Site-selective probing of cTAR destabilization highlights the necessary plasticity of the HIV-1 nucleocapsid protein to chaperone the first strand transfer

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Received January 19, 2013; Revised February 18, 2013; Accepted February 20, 2013

ABSTRACT

The HIV-1 nucleocapsid protein (NCp7) is a nucleic acid chaperone required during reverse transcription. During the first strand transfer, NCp7 is thought to destabilize cTAR, the (–)DNA copy of the TAR RNA hairpin, and subsequently direct the TAR/cTAR annealing through the zipping of their destabilized stem ends. To further characterize the destabilizing activity of NCp7, we locally probe the structure and dynamics of cTAR by steady-state and time resolved fluorescence spectroscopy. NC(11–55), a truncated NCp7 version corresponding to its zinc-finger domain, was found to bind all over the sequence and to preferentially destabilize the penultimate double-stranded segment in the lower part of the cTAR stem. This destabilization is achieved through zinc-finger–dependent binding of NC to the G10 and G50 residues. Sequence comparison further revealed that C/C15A mismatches close to the two G residues were critical for fine tuning the stability of the lower part of the cTAR stem and conferring to G10 and G50 the appropriate mobility and accessibility for specific recognition by NC. Our data also highlight the necessary plasticity of NCp7 to adapt to the sequence and structure variability of cTAR to chaperone its annealing with TAR through a specific pathway.

INTRODUCTION

The nucleocapsid protein (NCp7) of the human immunodeficiency virus type 1 (HIV-1) is a small basic protein resulting from the protease-mediated cleavage of the Gag polyprotein precursor. NCp7 is formed by two highly conserved CCHC zinc-finger motifs connected by a short flexible linker and flanked by poorly folded N- and C-terminal basic domains (1,2) (Figure 1). NCp7 is critical during the early and late steps of the HIV-1 viral life cycle (3–11). Most NCp7 functions rely on its RNA- and DNA-chaperone properties that direct the rearrangement of nucleic acids into their most thermodynamically stable structure (12–15). These chaperone properties are thought to play a critical role in reverse transcription (5,16,17). For instance, during the first strand-transfer of reverse transcription, NCp7 favours the transfer of the newly synthesized minus strong-stop DNA [(–) ssDNA] to the 3’-end of the viral RNA genome, by promoting the base pairing of the complementary repeat (R) sequences at the 3’-ends of the (–)ssDNA and genomic RNA, respectively (18). In vitro, the stable stem–loop (SL) of the transactivation response element (TAR) sequence within the R domain of the genomic RNA does not readily anneal with the complementary cTAR sequence of the (–)ssDNA in the absence of NCp7 (18–21). Addition of NCp7 strongly increases the rate and extent of the cTAR sequence of the (–)ssDNA and genomic RNA, respectively (18). In vitro, the stable stem–loop (SL) of the transactivation response element (TAR) sequence within the R domain of the genomic RNA does not readily anneal with the complementary cTAR sequence of the (–)ssDNA in the absence of NCp7 (18–21). Addition of NCp7 strongly increases the rate and extent of the cTAR/TAR annealing reaction (18–23) and switches the reaction pathway to direct the hybridization of these sequences through the end of their double-stranded (ds) stems (20,21). The main chaperone properties involved in promoting the first strand transfer have been characterized as two interconnected components: a destabilization activity and a hybridization-promoting activity. Promotion of the annealing of complementary sequences requires an efficient aggregation of nucleic acids (24). This is achieved by the numerous basic residues of NCp7 that decrease the electrostatic repulsion between complementary oligonucleotides by neutralizing the negatively
single-stranded sequences (44,45), as well as in the loop of the primer-binding site sequence of HIV-1 (46). These studies showed that binding of the intact ZF domain was responsible for a decrease of the DNA base stacking and for a strong restriction of the base mobility on a picosecond–nanosecond timescale. By selectively replacing different adenine positions of the cTAR sequence by 2-Ap (Figure 1), we aimed to picture the preferential destabilization sites and to gain insight into the molecular mechanism by which the NCp7 ZFs destabilize the cTAR stem–loop. Although NC(11–55) was found to bind all over the sequence, it preferentially restricted the dynamics and destabilized the bases of the penultimate ds-segment sandwiched between two C•A mismatches in the lower part of the cTAR stem. These NC(11–55)-induced changes are highly specific, as they were not observed when NC(11–55) was substituted by the SSHS–NC(11–55) mutant where the Cys residues in the ZFs were substituted by Ser residues to prevent the binding of zinc and the folding of the ZFs. These structural changes in this part of cTAR play likely a critical role in the ability of NCp7 to promote cTAR/TAR annealing through a specific zipping pathway.

Figure 1. The 2-Ap substituted cTAR (A) and NC(11–55) (B) sequences. The cTAR sequence from the Lai strain of HIV-1 was selectively modified either at position 9, 17, 21, 28, 35, 45, 49, 53 or 55 with 2-Ap.

MATERIALS AND METHODS

NC(11–55) (Figure 1) was synthesized as previously described (47) and stored lyophilized in its zinc-bound form. An extinction coefficient of 5.700 M⁻¹cm⁻¹ at 280 nm was used to determine its concentration. The oligonucleotides selectively labelled at different positions with 2’-deoxyriboyl-2-aminopurine (2-Ap) (Figure 1) were synthesized at a 1 μmol scale by IBA GmbH Nucleic Acids Product Supply (Göttingen, Germany) and purified by the manufacturer by double reverse-phase high-performance liquid chromatography.

Steady-state fluorescence

Experiments were performed in 25 mM Tris–HCl, pH 7.5, 30 mM NaCl and 0.2 mM MgCl₂ at 20°C. Excitation and emission spectra were recorded with a Fluorolog or FluoroMax spectrophluorometer (Jobin Yvon Instruments, S.A. Inc.). All fluorescence intensities were corrected for buffer emission, lamp fluctuations and instrumental wavelength-dependent bias. Kinetics of cTAR/TAR annealing were monitored in real-time using fluorescent doubly labelled cTAR sequences and non-labelled TAR (20). The cTAR sequences were doubly labelled with 5/6 fluorescein (Fl) and 5/6 tetramethylrhodamine (TMR), at their 3’- and 5’-ends, respectively. Excitation and emission wavelengths were at 480 and 520 nm, respectively, to monitor the Fl fluorescence restoration resulting from the formation of the cTAR/TAR duplex. Concentrations of 5’TMR-cTAR-3’Fl and TAR were 10 and 100–600 nM, respectively, to ensure pseudo-first order conditions. Both reactants in identical volumes coated by NC peptides were mixed together to trigger the reaction. The apparent rate constants kobs were determined from kinetic traces, as previously described (20). All fitting procedures were carried out with the Microcal Origin 6.1

charged phosphate groups (5.21,25). The annealing facilitator property of NCp7 is shared with many other basic proteins that are also able to promote the cTAR/TAR DNA hybridization (26–28). On the contrary, the ability to destabilize the cTAR sequence appears to be a hallmark of NCp7. NCp7 activates the transient opening (fraying) of the lower half of the cTAR stem (29–34) constituted by a series of short ds-segments separated by a bulge and two mismatches. This activation follows a non–two-state mechanism, which results in complex spatial and temporal fluctuations of the cTAR ends that greatly facilitate cTAR/TAR hybridization. This activity has been assigned to the conserved hydrophobic platform formed by the Val₁₃, Phe₁₆, Thr₂₄, Ala₂₅, Trp₃₇, Gln₄₅ and Met₄₆ residues at the surface of the folded zinc-finger (ZF) domains (1,2,35). More specifically, the Trp₃₇ residue plays a critical role through stacking with G residues (17,36–43). This hydrophobic platform has also been found critical for binding (35,38). Nevertheless, the precise molecular mechanisms by which NCp7 destabilizes cTAR remain poorly described. Here, we selectively modified the 57-nt cTAR stem–loop sequence of the Lai strain with 2-aminopurine (2-Ap) at different positions to investigate by fluorescence spectroscopy the structural and dynamic modifications that occur within cTAR in response to NCp7 binding. In contrast to most techniques that monitor only global structural changes, the exquisite sensitivity of 2-Ap fluorescence to its local environment allows region-specific characterization of the structural modifications within the cTAR hairpin sequence. The 2-Ap has been previously used to site-specifically characterize the NCp7-induced changes in the dynamics of short
software based on non-linear least-squares methods, applying the Levenberg–Marquardt algorithm.

**Time-resolved fluorescence spectroscopy**

Time-resolved fluorescence measurements were performed using the time-correlated single-photon counting technique described elsewhere (46). Briefly, excitation pulses at 315 nm were generated by a pulse-picked frequency-tripled titanium:sapphire laser (Tsunami, Spectra Physics) pumped by a diode-pumped solid-state Nd: YVO₄ doubled laser ( Millennia X, Spectra Physics). The emission was collected at 370 nm through a polarizer set at the magic angle and a 16-nm band-pass monochromator (H10, Jobin-Yvon). The single-photon events were detected with a micro-channel plate (R3809U, Hamamatsu) photomultiplier coupled to a pulse pre-amplifier (HFAC, Becker and Hickl GmbH) and recorded on a time correlated single photon counting board (SPC-130, Becker and Hickl GmbH). The full-width at half maximum of the instrumental response function was ~50 ps. For time-resolved anisotropy measurements, the fluorescence decay curves were recorded at vertical and horizontal positions as detailed in the Supplementary Materials. Experimentally measured fluorescence decays were de-convoluted with the instrumental response function and fitted to retrieve the most probable lifetime distribution using the maximum entropy method (48,49). As the resulting lifetime distribution evidenced four well-individualized lifetime clusters, see Supplementary Figure S3 the decays were reported as the sum of discrete exponentials according to \( I(t) = \sum a_i \exp(-t/\tau_i) \), where \( a_i \) is the relative amplitude associated to the fluorescence lifetime \( \tau_i \) representing a distribution of conformations in which 2-Ap experiences similar quenching rates. The mean lifetime \( \langle \tau \rangle \) was calculated according to \( \langle \tau \rangle = \sum a_i \tau_i \). The population \( a_0 \) of dark species was calculated by \( a_0 = 1 - \frac{\tau_{\text{free}}}{\tau_{\text{sample}} - \tau_m} \) where \( \tau_{\text{free}} \) is the lifetime of the free 2-Ap riboside, \( \tau_{\text{sample}} \) is the measured lifetime of 2-Ap within the oligonucleotide (either free or bound to NCp7), and \( \tau_m \) is the ratio of their corresponding quantum yields. The remaining amplitudes of each fluorescent population were recalculated according to \( a_{\text{res}} = a_i (1 - a_0) \). All fitting procedures were carried out both with the maximum entropy method Pulse5 software and a software based on an iterative application of the Marquardt’s method in non-linear least-squares analysis (kindly provided by G. Krishnamoorthy).

**Alignment of TAR sequences**

TAR sequence alignments were performed using the HIV sequence database from the Los Alamos National Lab HIV (http://www.hiv.lanl.gov/content/sequence/HIV/mainpage.html). A position-weighted matrix was then computed using the open-source statistical software R from the Comprehensive R Archive Network (http://www.R-project.org/) (50). The sequence logo (51) was plotted using the seqLogo package (http://www.bioconductor.org/packages/devel/bioc/html/seqLogo.html).

**RESULTS**

Site-specific characterization of the 2Ap-labelled cTAR sequences in the absence of NCp7

cTAR is an imperfect stem–loop characterized by a 3-nt internal loop, as well as by several bulged nucleotides and mismatches (Figure 1A). We selected labelling positions all along the cTAR Lai sequence. The 2-Ap in position 9 forms a mismatch with C within the lower part of the stem. Positions 28 and 35 are within the central and the internal loop, respectively, whereas positions 53 and 55 are flanking the G₃₄ bulge, and positions 17, 21, 45 and 49 are distributed all over the stem within the different ds-segments. The fluorescence spectra of the different 2-Ap-labelled cTAR sequences exhibited the same maximum emission wavelength (at ~370 nm) as the free 2-Ap nucleotide. In contrast, their quantum yields (QYs) were strongly reduced in comparison with the free probe (Figure 2), in line with static and dynamic quenching of 2-Ap by its close neighbours in the NAs (52–55). Although QYs were low in all labelled cTAR sequences, clear differences appeared as a function of the labelling positions. QYs were low when 2-Ap was inserted in ds-regions (positions 9, 17, 21, 45, 49, 53 and 55) that are known to favour base stacking, and thus the quenching of its fluorescence (53). The presence of guanine residues neighbouring 2-Ap was associated with a further decrease of the 2-Ap quantum yield, in full line with the most efficient quenching capabilities of Gs among bases (56). On the contrary, the presence of bulges or mismatches in the close vicinity of 2-Ap (positions 9, 53 and 55) was found to lead to somewhat higher QY values (57). By far, the highest QY value was observed when 2-Ap was inserted in the internal loop, suggesting a limited interaction of 2-Ap35 with its neighbours in this single-stranded segment of cTAR. In contrast to 2-Ap35, a significantly lower QY was observed for 2-Ap28, suggesting an efficient quenching of the probe by the G₃₂₉ residue in the loop. In line with nuclear magnetic resonance (NMR) data showing that the loop of cTAR is highly flexible (58), fast conformational fluctuations may drive 2-Ap28 from unstacked to stacked conformations, resulting in a low QY value. Taken together, we observed a strong dependence of 2-Ap fluorescence signals on the sequence and/or the structure of the DNA segments in which the 2-Ap is inserted. The 2-Ap signals easily discriminated stable regular ds-segments from positions destabilized by bulges or mismatches and clearly showed heterogeneous 2-Ap environments along the cTAR sequence.

To further characterize the local 2-Ap environment in the various labelled cTAR sequences, thermal melting curves were recorded using the 2-Ap fluorescence signal. These curves allow evidencing local conformations and dynamics of 2-Ap and its close neighbours (59). On melting, two types of transitions were observed. When the 2-Ap residue was located in ds-segments (positions 17, 21, 45 and 49), its fluorescence emission strongly increased during duplex melting (Figure 3A). This was ascribed to the progressive loss of base stacking as the temperature increases. The 2-Ap9 presented a similar transition pattern, suggesting that 2-Ap9 is likely base paired.
with C48. On the contrary, downward transitions were observed for 2-Ap53 and 2-Ap55 (Figure 3B), evidencing an extra-helical conformation of G54 when the last two ds-segments are formed (60). Indeed, such an extra-helical conformation is thought to limit the dynamical quenching of G54 with both 2-Ap53 and 2-Ap55, explaining a higher QY for the folded state than for the melted form. Moreover, the low melting temperature associated with the 2-Ap55 residue indicated that the last ds-segment is the least stable one. In addition, the broad melting transition observed with this residue further suggested a high propensity of the last ds-segment to transiently melt at room temperature, in line with previous reports (29,31).

More surprisingly, 2-Ap28 also underwent a downward melting transition (Figure 3B), probably as the consequence of the disappearance of the structural constraints within the apical loop when the upper ds-segment melted. Taken together, the range of melting temperatures of the different ds-segments of cTAR clearly supported a non–two-state melting transition (61,62) where ds-segments melted independently and where the upper part of the stem–loop seemed more stable than the lower part (Figure 3C).

Heterogeneity of the probe environment was further investigated using time-resolved experiments. In contrast to the mono-exponential fluorescence decay of free 2-Ap, the fluorescence decays of the 2-Ap-labelled cTAR sequences were multi-exponential. A satisfactory fit was obtained with 4 lifetime components, ranging from ~100 ps to nearly 10 ns, demonstrating the large conformational heterogeneity explored by the 2-Ap residues at each position (Table 1). The longest-lived component $t_4$ that likely corresponded to conformations where 2-Ap was either unstacked or extrahelical (63,64) varied from 6.1
parameters were determined in the absence of the extra-helical G54 conformation that prevented the stacking with its flanking residues. The three short-lived states, which may drive them from an unstacked to a stacked conformation, were largely dominated in the presence of 1–5 equivalents of NC(11–55) (+) or SSNS, NC(11–55) (M).

Beside the snapshot picture of the 2-Ap conformations distribution given by the time-resolved fluorescence decays, rotational dynamics of 2-Ap monitored through time-resolved fluorescence anisotropy provide information about the local and global dynamics of the cTAR sequences. Anisotropy decays were satisfactorily fitted using a three exponential model, with the exception of 2-Ap at position 55 for which only two components were sufficient. The short correlation time ($\tau_1$) of 2-Ap with respect to the strand backbone, whereas the longest rotational correlation time ($\tau_2$) described the whole tumbling of the molecule. The presence of an intermediate rotational correlation time $\tau_2$ (~1 ns) with an amplitude almost constant of about ~30% evidenced additional conformational fluctuations in cTAR. The amplitude of the short rotational correlation time reflected the conformational space explored by the 2-Ap residue during the excited time. The extent of the explored space was quantified by the cone semi-angle value $\theta_0$ (Table 1). The $\beta_1$ and $\theta_0$ values were low for 2-Ap residues located in ds-segments, but large for 2-Ap9 and 2-Ap49, indicating a high rotational freedom for the latter, in line with the higher propensity of mismatched bases to efficient quenching of 2-Ap28 by the G29 residue. In sharp contrast, a much more limited population (56%) of strongly stacked conformations was observed in the internal loop, probably because of the combination of a less efficient quenching by the flanking A together with constraints in the internal loop that prevent efficient quenching by the neighbour bases.
persist in the last ds-segments of the cTAR stem (dark grey).

ing 2-Ap residues at position 9, 45 and 49, whereas segmental motions largely decreases the segmental mobility of the DNA segments containing 2-Ap residues at position 9, 45 and 49, whereas segmental motions persist in the last ds-segments of the cTAR stem (dark grey).

Figure 4. Representation of the longest rotational correlation time $T_1$ in the absence and in the presence of NC(11–55). $T_1$ values for the 2-Ap residues inserted in the lower half of cTAR stem (hatched white bars) are significantly shorter than the values observed in the upper part of the sequence (hatched grey bars), indicating significant segmental motions in the lower part of the cTAR stem. In the presence of saturating concentrations of NC(11–55) (non hatched bars), NC(11–55) largely decreases the segmental mobility of the DNA segments containing 2-Ap residues at position 9, 45 and 49, whereas segmental motions persisted in the last ds-segments of the cTAR stem.

explore geometries outside from the helix. The values of the long rotational correlation time ($T_3$) ranged from 4 to 9 ns, depending on the labelled position. For 2-Ap residues in the upper half-part of the hairpin loop (at positions 17, 21, 28 or 35) (Figure 4), the $T_3$ value was ~8–9 ns, in full line with the theoretical correlation time calculated for the tumbling motion of a cylinder rod of length 95 Å and diameter 20 Å (66). In contrast, the $T_3$ values were substantially lower (~4–5 ns) for 2-Ap residues in the bottom of the cTAR sequence (positions 45, 49, 53 or 55) and were thus likely a combination of the overall tumbling of the cTAR sequence and of the segmental ‘fraying’ motion of the cTAR ends (30,34). Similarly to the $T_3$ value, the $T_2$ value was probably shortened in 2-Ap55 because of the larger segmental motions in the cTAR terminal ds-segment, making it impossible to resolve from $T_1$. This hypothesis was further supported by the unexpectedly high $\beta_1$ value observed for 2-Ap at position 55.

Effect of NC(11–55) on cTAR

Because of their low quantum yield, high concentrations of labelled NAs were required (typically 1–5 μM) to characterize the 2-Ap fluorescence parameters. This impeded us to use the full-length NCp7, which strongly aggregated nucleic acids, when added at these concentrations. The NC-induced destabilization of cTAR was thus investigated using NC(11–55), a mutant that lacks the major determinants of aggregation of the wild-type sequence but which is sufficient for specific binding and destabilization (24,35). Substitution of 2-Ap for an adenine in the cTAR sequence induced no detectable differences in the binding curves monitored through tryptophan quenching experiments, suggesting that 2-Ap did minimally affect NC(11–55) binding (see Supplementary Material and Supplementary Figure S4).

Fluorescence parameters of 2Ap-substituted cTAR in the presence of NC(11–55).

We investigated the effect of NC(11–55) with incremental addition of 1, 4, 7, 10 and 15 equivalents of peptide to the different 2-Ap-labelled cTAR sequences. NC(11–55) binding was confirmed by the progressive increase of the longest rotational correlation time $T_3$, as NC(11–55) concentration increased. $T_3$ reached values between 24 and 34 ns (Table 1 and Figure 4), confirming that ~10 NC(11–55) molecules bound to cTAR. When the 2-Ap residue was not located at the bottom of the stem, the $T_3$ values tended to be somewhat higher than the 30 ns value expected for a sphere with the same molecular mass (72 kDa) (66), suggesting that the global shape of the NC:cTAR complexes may differ from a sphere. With such a high-rotational correlation time, $T_3$ likely described the overall tumbling motion of the whole complex, with marginal contribution of segmental motions. Shorter $T_1$ values were only observed in the lower half part of the cTAR stem for the 2-Ap residues at positions 53 and 55 (Figure 4), suggesting that only the two terminal ds-segments kept a substantial mobility in the presence of NC(11–55). As a consequence, the binding of NC(11–55) to cTAR resulted in a significant decrease of the segmental mobility of the DNA segments containing 2-Ap9, 2-Ap45 and 2-Ap49, which were shown to undergo large segmental motions in the absence of NC(11–55).

Addition of NC(11–55) also increased the 2-Ap QY. In the presence of saturating concentrations of NC(11–55), the 2-Ap quantum yields varied from almost unchanged (2-Ap28) to an increase by a factor up to nearly 14 (2-Ap49) (Figure 2). The fluorescence changes were dependent on the amount of NC bound and on the position of the 2-Ap within the sequence. Time-resolved parameters further show that NC(11–55) did marginally change the lifetime values but induced a redistribution of the 2-Ap conformational states towards the longer-lived conformations (Table 1). The extent of this redistribution was dependent on the labelling position and on NC(11–55) concentration, being the most important at positions 9 and 49, showing that NC(11–55) strongly reduced base stacking at these two positions (Figure 5A). Stacked conformations were also decreased in positions 17, 21, 45, 53 and 55, albeit to a lesser extent, showing that NC(11–55) was able to bind and to decrease the base stacking all along the cTAR sequence (67). NCp7 seemed to be thus efficient to prevent the stacking of 2-Ap with its flanking bases, especially when 2-Ap is located close to a G. This likely resulted from the insertion of the neighbouring guanine into the hydrophobic platform at the top of the folded zinc-fingers of NC, which in turn prevents its collision with 2-Ap and thus, reduces its quenching efficiency (38,39,42,44). Moreover, the reduction of the amplitude associated with the shortest rotational correlation time $\Phi_1$, further indicates that NC(11–55) additionally restricts the base mobility on binding. Again, the $\Phi_1$ amplitude was the most decreased for 2-Ap at position 9 and 49, suggesting stable interactions of NC(11–55) with bases in the surrounding of 2-Ap in these two positions. The fluorescence parameters
associated to 2-Ap9 and 2-Ap49 suggested that the collisions of the 2-Ap with their neighbouring guanines (G10 and G50, respectively) were strongly prevented by NC(11–55).

Interestingly, in equimolar conditions where statistically one NC is bound per cTAR, a strong increase (~2-fold) in quantum yield was observed for 2-Ap49, whereas only marginal changes were observed for all other positions (Figure 2). The populations of stacked conformations and the cone semi-angle \( \theta_0 \) value of 2-Ap at these two positions were also significantly reduced (Figure 5A and B). These large fluorescence changes at a 1:1 stoichiometry suggest a high affinity of NC(11–55) for these positions. Therefore, it is likely that G10 and G50 constitute preferentially targeted guanines along cTAR and thus, constitute initial destabilization sites within the cTAR stem.

**Effect of (SSHS)2NC(11–55) mutant.**

To confirm that the decrease in base stacking and the restriction in base and/or segmental mobility observed on NC(11–55) binding resulted from a specific activity of NC ZFs on cTAR, we investigated the 2-Ap fluorescence parameters in the presence of (SSHS)2NC(11–55), a NC(11–55) mutant in which all cysteine residues have been changed for serines to prevent zinc binding, and thus the folding of the ZFs (68). Addition of saturating levels of (SSHS)2NC(11–55) affected poorly the mobility of cTAR bases, as seen from the small decrease of the \( \theta_0 \) value induced by (SSHS)2NC(11–55) at the different 2-Ap positions tested (Table 1). Moreover, the longest rotational correlation time was increased in the presence of (SSHS)2NC(11–55) [showing that (SSHS)2NC(11–55) was binding to cTAR] but remained significantly shorter than that observed in the presence of NC(11–55). Therefore, it is likely that G10 and G50 constitute preferentially targeted guanines along cTAR and thus, constitute initial destabilization sites within the cTAR stem.
K (70).

NC(11–55)

SSHS2NC(11–55)

which a sequence logo was derived (Figure 7). This

ure were used to calculate a position weight matrix from

The 587 sequences retrieved from the QuickAlign proced-

 subtype reference database (http://www.hiv.lanl.gov).

motifs within the HIV subtypes, we aligned the cTAR

whether these positions corresponded to conserved

recognition sites needed for efficient strand transfer, it is

notably through direct specific interactions with the G10

timescale into specific conformations (Figure 5B),

promoting the competent reactive species. This could be

the efficiency of the nucleation-limited reaction by

the cTAR reactive species, NC(11–55) also likely increases

transition free energy of base pair melting of cTAR.

(SSH$\text{S}_2$)NC(11–55) was at least one order of magnitude slower than that observed with NC(11–55).

Thermodynamic parameters of the reactions associated with the two NC mutants were determined (Table 2). The transition state enthalpy was estimated from the temperature dependence of the annealing reaction (Figure 6) according to:

$$\Delta H^\circ = -R \cdot d(\ln k_{obs})/d(1/T)$$

Compared with NC(11–55), the slow component of

(SSH$\text{S}_2$)NC(11–55) presented a largely increased $\Delta H^\circ$ value, indicating that more base pairs need to be melted for the reaction to proceed (Table 2). In contrast, the thermodynamic parameters observed with NC(11–55) were close to those associated with the fast component of (SSH$\text{S}_2$)NC(11–55). Together with the increase of the amplitude of this fast component with temperature (Figure 6), this suggests that this component may correspond to the (SSH$\text{S}_2$)NC(11–55)-promoted annealing of a cTAR population, which was partly melted by temperature before it was reacted with TAR. Thus, in full line with related studies describing the effect of NC on nucleic acid thermodynamics (71,72), NC(11–55) likely decreases the transition free energy of base pair melting of cTAR.

More than destabilizing the cTAR sequence to generate the cTAR reactive species, NC(11–55) also likely increases the efficiency of the nucleation-limited reaction by promoting the competent reactive species. This could be a consequence of the ability of NC(11–55) to freeze the bottom of cTAR stem at the picosecond–nanosecond timescale into specific conformations (Figure 5B), notably through direct specific interactions with the G$\text{G}_{10}$ and G$\text{G}_{50}$ residues.

Conservation of the cTAR sequence and structure

If G$\text{G}_{10}$ and G$\text{G}_{50}$ residues are critically involved in NCp7 recognition sites needed for efficient strand transfer, it is likely that these positions must be conserved. To check whether these positions corresponded to conserved motifs within the HIV subtypes, we aligned the cTAR sequence presented in Figure 1 to the LANL HIV subtype reference database (http://www.hiv.lanl.gov). The 587 sequences retrieved from the QuickAlign procedure were used to calculate a position weight matrix from which a sequence logo was derived (Figure 7). This

sequence logo is drawn such that the overall height of the stack indicates the information content at each position (derived from the Shannon entropy), whereas the height of the base symbols within the stack indicates their relative frequency (51,73,74). The information content, which can reach a maximal theoretical value of 2 bits for nucleic acids, is usually interpreted as an indicator of evolutionary conservation. The cTAR sequence seems to be well conserved, including the G$\text{G}_{10}$ and G$\text{G}_{50}$ residues. In contrast, residues A$\text{A}_{9}$ and T$\text{T}_{11}$ flanking G$\text{G}_{10}$ were associated to low information content as compared with the other residues of the cTAR ends, suggesting that the nature of the bases at these positions is of limited importance. As a result of this low base conservation, A$\text{A}_{9}$ was replaced by a T in one-third of the sequences. In addition, the C$\text{A}$48 residue was frequently replaced by an A or a T. Therefore, the A$\text{A}_{9}$–C$\text{A}_{48}$ mismatch is barely as probable as the canonical A–T base pair at this position (Figure 8B). However, a C$\text{A}_{11}$–A$\text{A}_{46}$ mismatch was also frequent, indicating that the position of the distal CA mismatch seems to be exchangeable on both sides of the G$\text{G}_{10}$–C$\text{A}_{47}$ base pair (data not shown). Similarly, a C$\text{A}_{5}$–A$\text{A}_{49}$ mismatch was also frequently found when the C$\text{A}_{5}$–A$\text{A}_{31}$ mismatch was absent. It resulted that albeit in different positions, CA mismatches were largely conserved, in close proximity of the conserved C$\text{G}_{5}$–G$\text{G}_{50}$ and G$\text{G}_{10}$–C$\text{A}_{47}$ base pairs, highlighting that their presence is required to finely tune the thermodynamic stability of the two terminal ds-segments of cTAR (75).

**DISCUSSION**

In this work, we investigated the steady-state and time-resolved fluorescence properties of cTAR labelled...
by 2-Ap residues at different positions. Fluorescence decay kinetics enabled the visualization of various 2-Ap conformers as snapshot pictures, whereas time-resolved fluorescence anisotropy provided complementary information on the associated rotational dynamics. This approach was thus well suited to monitor the structure and dynamics of the cTAR stem–loop and to probe the structural and dynamical perturbations induced on NC(11–55) binding. The base stacking patterns of 2-Ap seemed to be highly heterogeneous, adopting at least five conformations where the most stacked ones were the most represented. cTAR seemed to be highly dynamic at the picosecond–nanosecond timescale, especially in its lower half part that underwent segmental motions as the consequence of the fraying process that governs the dynamics of the cTAR stem ends (30,32,34). The terminal loop was found highly flexible, whereas the internal loop seemed to be highly constrained, in agreement with the slow micro-to-millisecond motions of the residues of the internal loop reported by NMR (58). Taken together, our data fit well with the NMR data of the top-half of cTAR (58) and the chemical probing experiments (67), as well as with the m-fold structure prediction of cTAR (76), except for the last ds-segment of cTAR that we found transiently annealed at 20°C in our conditions. Our data constitute thus a refined structural and dynamical model on the environment and the available conformational space for the base within the cTAR sequence.

Figure 7. Sequence logo of cTAR. Sequence logo resulting from 587 aligned cTAR sequences showed the globally strong conservation of the cTAR residues. Residues flanking the G10–C47 base pairs seemed to be among the least conserved.

Figure 8. Histogram of the mismatch frequencies for the different base pairs within cTAR. Mismatch base pairs are preferentially flanking the two well-conserved G10 and G50 residues identified as directly interacting with NC(11–55). These mismatch base pairs are conserved, suggesting a critical role of the mismatches for NC recognition.

Moreover, our data further suggest that the cTAR destabilization process is the result of the progressive coating of cTAR by NC(11–55). In contrast to 2-Ap9 and 2-Ap49 residues, which show subsequent changes in at least one of their fluorescence parameters at already low NC(11–55)/cTAR ratios, the 2-Ap45 residue exhibited fluorescence signal changes only for higher concentrations of NC(11–55), suggesting that the upper ds-DNA segment probed by 2-Ap45 can be destabilized only when its lower part is destabilized by NC(11–55). Moreover, as position 45 is only a few nucleotides apart from positions 9 or 49, the...
effects of NC on cTAR seem to be highly localized around its initial binding site. The latter is a prerequisite for transient interaction (78,79). Another consequence of this local NA remodelling is that effective chaperone activity requires a sufficient amount of bound NC, a situation which is achieved in the viral nucleoprotein complex context. The requirement for a sufficient level of coating to induce NA destabilization distant from its initiation site is in line with the fact that NC shows optimal chaperone activity when the target oligonucleotides are fully coated (21,29). This requirement also suggests that the initial and limited destabilization of cTAR can be extended by the binding of additional NC molecules, resulting in larger destabilized domains that likely persist long enough to efficiently react with the complementary TAR sequence. A progressive destabilization of cTAR stem as NC concentration increases was previously described (29) and likely explains the NC-coating dependent progressive switch in the cTAR/TAR annealing mechanism from a loop/loop mechanism to a zipper mechanism through the ends of their ds-stems (21).

The preferential binding of NC(11–55) to G10 and G50 residues showed that NC can discriminate and specifically target these two G residues within cTAR. Surprisingly, these guanines were located within motifs with low information content, suggesting that the nature of the bases flanking the targeted G seemed to be of limited importance for NC(11–55) recognition. However, the CA mismatches on the side of the G10–C47 and C7–G50 base pairs likely destabilize these base pairs and confer to the G10 and G50 residues the high degree of mobility (80), which was recently shown to be essential for recognition by NC (81). Thus, these intrinsic destabilizing CA motifs within the target DNA may facilitate the initial interaction of NC with the NA. The presence of mismatch base pairs seemed to play a more critical role in G recognition and discrimination than the nature of the NA sequence per se.

Similar conclusions have been previously made on MuLV Gag and NC showing that local context can considerably increase NC affinity for particular nucleic acid motifs (82). The conservation of the low stability of cTAR stem, probably co-evolving with NC chaperone properties, was shown to be critical for cTAR destabilization (31). This was confirmed by the early restoration of CA motifs within revertant viruses, which further highlights the importance of these destabilizing motifs in cTAR stem for the virus infectivity (83). Thus, this substantiates the hypothesis that NC destabilizes preferentially motifs with low stability, in line with the weak destabilizing activity of the NC ZFs (71,84). This finely tuned match between NC activity and the stability of the target NA sequence appears as an elegant way for NC to selectively chaperone a limited set of sequences within the HIV RNA genome and its DNA copies.

Moreover, the variability of the CA mismatch positions around G10 and G50 within the HIV-1 subtypes further highlights the necessary plasticity of NC to efficiently chaperone the first strand transfer. The flexible nature of NC must ensure sufficient plasticity to facilitate the correct and specific intercalation of the ZF aromatic residues to initiate the destabilization of cTAR, irrespective of the relative positions of the target G residues in respect with the CA mismatches. The poor sequence specificity and the high flexibility of NC likely ensure that NC can chaperone its various target nucleic acid sequences in the context of the high mutation rate of the virus. In addition, the high plasticity of NC and its unique ability to destabilize NA must provide NC a preferential and specific role in respect to other viral or cellular proteins (5,26,85,86) to timely and specifically enhance the cTAR/TAR annealing and to prevent self-priming (19). As HIV-1 has evolved to select such a complex interplay between NC and TAR structures and properties, the ability of NC to melt cTAR and to induce a specific

![Figure 9. Proposed mechanism for NC-induced destabilization of cTAR. NC(11–55) preferentially recognizes G10 and G50 and initiate the destabilization of cTAR. In the presence of higher concentrations, NC(11–55) can adjust to the heterogeneous structures and dynamics to bind all along cTAR and generate the reactive species needed to anneal the complementary TAR sequence.](https://academic.oup.com/nar/article-abstract/41/9/5036/2409011)

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annealing pathway through the bottom of these sequences is probably of major importance for the production of stable vDNA. As a result, therapeutic agents disturbing the fine-tuned equilibrium between cTAR stability and NC activity could probably significantly alter the specificity of the annealing pathway during the first strand transfer, and thus compromise the faithful and timely production of viral DNA (87–89).

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online: Supplementary Figures 1–5 and Supplementary Methods.

ACKNOWLEDGEMENTS

The authors are grateful to Hugues de Rocquigny for the peptide synthesis and purification. This article is dedicated to the memory of Dr Valérie Goldschmidt.

FUNDING

French agency against AIDS (ANRS); Agence Nationale de la Recherche [ANR-10-BLAN-1529]; University of Strasbourg. Funding for open access charge: Centre national de la recherche scientifique (CNRS).

Conflict of interest statement. None declared.

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