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Single-Molecule FRET Studies of Important Intermediates in the Nucleocapsid Protein Chaperoned Minus-Strand Transfer Step in HIV-1 Reverse Transcription

Running title: HIV-1 NC Chaperoned TAR DNA annealing

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Abstract

The minus-strand transfer step of HIV-1 reverse transcription is chaperoned by the nucleocapsid protein (NC), which has been shown to facilitate the annealing between the transactivation response element (TAR) RNA and complementary TAR DNA stem-loop structures. In this work, potential intermediates in the mechanism of NC-chaperoned TAR DNA/TAR RNA annealing have been examined using single molecule fluorescence resonance energy transfer. The interaction between TAR DNA and various DNA oligonucleotides designed to mimic the initial annealing step was monitored to capture potential intermediates along the reaction pathway. Two possible mechanisms of annealing were examined, namely nucleation through the 3'/5' termini, termed the "zipper" complex, or nucleation through the hairpin loops in a "kissing" complex. Intermediates associated with both mechanisms were observed in the presence of NC, and the kinetics of formation of these intermediates were also measured. Thus, the single molecule experiments support the notion that NC-assisted annealing of TAR DNA:TAR RNA may occur through multiple pathways.

Keywords: Single-molecule spectroscopy, Fluorescence resonance energy transfer, TAR DNA:RNA annealing

Abbreviations: NC, nucleocapsid protein; SMFRET, single-molecule fluorescence resonance energy transfer; TAR, transactivation response element; APD, avalanche photodiode; PEG, polyethylene glycol.

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INTRODUCTION

The HIV-1 nucleocapsid protein, NC, is composed of 55 amino acids, with two zinc-binding domains of the CCHC type (1-4). NC exhibits both non-specific and specific nucleic acid binding properties (5,6). It is thought that the zinc finger domain is responsible for sequence-specific base interactions (7) with a preference for TG rich regions (8-10), while the N terminal domain of NC interacts nonspecifically by electrostatic interactions (11,12).

NC plays multiple roles in the HIV-1 life cycle (5). In its role as a structural protein, NC stabilizes the virion by enhancing protein-protein interactions (13) and promoting the dimerization of viral RNA (14-18). NC also serves as a chaperone, by promoting several annealing reactions in the reverse transcription process. For example, NC has been shown to facilitate the annealing of the tRNA primer onto the primer binding site (19-21), annealing of complementary sequences in plus-strand transfer (22-25) and annealing of complementary repeat regions in minus-strand transfer (23,26-32).

The minus-strand transfer step in HIV-1 reverse transcription involves annealing of the newly synthesized minus-strand strong stop DNA to the complementary RNA located at the 3' end of the viral genome, so that reverse transcription can continue. This process, although thermodynamically favorable, does not occur with significant yield without the presence of NC (7,26). Both the RNA and DNA sequences can fold into stable hairpins known as trans-activation response element (TAR) RNA and TAR DNA, respectively (33-36). NC's importance in promoting minus-strand transfer has been well documented (5), however, the detailed mechanism of the annealing step in minus-strand transfer is not completely understood.

Several attempts have been made to provide more molecular-level details about NC's effects on oligonucleotide structure. Kinetic experiments suggest that NC's slight preference for binding to single stranded DNA could provide insight into the strand transfer process (37). Ensemble fluorescence resonance energy transfer (FRET) experiments conducted on TAR DNA have been used to show that NC destabilizes the double-helical structure (31,38,39). Further experiments detail a mechanism in which NC chaperones annealing by increasing conformational flexibility in the precursors, energetically opening pathways and thereby increasing the probability of encountering the most desired product (31,38-40).

In 2004, single-molecule FRET, SMFRET, was used to characterize the TAR DNA secondary structure in the presence of NC. It was determined that NC destabilizes the secondary structure in the 3'/5' terminal loop regions of the initially closed TAR DNA, termed the **C** form (41). This process is illustrated in Scheme 1, in which the four bulge/loop regions are indicated as L1-L4. The resultant system in the presence of NC is an equilibrium between **C** and the partially open TAR DNA/NC complex, termed the **Y** structure.

Recently, an in depth SMFRET study of the C/Y equilibration dynamics was undertaken, revealing a strong dependence of the FRET dynamics on both the NC and Mg concentrations (42) leading to several new insights on the C/Y interconversion mechanism. These single molecule results indicate that NC induces a highly dynamic Y structure, as compared to the relatively conformationally static C form. The C/Y dynamics were also observed to be significantly heterogeneous, indicating that the C/Y interconversion is not a simple two-state equilibration but rather must involve multiple conformational intermediates and/or multiple reaction pathways (42). The heterogeneity and the strong dependence of the dynamics on NC concentration is not unexpected since as many as 8 NC molecules can be simultaneously bound to TAR DNA(5). In this paper, a single NC concentration of 445 nM is used for the various annealing studies, in order to simplify the analysis and to mimic the relatively high NC concentration range believed to be relevant to the biological situation. In summary, single molecule FRET data strongly suggest that the Y structure corresponds to a range of secondary structures in the L1-L2 region and a range of numbers of bound NC molecules.

We have suggested that the Y structure might serve as a nucleation center for minus-strand transfer (41). In such a 'zipper' mechanism, nucleation occurs through either the 3' or 5' terminus in the NC-induced single-stranded form. In this model, the formation of such an intermediate increases the efficiency of forming a correctly annealed end product, and meanwhile decreases the probability of misfolded side products. The zipper mechanism is illustrated in Scheme 2 (left), where for brevity nucleation through both termini is shown.

The second mechanism that has been proposed is the kissing mechanism (43), in which nucleation occurs through the complementary single-stranded loop sequences. NC-induced kissing complexes have been found to be important in stabilizing RNA dimerization in the viral envelope (18,44-47). For minus-strand transfer this model involves the formation of inter-strand base pairs via loop-loop kissing, followed by formation of the extended DNA:RNA duplex. This mechanism also provides a specific nucleation site and decreases the likelihood of unfavorable side products. The formation of kissing complexes has been verified by NMR structural examinations of RNA/RNA interactions (48-50). The kissing mechanism is illustrated on the right in Scheme 2.

The current work represents an effort to apply SMFRET to characterize the two pathways that have been proposed for minus-strand transfer in HIV-1 reverse transcription. SMFRET was used to monitor the NC chaperoned interaction between TAR DNA and various DNA oligonucleotides that are analogs to the complementary RNA. Each DNA sequence was chosen for its specific complementarity to regions of TAR DNA characteristic of each of the types of pathways under consideration. Donor/acceptor labeled TAR DNA was used to monitor the effects of the interactions on the hairpin structure. Alternately, the acceptor dye was placed on the oligomers, which yields information about the annealing between the two species. The annealing kinetics of each reaction pathway were also compared.

MATERIALS AND METHODS

Coverslip and Chamber Preparation

Coverslips were cleaned in piranha solutions (25% H₂O₂ and 75% conc. H₂SO₄) for 1 hour, followed by various water (molecular biology grade), and acetone (HPLC grade) rinsing cycles. Dry, clean coverslips were then treated with vectabond/acetone 1% w/v solutions (Vector Laboratories, Burlingame, CA) for 5 min. Coverslips were then rinsed with H₂O and dried under a N₂ stream.

The clean coverslips were masked with patterned silicone films. The unprotected area was incubated with a 25% w/w polyethylene glycol (PEG) solution (MW 2000, Nektar Therapeutics, Huntsville, AL) containing 0.25% w/w biotinylated PEG (MW 5000, Nektar Therapeutics, Huntsville, AL) in a 0.1 M sodium bicarbonate solution (HyClone, Logan, UT) for 3h. The silicone templates were removed, the excess PEG rinsed with water, and the coverslips dried under a N₂ stream. Predrilled polycarbonate films with an adhesive gasket (Grace Bio-Labs, Bend, OR) were assembled on top of cleaned coverslips yielding a chamber with a total volume of ~5 μ L. The chamber was assembled on top of the PEG treated surface; the adhesive gasket adhering to the silicone template protected regions of the PEG treated coverslips. Inlet and outlet ports (NanoportTM, Upchurch Scientific, Oak Harbor, WA) were glued on top of the chambers.

DNA and Protein Preparation

All oligonucleotides were purchased in lyophilized form from TriLink Biotechnologies. The primary structure of each sequence used in the present study is detailed in Scheme 3, along with the complementarity between the TAR DNA and each oligonucleotide. Dye-labeled species were ordered with additional 5'T and/or 3' TTTT overhangs at the termini relative to the native form to prevent unwanted photophysics caused by G residue quenching (31,51,52).

Synthetic NC protein, with Met 46 mutated to norleucine to avoid oxidation to Met(O), was prepared chemically via Fmoc solid-phase synthesis carried out with aid of a 433A Peptide Synthesizer (Applied Biosystems, Framingham MA). The C-terminal residue, Asn, was anchored to PAL-PEG-PS resin via its side-chain (i.e. by coupling Fmoc-Asp-*O*tBu, as first residue). Deprotection/coupling cycles followed manufacturer recommendations, with all residues double coupled. The fully assembled peptide-resin was cleaved with reagent K (53). The crude product was purified by reversed-phase HPLC to give a material of ~ 87% homogeneity (as judged by analytical HPLC) (MALDI-MS; [M + H]_{calc.}: 6330.2, [M + H]_{found}: 6330.1). After lyophilization, the purified synthetic NC was reconstituted by assuming that the recovered white solid contained 100% peptide content. The solid was dissolved (0.6 mL/mg peptide) in reconstitution buffer (40 mM HEPES, 5 mM DTT, 0.1 mM tris(2-carboxyethyl)phosphine hydrochloride) that contained 3.0 equivalents of ZnCl₂, and lyophilized for long-term storage. It should be noted that the purified NC before reconstitution often contains a disulfide bond, based on mass spectrometric analysis. The

reconstitution buffer contains DTT that reduces the disulfides and allows correct folding of the Zn fingers to occur. The chaperone activity of the synthetic NC was shown to be higher than NC prepared previously from *E. coli* expression.

DNA Immobilization

The reaction chambers were incubated with streptavidin (Molecular Probes, Eugene, OR; 0.2 mg/mL in HEPES buffer) in 25 mM HEPES buffer, pH 7.3, 40 mM NaCl (buffer A) for 10 minutes, after which the chambers were rinsed with buffer A. In all cases, except for as noted in the text, the experiments were performed on TAR DNA immobilized through a biotin linkage at T28. The exceptions refer to the use of 'inverted' TAR, in which the biotin immobilization was performed at T64. The primary structure of each, including the respective location of the biotin, is listed in Scheme 3. Subsequently, the biotinylated TAR DNA solutions (25–50 pM) in buffer A were incubated in the reaction chamber for 20 min. The chambers were then filled with buffer A and Teflon tubing was adapted to the chamber ports. Two syringe pumps delivered buffer A/reactant solutions at a rate of 2 μ l/min. A 10-min equilibration period at a flow rate of 10 μ l/min was elapsed before measurements were taken under a new set of conditions. All solutions contained buffer A, 2 mM MgCl₂ (Ambion, Inc., Austin, TX), and an oxygen scavenger system (1% v/v 2-mercaptoethanol (Sigma-Aldrich, St. Louis, MO), 3% w/v β -D(+)-glucose (Sigma-Aldrich), 0.1 mg/mL glucose oxidase (Roche Applied Science, Hague Road, IN), and 0.02 mg/mL catalase (Roche Applied Science) (54,55).

Experimental Setup

The experimental setup has been described previously (41,56,57). Briefly, a closed-loop sample scanning stage (NPS-XY-100A, Queensgate, Torquay, Devon, U.K.) was used for imaging and sample positioning. Continuous wave excitation (514 nm, 5 to 10 μ W/ μ m²) from an argon ion laser (model Reliant 150m, Laser Physics, Inc., West Jordan, UT) was introduced via an optical fiber and was directed by a dichroic beamsplitter (530 DCLP, Chroma, Rockingham, VT) to the sample via a high numerical aperture oil immersion microscope objective (Zeiss Fluor, 100X, NA 1.3). Fluorescence was retrofocused using the excitation objective and split from scattered excitation light with a dichroic beamsplitter and a holographic Raman notch filter (Kaiser Optical Systems Inc., Ann Arbor, MI). Donor and acceptor fluorescence were separated with a second beamsplitter (Chroma 630 DCXR) and then directed to two avalanche photodiode (APD) detectors (Perkin Elmer Optoelectronics SPCM-AQR-15, Vaudreuil, Quebec, Canada). The intensity time trajectories were acquired with 1 ms time resolution.

Data Analysis

Single hairpin fluorescence intensity time trajectories were recorded using separate donor and acceptor channels. The signals were corrected for background emission/noise and donor/acceptor cross talk due to overlapping emission as previously described (41). Blinking events (reversible acceptor photobleaching) were removed prior to data analysis as previously described (41). The corrected intensity trajectories for the donor and acceptor channels, $I_A(t)$ and $I_D(t)$, respectively, were binned in the appropriate widths,

referred to as τ_B . In this way the single molecule FRET efficiency, $E_{FRET}(t)$ can be defined as:

$$E_{FRET}(t) = \frac{I_A(t)}{I_A(t) + I_D(t) \times \frac{f_A \times h_A}{f_D \times h_D}} \quad (1)$$

where ϕ_i is the quantum yield of the respective dye, and η_i is the efficiency of each detector. What is measured, however, is the apparent FRET efficiency, E_A , defined as:

$$E_A = \frac{I_A}{I_A + I_D} \quad (2)$$

In order to convert E_A into E_{FRET} , it is necessary to determine the correction factor describing the dye/detector efficiencies. The correction factor was obtained by comparing emission under 100% and 0% energy transfer conditions, and was found to be ~ 1 in the current experiment. Thus, the value obtained in each case for E_A represents almost exactly the true value of E_{FRET} . E_A was calculated for each binned point over the entire trajectory. Filtering of low E_A systems (e.g. 0.4) was not performed because of shot noise limits.

E_A autocorrelation amplitudes were estimated, as described in a previous study (41) using the single molecule cross correlation curves and the following expression,

$$C(\mathbf{t}) = \frac{\langle \mathbf{d}I_D(t) \mathbf{d}I_A(t+\mathbf{t}) \rangle}{\langle I_D \rangle \langle I_A \rangle} = \frac{\langle I_D(t) I_A(t+\mathbf{t}) \rangle}{\langle I_D \rangle \langle I_A \rangle} - 1 \quad (3)$$

$$E_A \text{ Autocorrelation}(\mathbf{t}) = C(\mathbf{t}) \left(- \frac{\langle I_D \rangle}{\langle I_A \rangle} \right) \quad (4)$$

where τ is the lag time. The E_A autocorrelation amplitude is a measure of the variance of the nucleic acid 3'/5' end-to-end distance FRET fluctuations. It is given by $C(\delta\tau)$ where $\delta\tau$ is the time spacing in the $E_{FRET}(t)$ measurements. It is the earliest value of $C(\tau)$ that is not affected by shot noise. For the qualitative purpose of the current work we simply report the order of magnitude of the observed autocorrelation amplitudes, and do not report the correlation time (typically a few ms) or the functional form of the observed autocorrelation decay, which is beyond the scope of the present study. For many of the observations the autocorrelation amplitudes were undetectably small (i.e. $< 5 \times 10^{-4}$) indicating a relatively static structure analogous to a DNA duplex structure in the absence of NC. In other cases, relatively large autocorrelation amplitudes were observed indicating large amplitude TAR DNA secondary structure fluctuations.

Single molecule kinetic experiments of annealing of donor-labeled TAR DNA to the zipper mechanism mimic, A-27-A and the kissing mechanism mimic, A-24-B (Fig. 4), were accomplished by recording multiple single molecule confocal images of a $30 \mu\text{m} \times 30 \mu\text{m}$ regions at various time after introduction of the DNA mimics to a sample cell containing immobilized donor-labeled TAR DNA. Each image frame contained 100-200 single molecule spots. The total scan time for each images frame was 128 sec. The

donor and acceptor images were collected simultaneously. Photobleached molecules were eliminated from the statistical analysis. The data were analyzed automatically by a computer program that determined the location and intensity of each molecule in the donor and acceptor images for each frame.

A FRET value was calculated for each molecule using the observed intensities, and corrected for crosstalk and background intensity. For each molecule in each frame, the FRET value was used to classify each molecule as either un-annealed donor-labeled TAR DNA or the annealed product. FRET histograms for each frame showed well resolved peaks for annealed and unannealed TAR DNA, that are analogous to Fig. 3. The reported reaction time, τ , was determined by fitting the % annealed molecules vs. time curves to a first-order kinetic model, with a rate constant equal to $1/\tau$.

RESULTS AND DISCUSSION

Columns 1 and 2 of Fig. 1 summarize the results of a SMFRET analysis of donor/acceptor labeled TAR DNA alone and in the presence of NC, respectively. The individual single molecule E_A histograms (obtained from the single molecule E_A trajectories) can be combined in an ‘ensemble’ histogram. In this case, the term ‘ensemble’ does not imply that the E_A values have been ensemble averaged. The resultant ensemble E_A histograms, gathered from 20-100 single molecules and comprised of thousands of occurrences, are normalized to a common scale. Ensemble E_A histograms for donor/acceptor labeled TAR DNA in the absence of NC are shown in column 1 of Fig. 1. The data are presented at three different binning times, τ_B : 5, 25, and 250 ms. The near unity E_A value for donor/acceptor labeled TAR DNA in the absence of NC indicates that it is in its **C** TAR form, shown in Scheme 1. The 5 ms τ_B data are broadened primarily by shot noise, but these effects are virtually eliminated once the data have been binned into 250 ms widths. The small broadening in the distribution is due to experimental/data analysis errors. The ~ 1 E_A , narrow distribution width, and zero E_A autocorrelation amplitude (not shown) support the model of **C** TAR as a static system with intact secondary structure.

In contrast, when 445 nM NC is added (Fig. 1, column 2) the histogram of E_A values becomes bimodal, indicating a dynamic distribution of both **C** and **Y** forms. Our previous results have shown that the NC-induced structural transformation between **C** and **Y** form (41) occurs on a ~ 5 ms time scale. Compared with the results in the absence of NC (Fig. 1, columns 1 and 2, bottom), the significant broadening of the E_A histogram at 250 ms τ_B in the presence of NC further indicates there is a slow fluctuation occurring at >250 ms. These results, and the detailed analyses presented elsewhere(41,42), reveal that there are considerable NC-dependent heterogeneous fluctuation in the TAR DNA secondary structure. This dynamic heterogeneity was found to be highly dependent on NC concentration, and suggests that the effects of NC are cooperative (i.e., multiple NC molecules are required for even the simplest dynamics described in ref. (42)). In the annealing experiments presented below, the NC concentration was held constant at 445

nM, two orders of magnitude above the limiting concentration, assuming an approximate binding ratio of 1 NC per 7 nucleotides (58).

The critical dependence of NC activity on the TAR DNA secondary structure can also be observed by introducing a single-stranded noncomplementary oligonucleotide to the system. The presence of the noncomplementary oligonucleotide, 12-N, at high concentration (100nM), only affects the relative distribution of **C** and **Y** forms as indicated by the histograms shown at different τ_B in column 3 of Fig. 1. The ensemble E_A values show that the TAR DNA structure has shifted towards the **C** form. These results suggest that the effect of 12-N is to reduce the activity of NC. NC has a high binding efficiency for both TAR DNA and for the single-stranded 12-N. The lower effective concentration of NC results in increased occurrence of the **C** form of TAR DNA. These results are consistent with detailed NC concentration-dependent studies presented elsewhere (41,42).

The situation is different for the case of the complementary oligomer, 12-C. An additional feature ($E_A \sim 0.5$) is present in the E_A distribution shown in column 4 of Fig. 1, which corresponds to the binding between 12-C and TAR DNA. The process, however, is reversible on a ~ 10 s time scale (see further results on A-13-C, discussed below). Retention of separate species at high τ_B indicates that the lifetimes of the bound and unbound systems of 12-C with TAR DNA are longer than 250 ms. Additionally, a similar shift from **Y** TAR to **C** TAR can be observed as described for 12-N above. The cause is similar: competition for NC by the single-stranded 12-C decreases the activity of NC towards destabilizing L1-L2. These results indicate that nucleation with a complementary oligonucleotide can occur at the 3' end of the **Y** form of TAR DNA, providing support for the zipper pathway.

The dynamic equilibrium for the binding of shorter oligomers to TAR DNA is demonstrated in Fig. 2. Here, the E_A histograms for the reversible binding of 12-C to donor/acceptor labeled TAR DNA are compared for two different oligomer concentrations. In each case the histogram indicates the presence of three distinct species corresponding to **C** TAR, **Y** TAR, and the bound 12-C/TAR DNA complex. The area of each peak in the histogram reflects the relative concentration of each form at equilibrium. When the 12-C concentration is increased to 100 nM, amplitude ratios shift to increased populations of bound and **C** forms relative to the **Y** form. This is due both to a higher 12-C concentration induced increase in yield, in the case of the bound form, and to decreased activity of NC, in the case of the **Y** and **C** forms.

To further examine this process, the interaction between donor labeled TAR DNA and acceptor labeled A-27-A (Scheme 3) was investigated. This construct was designed to mimic zipper nucleation at both termini. In contrast to the results obtained with 12-C, the reaction between TAR DNA and A-27-A occurs irreversibly within the time frame of the experiment, and results in a narrow distribution of E_A at ~ 1 (Fig. 3 A). The high E_A is consistent with the structure illustrated (Fig. 3 A, left) in which the two dyes are in close proximity due to complete annealing. An undetectably small E_A autocorrelation amplitude was observed indicating a static, fully annealed product. These results provide

support for a stable structure that is an excellent candidate for an intermediate occurring by the zipper mechanism.

When the analogous experiment was performed using a DNA oligonucleotide designed to interact via the kissing mechanism, A-24-B, the formation of a stably annealed product was also observed (Fig. 3 B). These data indicate that the kissing mechanism is also viable. However, the formation of the duplex via the kissing interaction required a longer time relative to the zipper construct (see below for further discussion), and in fact did not react to completion within the experimental observation time (~4 hours; see discussion of Fig. 4 for further detail). This is indicated by the retention of unreacted donor-only emission observed in the E_A histogram in Fig. 3 B (center column). Representative single molecule E_A trajectories, $E_A(t)$, for reacted and unreacted molecules are shown in the right column and its inset. The time scale for the formation of the kissing loop intermediate was similarly slow when inverted TAR DNA was used. The $\langle E_A \rangle$ of ~0.8 for the annealed product is consistent with an increased distance between the two dyes, as illustrated (Fig. 3 B, left). Like the zipper reaction mentioned above, the reaction was irreversible within the experimental time frame. However, both the relatively broad E_A distribution and a relatively large E_A autocorrelation amplitude ($\sim 10^{-3}$) suggest that the resultant annealed structure is highly dynamic, compared to the product formed when TAR DNA was annealed with the zipper mimic discussed earlier. This is most likely due to residual fluctuations in the un-annealed 3' and 5' termini of TAR DNA, as observed earlier.

In contrast to the irreversible reactions observed in zipper and kissing reactions, reversible binding was observed when SMFRET was performed using shorter acceptor-labeled oligomers complementary to either the L1-L2 region (A-13-C) or the L3-L4 region (A-14-D). The ensemble E_A histograms for each reaction are shown in Fig. 3 C and D, respectively. Single molecule trajectories in the right column for the nucleation of these shorter oligomers indicate binding reversibility, similar to the results described earlier for 12-C (Fig. 1). Thus, in the ensemble E_A histogram for each reaction, the relative distribution of bound and unbound states reflects the equilibrium process. The K_{eq} for the reactions shown in Fig. 3 C and D were calculated to be 0.62 and 1.19, respectively. The ensemble E_A histograms for these experiments, shown in the center column of Fig. 3 C and D, yield evidence that in the presence of NC, the annealing can occur at any nucleation site along the TAR DNA hairpin. This is despite previous observations that the secondary structure of TAR DNA is not measurably weakened in the presence of NC in any region other than L1-L2 (41). These experiments also verify that the annealing processes are not perturbed by dye location. Non-specific binding can be ruled out because no reaction was observed when acceptor-labeled noncomplementary A-13-N was used, as shown in Fig. 3 E.

The results presented in Fig. 3 demonstrate that annealing can be initiated at multiple locations on the TAR DNA hairpin. Also, the relative yield of the kissing product versus the zipper product observed at room temperature suggests that the kinetic pathways for each type of intermediate may be different. In order to evaluate this hypothesis in more depth, the kinetics of formation of each type of product was measured, as shown in Fig.

4. As can be clearly observed, the zipper product was formed more rapidly than the kissing product, with reaction lifetimes of 7 minutes, and 37 minutes, respectively. The analogous two reactions were also performed using inverted donor-labeled TAR DNA, yielding first order lifetimes of <3 minutes and 21 minutes for the zipper and kissing products, respectively. (The ~ 3 minute value is close to the instrument response time which is limited by the confocal scan time per image, i.e. 2 min.).

These kinetic results are consistent with our previous observation that in the presence of NC, the secondary structure in the L1-L2 regions of TAR DNA is disrupted (41,42), allowing for more rapid annealing by the zipper mechanism. Interestingly, the kissing product formation lifetime was faster when the immobilization site was moved away from the hairpin region. This helps to confirm that the kissing product forms through inter-hairpin interactions, because biotin immobilization in the hairpin loop would be expected to inhibit loop-loop interactions.

These results demonstrate that the zipper product is the more efficient pathway for the annealing of TAR DNA with DNA oligonucleotides. It is not the only pathway, however, and several types of evidence suggest that the kissing intermediate should also be considered as a candidate for TAR/TAR annealing (18,43). Also, MFOLD predicts that the kissing intermediate is more thermodynamically stable than the zipper intermediate(59-62). The competition between kinetic and thermodynamic reaction control was tested using a competitive annealing experiment, in which both zipper and kissing intermediates were allowed to react simultaneously with TAR DNA. When both A-27-A and A-24-B were reacted with donor-labeled TAR DNA, the E_A histogram, shown in Fig. 5 A, suggests that the zipper product dominates. The results of a control experiment (Fig. 5 B), in which only the kissing oligomer was dye-labeled, suggests that both types of products were formed, however. Thus, although the zipper product forms more rapidly, as the kinetically favored pathway, the kissing product forms as well, consistent with its predicted thermodynamic stability.

CONCLUSIONS

The mechanism of NC chaperoned TAR DNA/TAR RNA annealing has been examined using TAR DNA and DNA oligonucleotides mimicking the potential annealing intermediates. In all cases, the previously reported Y structure of TAR DNA is preserved in the presence of NC and various oligonucleotides. It was found that annealing occurs through both zipper and kissing intermediates, although kinetic experiments indicate that the zipper product formation occurs more rapidly. In competitive reaction experiments, both types of intermediates are in fact observed. The results support a model of minus-strand transfer in which NC promotes the formation of multiple intermediates through which the fully formed product may occur.

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CAPTIONS

Scheme 1 Previously, SMFRET was used to demonstrate that, in the presence of NC, the TAR DNA hairpin is ‘open’ through the terminal two loop regions (the four loops are indicated above by L1-L4) (41). The partially open form that exists in the presence of NC is termed the **Y** Form.

Scheme 2 The two intermediates in the NC chaperoned annealing of TAR DNA and TAR RNA that are examined in the current study are illustrated. *Left:* A zipper mechanism can occur if annealing initiates through the 3’ or 5’ terminus (or both simultaneously, as shown in the case of a **Y:Y** complex). *Right:* A kissing mechanism occurs if annealing is initiated through interaction between the bottom loop regions.

Scheme 3 Proposed secondary structure of TAR DNA (right) and sequences of complementary and noncomplementary DNA oligonucleotides used in this study (left) are shown. Right: Region A, B, C and D represent the regions complementary to the DNA oligonucleotides shown on the left. Left: Oligonucleotides preceded by ‘A’ are Cy5-labeled. A-13-N and 12-N are oligonucleotides designed to be non-complementary to TAR DNA, as defined by fewer than 4 consecutive Watson-Crick base pairs.

Figure 1 Ensemble E_A values of doubly-labeled TAR DNA at three different τ_B under different reaction conditions are shown. (*Column 1*) E_A histogram before the addition of NC. (*Column 2*) E_A histogram after addition of 445 nM NC. (*Column 3*) E_A in the presence of 445 nM NC and 100 nM noncomplementary 12-N oligomer. (*Column 4*): E_A in the presence of 445 nM NC and 100 nM complementary 12-C oligomer. All experiments were performed at 2 mM Mg^{2+}

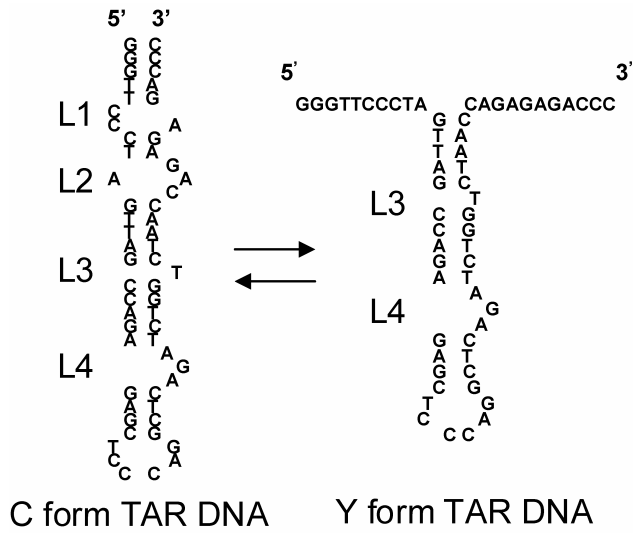
Figure 2 A summary of the reaction between doubly-labeled TAR DNA and oligonucleotide 12-C at two concentrations is shown. The ensemble distributions of E_A were obtained at 2 mM Mg^{2+} , 445 nM NC, and 10 nM 12-C (top) or 100 nM 12-C (bottom). At higher [12-C], the structure of TAR is shifted from **Y** states to bound states. The data are presented using $\tau_B = 10$ ms.

Figure 3 (A-E) Ensemble E_A histograms (*Middle*) and representative single molecule E_A time trajectories (*Right*) are shown for the reactions shown at left. All measurements were performed using donor-labeled TAR DNA and acceptor-labeled oligonucleotides in the presence of 445 nM NC and 2 mM Mg^{2+} . The data are presented using $\tau_B = 50$ ms. The oligonucleotides used are: (A) 10 nM A-27-A, (B) 10 nM A-24-B. The inset shows that no observable binding occurs in a minor population of the single molecules measured. (C) For 100 nM A-13-C, the distributions of E_A and the E_A trajectory exhibit reversibility. (D) Similar reversible binding was observed with A-14-D, and (E) 100 nM A13-N. No binding was observed for 100 nM A-13-N.

Figure 4 Kinetic curves for the annealing rates of the zipper mimic, A-27-A, vs. the kissing mimic, A-24-B, with donor-labeled TAR DNA. These experiments were performed in the presence of 445 nM NC and 2 mM Mg^{2+} . The oligonucleotide concentrations were 10 nM in both cases. Analogous reactions with inverted donor-labeled TAR DNA are also included in the figure.

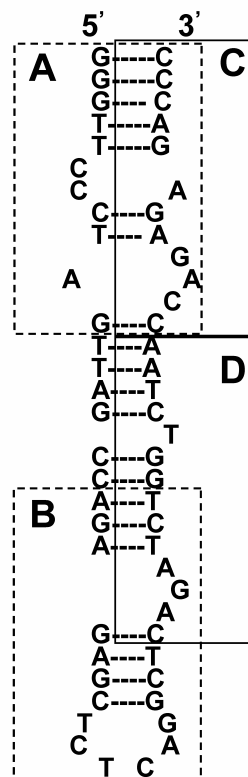
Figure 5 (A) An ensemble E_A histogram is shown for the simultaneous reaction of A-27-A and A-24-B with donor-labeled TAR DNA at $[Mg^{2+}] = 2$ mM and $[NC] = 445$ nM. The $\langle E_A \rangle$ of 0.92 is consistent with A-27-A binding. (B) An ensemble E_A histogram for the reaction of 23-A and A-24-B with donor-labeled TAR DNA under similar reaction conditions results in an $\langle E_A \rangle$ of 0.75, consistent with A-24-B binding. The data are presented using $\tau_B = 10$ ms.

SCHEMES



SCHEME 1

Oligonucleotide	Primary Structure
donor/ acceptor-labeled TAR DNA	5'-Cy3-TGGGTTCCCTAGTTAGCCAGAGAGCTCT(biotin)CAGGCTCAGATCTGGTCTAACCAGAGAGACCCTTTT-Cy5-3'
donor-labeled TAR DNA	5'-Cy3-TGGGTTCCCTAGTTAGCCAGAGAGCTCT(biotin)CAGGCTCAGATCTGGTCTAACCAGAGAGACCCTTTT-3'
inverted donor-labeled TAR DNA	5'-Cy3-TGGGTTCCCTAGTTAGCCAGAGAGCTCTCAGGCTCAGATCTGGTCTAACCAGAGAGACCCTTTT(biotin)-3'
12-N	5'-GCCGTA AAAATTT-3'
12-C	5'-GGGTCTCTCTGG-3'
23-A	5'-GGGTCTCTCTGGCTAGGGAACCC-3'
A-27-A	5'-GGGTCTCTCTGGCTAGGGAACCCTTTT-Cy5-3'
A-24-B	5'-AGATCTGAGCCTGAGAGCTCTCTT-Cy5-3'
A-13-C	5'-Cy5-TGGGTCTCTCTGG-3'
A-14-D	5'-Cy5-TTAGACCAGATCTG-3'
A-13-N	5'-GCCGTA AAAATTT-Cy5-3'



SCHEME 3

FIGURES

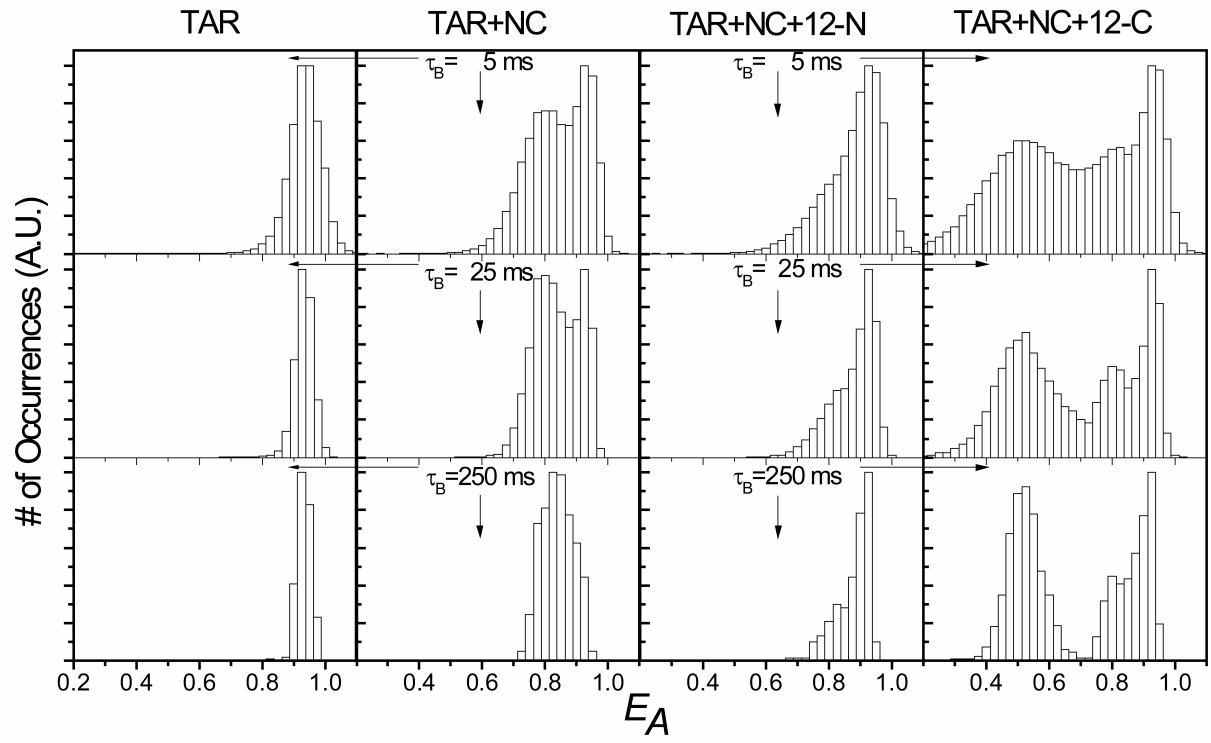


FIGURE 1

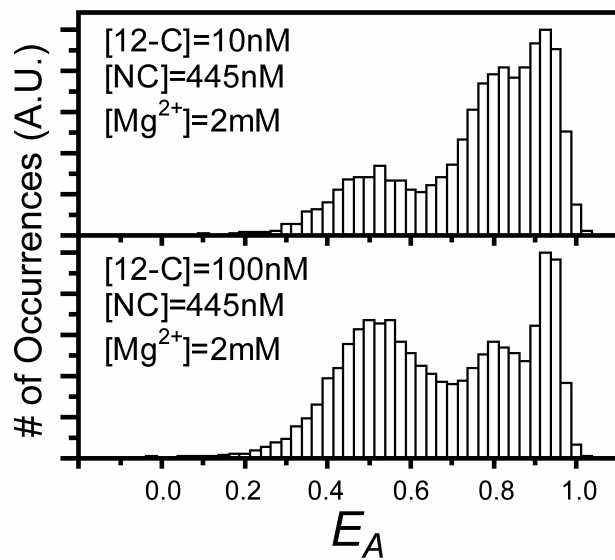


FIGURE 2

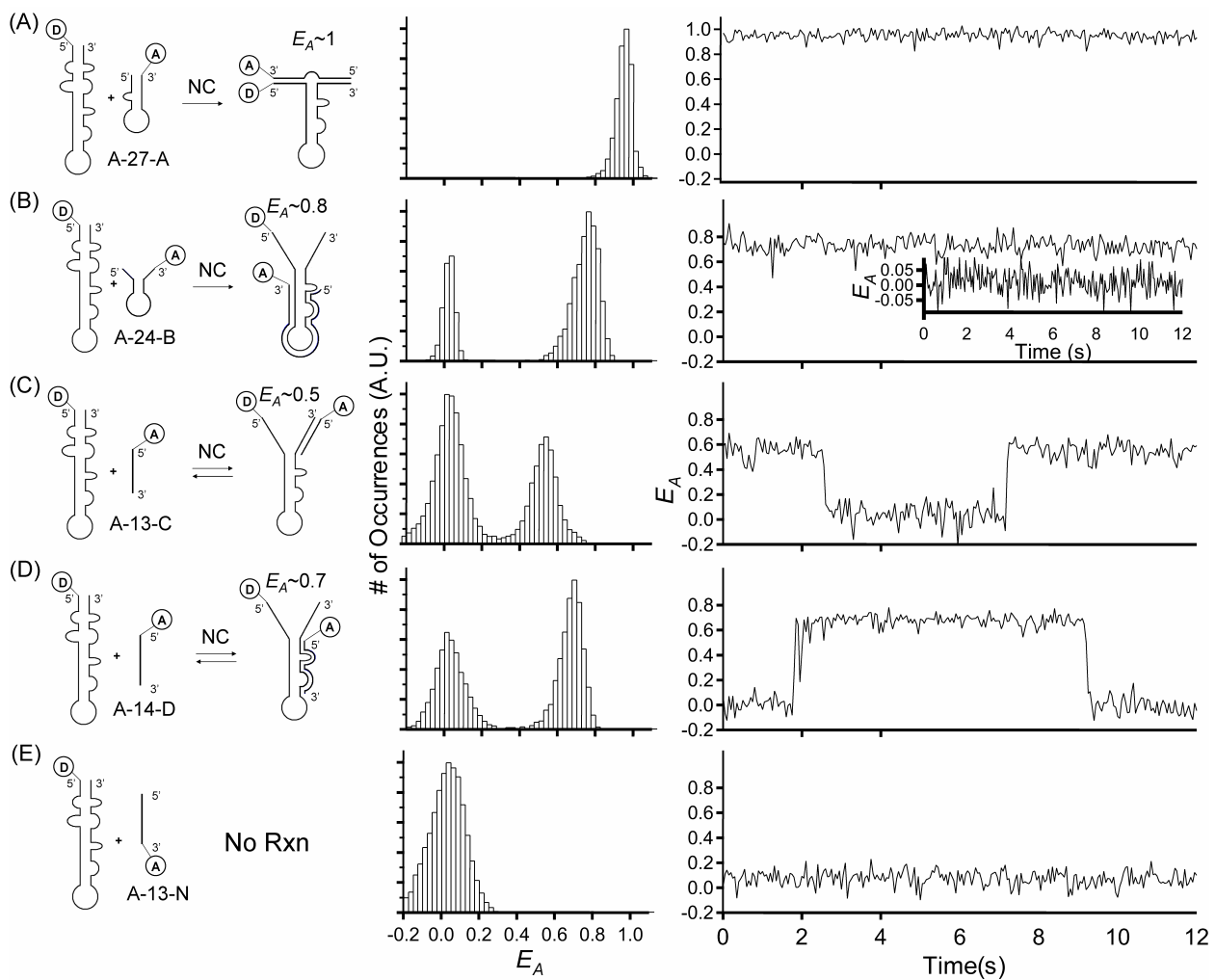


FIGURE 3

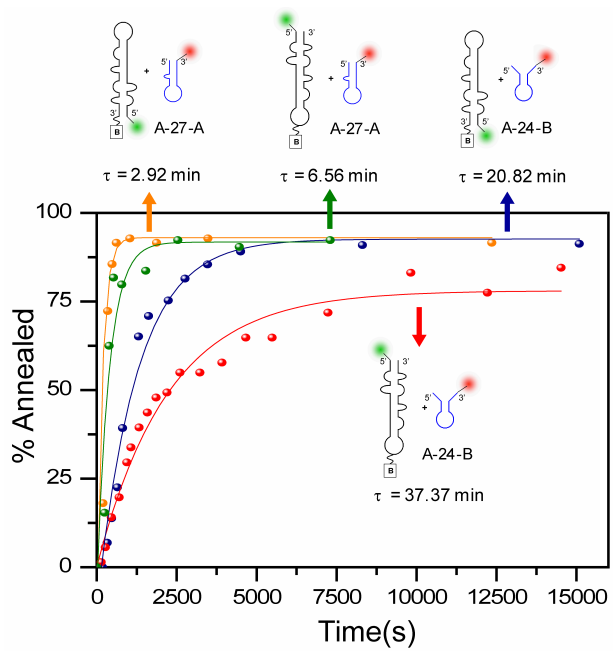


FIGURE 4

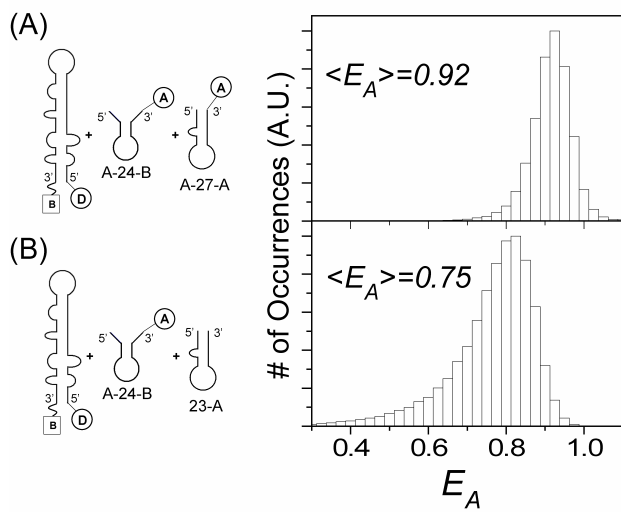


FIGURE 5