luciferase under long terminal repeat (LTR) promoter control when infected by HIV-1. All new compounds with the exception of 2E showed significantly greater antiviral activity in this assay than our previously reported compound 1Z, with compound 4E exhibiting the most potent activity (Figure 3). Data shown in the figure are a composite from two experiments; additional data may be found in the Supporting Information. While factors governing activity in this assay are complex, the enhanced potency of the N-methyl compounds likely is a function of both their enhanced affinity for the FSS RNA and improved cell permeability. To ensure reproducibility of the observed biological effect, infectivity inhibition assays were repeated for select compounds in a second laboratory (Supporting Information S27).

\begin{table}[h]
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\small
\begin{tabular}{|c|c|c|c|c|}
\hline
compound & $K_D$ (nM) & $k_1$ (1/M/s) & $k_2$ (1/s) \\
\hline
1Z$^a$ & 71 & $(2.2 \pm 0.8) \times 10^4$ & $(1.4 \pm 0.1) \times 10^{-3}$ \\
3Z$^b$ & 6.7 ± 3.7 & $(3.2 \pm 1.5) \times 10^4$ & $(2.7 \pm 1.8) \times 10^{-3}$ \\
4E$^c$ & 13 ± 5.2 & $(1.6 \pm 0.7) \times 10^4$ & $(1.9 \pm 0.5) \times 10^{-3}$ \\
\hline
\end{tabular}
\caption{SPR Confirms High Affinity Binding and Increased On Rate}
\end{table}

\textsuperscript{a}The values for compound 1Z were previously reported.\textsuperscript{19} SPR was carried out in 150 mM NaCl and 10 mM HEPES. Injections were repeated twice for consistency. Curves were fit individually to a mass transfer model, and the error reported is standard deviation of all fits.

\textbf{N-Methyl Compounds Increase Gag-Pol Levels in Virions.} We next assayed viral protein production as a function of added compounds 2–4 to determine if inhibition of

\begin{figure}[h]
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\includegraphics[width=\textwidth]{figure2.png}
\caption{Cell permeability for HIV FSS-binding compounds, as measured by flow cytometry. HEK293T cells were incubated for 24 h in the presence of compound. Cells were then gated for negative propidium iodide and positive compound fluorescence. The mean intensity was calculated and plotted for each concentration (blue, 5 μM; red, 15 μM; dashed lines are provided for comparison to 1Z). Error bars represent the rCV (robust coefficient of variation, or standard deviation divided by the median).
\end{figure}
Articles

infectivity correlated with increased amounts of frameshift products. Viral particles were collected and their proteins analyzed by SDS-PAGE and Western blotting. The primary antibody reacted with the blots recognized p160, the full length Gag-Pol polyprotein that is only made if the −1 frameshift occurs, as well as with p55 and p24, two forms of Gag that are made with or without frameshifting (Figure 4). The ratio of p160 to p24+p55 was determined as a semiquantitative measure of frameshifting. We observed that this ratio increases in a dose-dependent manner and correlated strongly with the ability of individual compounds to inhibit the infectivity of pseudotyped virions. As these ratios could be affected by packaging bias, we also examined Gag/Gag-Pol ratios directly in the cell lysates of producer cells (Supporting Information). We observed similar trends to those obtained from virus particles. In tandem, these results are consistent with a direct concentration-dependent enhancement of FSS RNA-binding compounds on frameshifting.

To control for the possibility that FSS-binding compounds might also interact with viral protease, results were compared with an analogous assay including the FDA-approved protease inhibitor, indinavir. We observed that indinavir increased the amount of p55 visible on the blot, while dramatically decreasing the amount of p24. No buildup of p160 was observed. These results strongly contrast with those observed for compounds 2–4. We further confirmed that compound activity is independent of HIV protease via a commercial protease inhibition assay (Supporting Information Figure S-9). The compounds do not show any significant protease inhibition up to 100 μM.

DISCUSSION

While several authors have hypothesized that compound-dependent frameshift enhancement could serve as an effective means of limiting viral infectivity, the lack of high affinity, sequence-selective frameshift RNA-binding compounds has made testing this hypothesis a challenge. Compounds we reported previously reduced the infectivity of pseudotyped virions and enhanced frameshifting, but their modest selectivity for the FSS RNA suggested that the antiviral activity could be due to a variety of mechanisms. Here, we find that antiviral activity and changes in Gag/Gag-Pol in produced pseudotyped virus particles strongly correlates with affinity for the HIV-1 FSS RNA, as well as cell permeability.

The enhanced on rates observed by SPR for 3Z and 4E relative to the non-N-methylated compound 1Z are intriguing. Very little is known about the relationship of binding kinetics and selectivity for RNA-targeted molecules. This is in contrast to proteins, for which a consensus is emerging that selectivity and efficacy are direct functions of the off rate (frequently described as the “residence time”). For RNA, and nucleic acids generally, such a broad generalization is probably not possible, as there are clearly nonselective binders (for example, many intercalators) with slow off-rates. Here, we observe an increase in affinity and selectivity for compounds with enhanced on-rates, but comparable off-rates. We hypothesize that, as with other N-methyl peptides, the enhanced on-rates are the result of differences in the ground-state conformational ensemble in which the proportion of binding-favorable conformers are enhanced. Structural studies designed to test that hypotheses are in progress.

To date, these compounds show the highest affinity for the FSS-RNA and are the only class of molecules that alters the products of frameshifting and inhibits viral infectivity. Tetramethylated compound 4E has the greatest increase in

Figure 3. N-methyl compounds demonstrate enhanced antiviral potency. HEK293T producer cells were transfected for 4 h with pro-viral and VSV-G plasmids to produce single cycle infectious particles. The producer cells were then incubated for 24 h in the presence of compound, in triplicate for each condition. Virions were harvested, normalized, then incubated with TZM-bl reporter cells. Luciferase production from the reporter cells was measured and normalized to values obtained from control cells. Error bars indicate standard deviation.

Figure 4. Virions collected and their proteins harvested and run on an SDS-PAGE gel. The transferred blot was then probed with an anti-p24 antibody, which will mark p24, as well as its precursor proteins p55 and p160. The ratio of p160 to p24+p55 was calculated by scanning densitometry (UVP VisionWorks software). The 20 μM lane for 4E was loaded with half as much protein due to a limiting quantity of virions.
hydrophobicity relative to 1 and also was the most cell-permeable. Further, 4E showed the greatest ability to inhibit the production of infectious virions. Interestingly, 4E also affected not only the ability to make infectious virions but also severely limited the intracellular replication of the virus, as can be visualized via GFP production from the proviral plasmid in HEK 293T producer cells (Figure 5). Thus, both the compound interferes with production of virus particles and the particles produced are less infective. While the availability of the compound increases with hydrophobicity, and appears to be the major limiting factor for antiviral effects, the discrepancy between compound 2E and 3E indicates that the increase in hydrophobicity is not the only governing factor for cell penetration. Based on the various patterns of N-methylation, it is not unreasonable to hypothesize that the differences are due to changes in the conformational ensembles of these molecules. More broadly, these results support the general hypothesis that synthetic molecules can have useful biological effects, when targeted to RNAs believed to have regulatory roles in protein recoding. Given the breadth of critical RNA-regulated frameshifting events discovered in viruses, 38–40 bacteria, 41,42 and even humans, 43,44 further exploration of this strategy in the context of understanding basic biology and the development of therapeutics is warranted.

**MATERIALS AND METHODS**

FSS-RNA and DNA were purchased from Integrated DNA Technologies and assessed for purity and structure using thermal melting protocols. Yeast tRNA was purchased from Sigma-Aldrich and used without further purification. Amino acids were purchased from P3Bio and used as received.

**Fluorescence Titration.** Fluorescence titrations were carried out in 20 mM HEPES, 150 mM NaCl, and 5 mM MgCl₂ using a Horiba FluoroMax4 fluorometer. All titrations started at a volume of 500 μL with the compound at 20–500 nM. The concentration was chosen based on a preliminary titration using 1 μM compound to get a first approximation of the binding affinity and then lowered to more accurately measure the affinity. RNA was then titrated in from a high concentration stock (0.5–10 μM for HIV-1 FSS RNA; 10 μM for HIV-1 FSS DNA and tRNA) in 1–10 μL increments. After RNA was added, the solution was thoroughly mixed in the cuvette via pipetting and allowed to stand for 10 min to reach equilibrium. Each measurement was taken three times with a 1 min waiting period between scans to confirm equilibrium was reached. Intensities were corrected for dilution. All titrations were completed twice.

**Surface Plasmon Resonance.** SPR was conducted using a Biacore X (Biacore, Inc.). Compounds were injected at a flow rate of 30 μL/min in 10 mM HEPES and 150 mM NaCl for 2 min. Flow was then maintained for at least 1 h to allow for dissociation. Each injection was repeated twice for consistency. Each trace was fit individually to a mass transfer model.

**Cell Toxicity.** HEK293T cells were plated at 1 × 10⁴ cells/well in a 96 well plate in DMEM with 10% fetal bovine serum and 1% anti–anti (penicillin-streptomycin + fungizone) at 37 °C. After cells were allowed to adhere for 6 h, they were then incubated with the compound for 24 h in triplicate. A total of 10 μL of WST-1 premix (Clontech) was added and incubated for 1 h followed by measurement using a PerkinElmer EnSpire plate reader.

**Flow Cytometry.** HEK293T cells were grown in DMEM with 10% fetal bovine serum and 1% penicillin-streptomycin at 37 °C to 80% confluency in a 12-well plate and treated with the compound for 24 h. Cells were trypsinized, pelleted, and washed twice with DPBS (Gibco). Cells were then resuspended in 300 μL ice cold DPBS and incubated with 5 μL of propidium iodide to stain dead cells. A total of 10 000 events were collected using a BD LSR-II flow cytometer.

**HIV-1 Infectivity.** The antiviral activity of all RNA-targeted compounds was measured by single-round infectivity assay with pseudotyped HIV-1 using HEK293T producer cells. The HIV-1 proviral vector (pDHIV3-GFP) includes all HIV-1 NL4–3 genes except nef (replaced with GFP) and env, thus preserving gag and pol, and the frameshift required for production of the Gag-Pol polyprotein. A single-round infectivity assay was conducted by transient transfection of the viral vector with VSV-G coat protein vector at a ratio of 1:0.5 using Fugene HD (Promega). The virus producer cells were dosed with compounds 4 h after transfection, and viral particles were harvested from the media 24 h after transfecting by filtering through a 0.45-μm syringe filter. Viral load was normalized with a p24 ELISA (PerkinElmer). The infections were performed using TZM-bl reporter cells containing stably integrated firefly luciferase, the expression of which is driven by the HIV-LTR promoter. Therefore, luciferase is expressed upon successful HIV infection. Triplicate infections in 96-well plates at 10 000 cells/well with 500 pg p24/well proceeded for 48 h before the addition of SteadyGlo reagent (Promega) to each well for 30 min. Luminescence was measured as a quantitative metric for changes in viral infectivity in the presence of a compound.

**Protease Inhibition.** The protease inhibition kit was purchased from Protein One and used as instructed. HIV-1 Protease was incubated at RT with no compound, a protease inhibitor, or compound for 90 min in the presence of the peptide FRET substrate. The protease inhibition kit was purchased from Protein One and used as instructed. HIV-1 Protease was incubated at RT with no compound, a protease inhibitor, or compound for 90 min in the presence of the peptide FRET substrate. The infections were performed using TZM-bl reporter cells containing stably integrated firefly luciferase, the expression of which is driven by the HIV-LTR promoter. Therefore, luciferase is expressed upon successful HIV infection. Triplicate infections in 96-well plates at 10 000 cells/well with 500 pg p24/well proceeded for 48 h before the addition of SteadyGlo reagent (Promega) to each well for 30 min. Luminescence was measured as a quantitative metric for changes in viral infectivity in the presence of a compound.

**ASSOCIATED CONTENT**

**Supporting Information**

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acschembio.5b00682.

Synthesis of compounds, compound characterization, fluorescence titrations, characterization of nucleic acid (DNA and RNA) sequences, surface plasmon resonance measurements, cell toxicity measurements, protease inhibition assay, flow cytometry data, fluorescence microscopy, additional Western blotting experiments, and replicate viral infectivity assay data (PDF)

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**Notes**

The authors declare no competing financial interest.

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