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Comparative Structural Effects of HIV-1 Gag and Nucleocapsid Proteins in Binding to and Unwinding of the Viral RNA Packaging Signal

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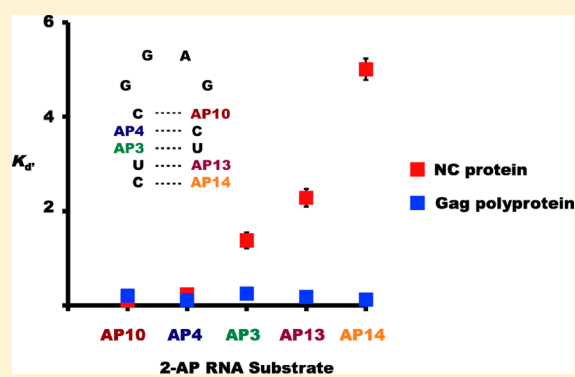
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S Supporting Information

ABSTRACT: The major RNA binding region of the HIV-1 Gag polyprotein is the nucleocapsid (NC) domain, which is responsible for the specific capture of the genomic RNA genome during viral assembly. The Gag polyprotein has other RNA chaperone functions, which are mirrored by the isolated NC protein after physiological cleavage from Gag. Gag, however, is suggested to have superior nucleic acid chaperone activity. Here we investigate the interaction of Gag and NC with the core RNA structure of the HIV-1 packaging signal (Ψ), using 2-aminopurine substitution to create a series of modified RNAs based on the Ψ helix loop structure. The effects of 2-aminopurine substitution on the physical and structural properties of the viral Ψ were characterized. The fluorescence properties of the 2-aminopurine substitutions showed features consistent with the native GNAR tetraloop. Dissociation constants (K_d) of the two viral proteins, measured by fluorescence polarization (FP), were similar, and both NC and Gag affected the 2-aminopurine fluorescence of bases close to the loop binding region in a similar fashion. However, the influence of Gag on the fluorescence of the 2-aminopurine nucleotides at the base of the helix implied a much more potent helix destabilizing action on the RNA stem loop (SL) versus that seen with NC. This was further supported when the viral Ψ SL was tagged with a 5' fluorophore and 3' quencher. In the absence of any viral protein, minimal fluorescence was detected; addition of NC yielded a slight increase in fluorescence, while addition of the Gag protein yielded a large change in fluorescence, further suggesting that, compared to NC, the Gag protein has a greater propensity to affect RNA structure and that Ψ helix unwinding may be an intrinsic step in RNA encapsidation.



The major structural protein of retroviruses, including HIV-1, is the Gag polyprotein, which can form virus-like particles when expressed alone in cells.¹ Gag has four major domains into which it is cleaved during viral maturation, a process associated with viral budding from the cell.^{2,3} In HIV, one of these four domains has two zinc knuckle motifs and on cleavage forms the nucleocapsid protein (NC) that coats the RNA genome within the mature virion.^{2,4} The RNA chaperone functions of NC have been extensively studied;^{5–9} however, the ability of NC to fully unwind nucleic acid helices is not certain. Early circular dichroism (CD) and nuclear magnetic resonance (NMR) spectroscopic evidence for binding of NC to tRNA suggested that NC alters base stacking interactions, but it was unclear whether unwinding of the nucleic acid helices was occurring.^{10–12} Musier-Forsyth et al. used several techniques to monitor the binding of NC to tRNA, and although helix unwinding of the tRNA was not observed, there was significant unstacking of the bases within the tRNA acceptor–T Ψ C minihelix.^{9,13} Further work on the secondary structure of the cTAR^{14–17} and the primer binding site (PBS) stem loop (SL)¹⁸

again demonstrated the ability of NC to interfere with base stacking but without completely opening the helix. Known RNA annealing activities of NC include promoting dimerization of the RNA genome and positioning of the tRNA primer on the PBS. However, these processes occur in immature virions prior to the release of NC from the precursor Gag polyprotein,^{2,19} suggesting that it is the uncleaved Gag that may be the effector molecule. Similarly, the highly specific capture of the viral genomic RNA from the vast excess of similar cellular RNAs must occur before budding and processing of the polyprotein, again with Gag necessarily being the active moiety involved. Gag has been less studied as an RNA chaperone than NC, although the NC domain of the Gag polyprotein is in part similar in structure to the free NC protein.^{20,21}

Gag has also been shown to have superior RNA chaperoning ability compared to that of NC in strand transfer assays during

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Aliquots of FAM WT, AP10, AP4, AP3, AP13, or AP14 RNA and NC or GagΔP6 protein were allowed to defrost on ice prior to being used. All RNA oligonucleotides were heated at 95 °C for 5 min and snap-cooled on ice for a further 5 min, before the addition of buffer to give an RNA concentration of 1 μM [25 mM NaOAc (pH 6.5), 2 mM MgCl₂, and 0.1 mM ZnCl₂]. Five microliters of the RNA substrate was added to the wells of a 384-well microplate (low-volume flat bottom black NBS treated, Corning 3820) before 5 μL of NC or GagΔP6 protein was added to the wells containing the RNA at concentrations between 0 and 20 μM. The microplate was incubated for 45 min before measurements were taken on a Pherastar⁺ plate reader (BMG LabTech) at 25 °C. For the steady-state measurement, a 320 nm excitation filter (to minimize protein fluorescence) and a 380 nm emission filter were used. For FP measurements, a 485 nm excitation filter and a 520 nm emission filter were used. Each measurement is the mean of three independent experiments, and errors are one standard deviation (SD). Experimental titration data were modeled using a non-linear least-squares fit of a single-site, two-state binding model equation (eq 1).³¹ Dissociation constants (K_d) and apparent dissociation constants (K_d') were estimated from the total change in polarization or the total change in 2-aminopurine fluorescence intensity, $(I - I_0)/(I_f - I_0)$, plotted as a function of the total NC or GagΔP6 concentration.

$$y = \frac{[R_{\text{tot}} + P_{\text{tot}} + K_d - \sqrt{(R_{\text{tot}} + P_{\text{tot}} + K_d)^2 - 4(R_{\text{tot}}P_{\text{tot}})}]}{2R_{\text{tot}}} \quad (1)$$

where I is the measured fluorescence intensity/polarization, I_0 is the initial fluorescence intensity/polarization of the RNA substrate in the absence of protein, I_f is the fluorescence intensity/polarization of the RNA substrate when saturated with protein, R_{tot} is the total RNA concentration, P_{tot} is the total protein concentration, and K_d is the dissociation constant (or K_d' as the apparent dissociation constant).

Because of the high NC concentrations required to saturate the system when using AP3, AP13, and AP14, the assumption was made that I_f during NC titration would be the same as I_f during GagΔP6 titration. This assumption was based on the very similar experimental I_f gained for both AP10 and AP4 SLs, from saturation with NC and GagΔP6 titrations (Figures 6–10 of the Supporting Information).

Fluorescence Intensities of the Molecular Beacons FQ WT and FQ Control with Titrations of NC and GagΔP6 Protein. These experiments were essentially conducted as described above. Five microliters of the snap-cooled molecular beacon FQ WT or FQ Control at a concentration of 0.2 μM was added to the wells of a 384-well plate followed by addition of 5 μL of either the protein buffer, NC, or GagΔP6 (0–20 μM). The microplate was incubated for 45 min before measurements were taken at 25 °C using a Pherastar⁺ plate reader (BMG LabTech) with a 510 nm excitation filter and a 540 nm emission filter. The FQ WT and FQ Control RNA in their native and chemically denatured (9 M urea) states were corrected for screening and buffer effects and normalized as 0 and 100% intensity, respectively; the signal gained from the unfolding of FQ WT and FQ Control by GagΔP6 or NC is described in comparison to these native and denatured states. Each measurement is the mean of three independent experiments, and

errors are one SD and the data modeled with eq 1 as described above.

RESULTS

Design and Structural Validation of 2-Aminopurine-Modified RNA Oligonucleotides AP10, AP4, AP3, AP13, and AP14. The 14-nucleotide RNA SL used in this study is the native SL3 found in the Ψ-packaging domain of HIV-1. SL3 consists of a 5 bp stem with a four-base GGAG loop and as such is part of the GNAR family of tetraloops.⁴ To study the interactions of both NC and GagΔP6 proteins on the overall structure of SL3, we systematically incorporated a 2-aminopurine-modified nucleotide at each of the 5 bp comprising the stem of SL3 (Figure 1). This gave a series of five 2-aminopurine-modified RNAs, which are capable of giving single-nucleotide resolution of the binding of NC and GagΔP6. Because of the differences in hydrogen bonding and sterics of the 2-AP:C and 2-AP:U base pairs compared to those of the native G:C and A:U base pairs, it was important to biophysically characterize the 2-aminopurine-modified RNA substrates compared to the WT RNA (Figures 2–5 of the Supporting Information).

The melting temperature (T_m) of WT (67.9 ± 0.5 °C) SL3 RNA indicates the formation of a highly thermally stable SL (Figure 2 of the Supporting Information). Furthermore, a magnesium titration of WT SL3, measured by CD (Figure 4 of the Supporting Information), indicates that very little structural change is observed over the range of magnesium concentrations measured (0–10 μM). The high thermal and structural stability of SL3 fits well with previous studies of RNA tetraloops.³²

2-AP modification to nucleotides at positions 10 and 4 created the modified SLs AP10 ($T_m = 53.5 \pm 0.8$ °C) and AP4 ($T_m = 43.2 \pm 0.4$ °C). The large differences in the melting temperatures (ΔT_m) of AP10 ($\Delta T_m = 14.4$ °C) and AP4 ($\Delta T_m = 24.7$ °C) highlight the importance of the closing G-C base pair in the thermal stability of RNA tetraloops.^{33–36} Modifications at positions 3, 13, and 14 gave SLs AP3 ($T_m = 63.2 \pm 0.4$ °C), AP13 ($T_m = 64.3 \pm 0.8$ °C), and AP14 ($T_m = 62.1 \pm 0.8$ °C). The relatively small differences in ΔT_m values for SLs AP3 ($\Delta T_m = 4.7$ °C), AP13 ($\Delta T_m = 3.6$ °C), and AP14 ($\Delta T_m = 5.8$ °C) indicate that modifications toward the lower part of the SL yield little change in their overall thermal stability. The thermal denaturation data show that at 25 °C the majority of the RNA population is in the folded SL structure, which applies to all of the 2-aminopurine-modified SLs. This is further supported by the CD spectra obtained at 25 °C, where all the 2-aminopurine-modified SLs suggest a structure closely related to that of the WT RNA (Figure 4 of the Supporting Information).

Binding of GagΔP6 and NC to Wild-Type SL3 Measured by Fluorescence Polarization. Each 2-aminopurine-modified RNA SL can provide detailed insight into the effects that binding of GagΔP6 and NC has on the local structure of SL3, while together the 2-aminopurine series can illustrate how the binding of the viral proteins affects the global structure of SL3. The 2-aminopurine modifications provide only apparent dissociation constants (K_d') relating to each 2-aminopurine nucleotide. Dissociation constants for GagΔP6 and NC with SL3 as a whole were determined by measurement of the changes in the fluorescence polarization (FP) as protein was titrated against a 5'-FAM-labeled SL3 (FAM WT).

The determination of the K_d values for both GagΔP6 (0.086 ± 0.002 μM) and NC (0.117 ± 0.003 μM) was important

Table 1. Relative Melting Temperatures, Quantum Yields, and Dissociation Constants of the 2-AP RNA Substrates Used in This Study

quantum yield										apparent dissociation constant (K_d) ^h				
UV thermal denaturation				folded ^d		unfolded ^e			$\Delta\Phi^f$	NC		GagΔP6		
substrate	T_m (°C) ^a	1 SD	ΔT_m (°C) ^b	substrate	Φ^c	1 SD	Φ	1 SD		substrate	K_d (μM)	1 SD	K_d (μM)	1 SD ^g
WT	67.9	0.4	—	2-AP	0.680	0.680	0.690	0.002	—	Fam-WT	0.117 ⁱ	0.003	0.086 ⁱ	0.002
AP10	53.5	0.5	14.4	AP10	0.190	0.006	0.117	0.007	0.073	AP10	0.10	0.04	0.21	0.03
AP4	43.2	0.4	24.7	AP4	0.008	0.0004	0.080	0.003	0.072	AP4	0.23	0.04	0.11	0.04
AP3	63.2	0.4	4.7	AP3	0.002	0.0002	0.020	0.001	0.018	AP3	1.4	0.2	0.25	0.05
AP13	64.2	0.8	3.7	AP13	0.004	0.0002	0.038	0.001	0.034	AP13	2.3	0.2	0.16	0.04
AP14	62.1	0.9	5.8	AP14	0.039	0.004	0.187	0.002	0.148	AP14	5.0	0.2	0.13	0.08

^aMelting temperatures were determined in 25 mM NaOAc (pH 6.5), 25 mM NaCl, and 0.1 mM MgCl₂ at an RNA substrate concentration of 5 μM. ^bDifference in melting temperature compared to that of wild-type SL3. ^cRelative quantum yields were determined using 2-aminopurine ($\Phi = 0.68$) under folded and unfolded conditions at an RNA substrate concentration of 0.5 μM. ^dFolded conditions included 25 mM NaOAc (pH 6.5), 25 mM NaCl, and 0.1 mM MgCl₂. ^eUnfolded conditions included 25 mM NaOAc (pH 6.5), 25 mM NaCl, and 9 M urea. ^fDifference in quantum yield under folded and unfolded conditions. ^gEach measurement is the mean of three independent experiments, and errors are one SD. ^hApparent dissociation constant (K_d) measured by 2-aminopurine intensities. ⁱDissociation constant (K_d) measured by FP.

(Table 1 and Figure 2A) as it rules out any differences seen from the K_d values determined when using the 2-aminopurine-modified RNA substrates, being an effect from the difference in the binding affinities of the Gag and NC proteins.

Binding of GagΔP6 and NC to the Series of 2-Aminopurine-Modified RNA Stem Loops. The K_d value for each 2-aminopurine-modified nucleotide can be combined to determine whether the effects of GagΔP6 and NC binding encountered at the top of the stem are translated equally through the 5 bp to the bottom of the stem and therefore whether the two proteins affect the SL in a similar fashion at comparable concentrations.

If one starts with AP10 and AP4 where the 2-aminopurine modifications are positioned at the top of the stem, the K_d values for NC (for AP10, $K_d = 0.10 \pm 0.04$ μM, and for AP4, $K_d = 0.23 \pm 0.04$ μM) and GagΔP6 (for AP10, $K_d = 0.21 \pm 0.02$ μM, and for AP4, $K_d = 0.11 \pm 0.04$ μM) are not only similar to each other but also similar to the K_d values determined by FP for the binding of NC ($K_d = 0.117 \pm 0.003$ μM) and GagΔP6 ($K_d = 0.086 \pm 0.002$ μM) to FAM WT. Moving down the stem to the third base pair below the GGAG tetraloop reveals a difference between the K_d values for NC ($K_d = 1.4 \pm 0.2$ μM) and GagΔP6 ($K_d = 0.25 \pm 0.04$ μM) for the AP3 RNA substrate. This difference in K_d values between NC and GagΔP6 continues to increase at the fourth and fifth base pair below the GGAG tetraloop, with RNA substrates AP13 (for NC, $K_d = 2.3 \pm 0.2$ μM, and for GagΔP6, $K_d = 0.18 \pm 0.04$ μM) and AP14

(for NC, $K_d = 5.0 \pm 0.2$ μM, and for GagΔP6, $K_d = 0.13 \pm 0.6$ μM), respectively (Table 1 and Figures 6–10 of the Supporting Information).

Binding of GagΔP6 and NC to the Molecular Beacons FQ WT and FQ Control. To further describe the unwinding ability of GagΔP6 compared to that of NC, an assay involving an RNA molecular beacon (FQ WT) was established. FQ WT ($T_m = 72.2 \pm 0.6$ °C) was titrated against either GagΔP6 or NC, in the same manner described for the 2-aminopurine-modified RNA substrates. Addition of NC to FQ WT ($K_d = 0.33 \pm 0.06$ μM) results in an increase in the magnitude of the signal of ~15% at 10 μM, whereas the unwinding of FQ WT ($K_d = 0.15 \pm 0.02$ μM) by GagΔP6 leads to an increase in the magnitude of the signal of ~60% at 10 μM, when compared to that of chemically denatured FQ WT (Figure 3B).

The ability of NC to push a structured nucleic acid population toward a more unfolded or folded state has been described previously, and the slight increase in the fluorescence from the FQ WT RNA is most probably in response to this shift in the population.¹⁶ Addition of the GagΔP6 protein results in a significant change in the fluorescence intensity and further supports the idea that GagΔP6 has better SL destabilizing and unwinding ability than NC.

DISCUSSION

The biophysical characterization of the WT RNA shows that the GGAG tetraloop forms a highly stable hairpin structure.

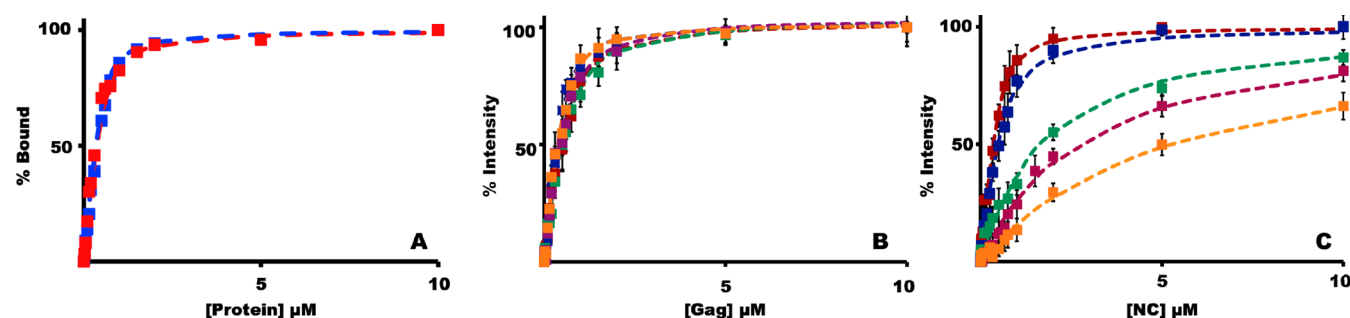


Figure 2. Titration curves of Fam WT and 2-aminopurine-modified RNA SLs AP10 (dark red, top), AP4 (dark blue), AP3 (green), AP13 (dark red, bottom), and AP14 (orange) with GagΔP6 and NC. The changes in (A) fluorescence polarization [GagΔP6 (blue) and NC (red)] or (B and C) 2-aminopurine fluorescence intensity, $(I - I_0)/(I_t - I_0)$, were plotted as a function of the total NC or GagΔP6 concentration with an RNA substrate concentration of 0.5 μM. The dashed lines in panels A–C of Figure 3 correspond to the fits of the experimental points with eq 1.

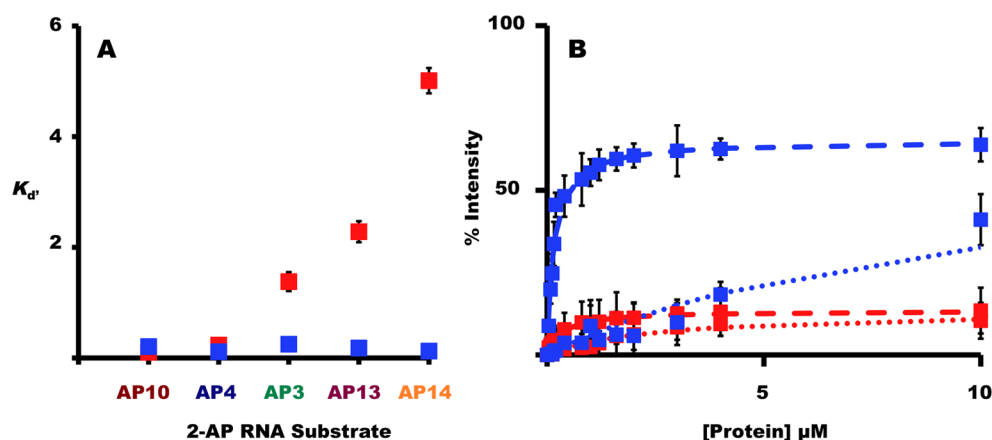


Figure 3. (A) Plot of apparent dissociation constant (K_d) for GagΔP6 (blue) and NC (red) vs 2-aminopurine positional modification within the RNA substrates. Starting with AP10 as the closing base pair to AP14 as the terminal base pair. (B) Plot of the change in fluorescence intensity of FQ WT (---) and FQ Control (···) upon addition of GagΔP6 (blue) or NC (red) with an RNA substrate concentration of 0.1 μ M. Fluorescence intensities, $(I - I_0)/(I_f - I_0)$, were plotted as a function of the total NC or GagΔP6 concentration. The dashed lines in panel B correspond to the fits of the experimental points to eq 1. Because of the high protein concentration that would be required to saturate the FQ control RNA, the dotted lines are illustrative, not actual fits of the experimental data.

The high thermal stability of SL3 has been reported previously by Borer et al. ($T_m < 70$ °C)³⁷ and Scholes et al. ($T_m = 86.4$ °C with Mg^{2+} , and $T_m = 76.8$ °C without Mg^{2+}),³⁸ with both groups using a 3 bp extended SL3.

The characterization of the modified RNAs demonstrated that a 2-aminopurine inserted at or just below the closing C-G base pairs has a stronger effect on the thermal stability of the SL than modifications made further down the stem. The ΔT_m values for the series of 2-aminopurine-modified SLs ranged from 3.6 to 24.7 °C, when compared to that of WT SL3 (Figure 2 of the Supporting Information). All the 2-aminopurine RNAs showed T_m values of >40 °C. Furthermore, CD spectra showed an A-form RNA structure for all the 2-aminopurine-modified SLs at 25 °C (Figure 3 of the Supporting Information), indicating that the 2-aminopurine series exhibit a SL structure that closely resembles the structure of WT SL3 (Figure 3 of the Supporting Information).

The quantum yield determinations for the 2-aminopurine RNA substrates in folded and unfolded states again support a SL structure (Figure 5 of the Supporting Information). AP10 like AP3, AP4, and AP13 has two neighboring nucleotides, yet AP10 shows a very strong emission spectrum compared to those of AP3, AP4, and AP13. The structure of the GGAG tetraloop, determined by NMR spectroscopic studies,⁴ shows that the G nucleobase at position 9 of the tetraloop is oriented away from the G nucleobase at position 10. Therefore, the limited base stacking of the G nucleobases at position 10 results in stronger emission spectra for AP10 than for any of the other 2-aminopurine-modified SLs.

The binding of NC to SL3 has been extensively investigated using a variety of biophysical techniques,^{37–45} whereas binding of Gag to SL3 has received far less attention. There are several published studies, using fluorescence anisotropy (FA), that compare the binding of NC and Gag to different RNAs. Musier-Forsyth et al. used a 37-nucleotide RNA minihelix^{lys3} (for NC, $K_d = 0.092 \pm 0.009$ μ M, and for GagΔP6, $K_d = 0.07 \pm 0.02$ μ M);²³ Williams et al. used a 10-nucleotide (UG)₅ repeat (for NC, $K_d = 0.012 \pm 0.002$ μ M, and for GagΔP6, $K_d = 0.019 \pm 0.008$ μ M),⁶ and Wu et al. used a 20-nucleotide (AGC UGC UUU UUG CCU GUA CU) sequence from the extreme 3' terminus of U3 (for NC, $K_d = 0.10 \pm 0.02$ μ M, and for GagΔP6,

$K_d = 0.01 \pm 0.02$ μ M).⁵ The differences found in the binding of Gag and NC to NA have previously been discussed in the literature.⁷

As NC, and therefore the NC domain within GagΔP6, binds to the GGAG tetraloop at the top of the stem,^{4,44} the similarity between the K_d values determined for AP10 and AP4 and the K_d values determined by FP above (with NC and GagΔP6) are consistent with the assumption that the local environments of the 2-aminopurine nucleotide, in AP10 and AP4, are similar when bound to the free NC or to the NC domain within the GagΔP6 protein.

Further down the stem RNA substrates AP3, AP13, and AP14, the similarities among the K_d values determined for GagΔP6 and NC start to diverge with the distance of the base pair from the GGAG tetraloop (Figure 3A). The K_d values for GagΔP6 with AP3, AP13, and AP14 all remain comparable with each other as well as with the K_d values determined for AP10 and AP4.

Given that the NC footprint is approximately six to eight nucleotides and assuming a 1:1 RNA:NC stoichiometry,⁴⁴ the nucleotides at the bottom of the stem (AP3, AP13, and AP14) are less involved in a direct interaction with the free NC. The intensity change to the AP3, AP13, and AP14 nucleotides may be more of a response to the destabilization of the base pairs caused by the accommodation of NC on the top of the loop or due to a second, nonspecific binding of NC further down the SL.

During the titrations of NC and GagΔP6 with FQ WT, the data from both titrations show little change in signal intensity at higher protein concentrations (Figure 3B). An expectation of a second nonspecific binding of NC to FQ WT might be the increased level of destabilization of the stem loop and therefore an increase in the magnitude of the signal measured as seen with the 2-aminopurine substrates mentioned above. The fact that with the FQ WT substrate there is very little signal change at higher NC concentrations suggests that a second NC binding is not occurring, but also the possibility that further opening of the stem loop does not occur during the second NC binding or that the fluorophore–quencher system interferes with this nonspecific binding.

The destabilization of nucleic acid structure by NC has been demonstrated previously by time-resolved fluorescence studies on a molecular beacon based on the DNA TAR SL; the population of the DNA TAR SL moved toward a more unfolded or folded state upon addition of NC.¹⁴ To ensure the change in signal observed for the NC is not due to an artifact of the system used in this study, a second molecular beacon, **FQ Control** ($T_m = 75.1 \pm 0.3$ °C), was employed in which the stem was kept identical to **FQ WT** but the loop was changed from a GGAG to a GAAA tetraloop. Previous studies determined that the SL3 GAAA mutation removed the high-affinity binding of NC to the tetraloop.⁴⁶ The titration of **FQ Control** with NC showed that nonspecific binding of the protein to the SL is possible (Figure 3B). We would argue that because of the size of the RNA chosen in this study, it is improbable that two NC proteins could fit onto the 14-nucleotide stem.

However, it is apparent that the initial, specific, and high-affinity interaction between NC and SL3 has little effect on the 2-aminopurine intensities (**AP3**, **AP13**, and **AP14**) toward the bottom of the stem, which is in agreement with the NMR solution structures of NC bound to an extended SL3. The NMR structure shows a single binding of NC and shows that this single NC is not enough to unwind the base pairs at the bottom of the stem.⁴⁴

The footprint of GagΔP6 has not been established but is unlikely to be smaller than that of NC,⁷ meaning the nucleotides at the bottom of the stem (for **AP3**, **AP13**, and **AP14**) would be more directly involved in the binding of GagΔP6. Our results support this, as all five 2-aminopurine-modified SLs as well as the molecular beacon **FQ WT** show consistent K_d values with GagΔP6. Again, the assumption is for a 1:1 binding of GagΔP6 to SL3, but there is the possibility that Gag exists in a multimeric state before the introduction of the RNA substrate or the RNA substrate acts as a template for the formation of higher-order structure. Rein et al. showed that GagΔP6 does dimerize in solution with a K_d of 5.5 μM,⁴⁷ and that aggregation of GagΔP6 is achieved by a nucleic acid template.⁴⁸

The binding of more than one GagΔP6 on or near the RNA substrate would create a higher local concentration of the NC domain around the RNA substrate. To estimate stoichiometries of binding of GagΔP6 to the 2-AP RNA substrates, the data were modeled according to the method by Doudna et al.⁴⁹ Models for 1:1 and 1:2 RNA:GagΔP6 stoichiometries were also plotted as a comparison and illustrate that the data gained from the titration of GagΔP6 into the 2-AP substrates are best described by a 1:1 stoichiometry (Figures 6–10 of the Supporting Information). Furthermore, all the K_d values reported by the 2-aminopurine intensities are at least 20-fold below the K_d for Gag dimerization (Table 1).⁴⁷

Gag has been shown to have a NA binding region in the matrix domain,²³ again the short SL used in our experiments is unlikely to be able to bind both NC and matrix regions on Gag simultaneously, and the dominant effect of NC means that effectively our system is studying NC domain binding alone. Further work to include larger RNAs and Gag mutants to further clarify the points made above and to provide further information about a possible mechanism for the unwinding of nucleic acid structure by Gag is ongoing.

For these reasons and because of the evidence of the molecular beacon experiments, we believe that the results presented here indicate that a much greater destabilizing effect is introduced through the stem to accommodate the binding of a single GagΔP6 to the top of the tetraloop, compared to NC.

CONCLUSION

The implication of a greater destabilizing effect by GagΔP6 on SL3 is that GagΔP6 shifts the population further toward a more unfolded state, suggesting that GagΔP6 is more able to alter NA structure than NC. The data are consistent with a model in which the Gag protein has the ability not only to bind to ssRNA, which is a function well established for the NC protein, but also to unwind the RNA helices to which it binds during the process of genome encapsidation. RNA structure in HIV is highly likely to be dynamic, with different regions adopting different structures during trafficking through the cell to accommodate the functional requirements of the RNA at different times (transactivation, nuclear export, translation and packaging, etc.). The viral genome is believed to be captured in the cytoplasm and transported to the plasma membrane chaperoned by relatively few Gag molecules² and at this stage as at many others is likely also complexed with cellular RNA binding proteins, many of which are still being identified. Structural switches also seem to be critical for this process, with accumulating evidence indicating that the viral genomic RNA can undergo quite striking changes in conformation to fulfill its function and these are undoubtedly triggered by RNA–protein interactions. The specific ability of Gag, rather than NC, to effect structural change after binding SL3 also prevents any prematurely cleaved NC within the cell from deflecting the RNA from its pathway to be packaged, as it will not trigger the same changes with their functional consequences.

ASSOCIATED CONTENT

Supporting Information

Additional materials and methods and further results and discussion of UV thermal melting, circular dichroism, quantum yield, and stoichiometry determination as well as individual plots for the titration of all 2-aminopurine-modified RNA substrates with NC and GagΔP6. This material is available free of charge via the Internet at <http://pubs.acs.org>.

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Notes

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ABBREVIATIONS

2-AP, 2-aminopurine; $\Delta\Phi$, difference in quantum yield; ΔT_m , difference in melting temperature; Φ , quantum yield; A, adenine; BHQ, black hole quencher; C, cytidine; CD, circular dichroism; FAM, 5(6)-carboxyfluorescein; G, guanine; HIV-1, human immunodeficiency virus type 1; I , measured fluorescence intensity/polarization; I_f , final fluorescence intensity/polarization; I_0 , initial fluorescence intensity/polarization; K_d , dissociation constant; K_d , apparent dissociation constant; Tet, tetrachlorofluorescein;

T_m , melting temperature; NC, nucleocapsid; PBS, primer binding site; P_{tot} , total protein concentration; R_{tot} , total RNA concentration; U, uridine; UTR, untranslated region; UV, ultraviolet.

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