Interaction of the HIV-1 frameshift signal with the ribosome

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Received March 14, 2009; Revised September 1, 2009; Accepted September 2, 2009

ABSTRACT

Ribosomal frameshifting on viral RNAs relies on the mechanical properties of structural elements, often pseudoknots and more rarely stem-loops, that are unfolded by the ribosome during translation. In human immunodeficiency virus (HIV)-1 type B a long hairpin containing a three-nucleotide bulge is responsible for efficient frameshifting. This three-nucleotide bulge separates the hairpin in two domains: an unstable lower stem followed by a GC-rich upper stem. Toeprinting and chemical probing assays suggest that a hairpin-like structure is retained when ribosomes, initially bound at the slippery sequence, were allowed multiple EF-G catalyzed translocation cycles. However, while the upper stem remains intact the lower stem readily melts. After the first, and single step of translocation of deacylated tRNA to the 30 S P site, movement of the mRNA stem-loop in the 5' direction is halted, which is consistent with the notion that the downstream secondary structure resists unfolding. Mechanical stretching of the hairpin using optical tweezers only allows clear identification of unfolding of the upper stem at a force of 12.8 ± 1.0 pN. This suggests that the lower stem is unstable and may indeed readily unfold in the presence of a translocating ribosome.

INTRODUCTION

The ribosome translocates mRNA and bound tRNA molecules accurately in order to maintain the reading frame. This process results in movement of the ribosome along the mRNA by three nucleotides toward the mRNA’s 3’-end. Translocation of mRNA and tRNAs is a property of the ribosome itself (1,2), however binding of elongation factor G (EF-G) and subsequent hydrolysis of GTP strongly catalyzes it (3). Although the ribosome acts as its own helicase, stable folded structures within the coding regions of mRNA affect the rate of translocation, and more seriously, may trigger a change of reading frame (4,5). Such frameshifting mRNA elements play a crucial role in the translational control of viral proteins via –1 programmed ribosomal frameshifting (–1 PRF) where the reading frame has shifted by one base toward the mRNA 5’-end. The –1 PRF requires both the mRNA slippery sequence at the ribosome coding sites as well as a downstream structural element that resists unfolding, representing a physical barrier to the mRNA translocation machinery. While in many cases the downstream barrier constitutes a hairpin (H)-type pseudoknots (6,7), on a rare occasion it can also be a simple stem–loop structure (8–11). It is of interest to note that although these pseudoknots or stem–loops also trigger ribosomal pausing at the slippery site, consistent with notion of them acting as physical barriers, the extent of pausing shows no correlation with frameshift efficiency (12).

The crystal structure of the ribosome in complex with mRNA has revealed that the mRNA is in a single-stranded conformation in the narrow downstream tunnel (13,14). The ribosome therefore has to unwind mRNA secondary structure through its mRNA helicase activity (15,16). A mechanistic basis for mRNA helicase activity has been proposed involving ribosomal proteins S3, S4, S5 at the mRNA entrance and rotational movement of the head of the 30 S subunit (13,15). The 9 Å and the torsional restraint models (17) propose that –1 PRF is dependent on the mechanical tension induced when a pseudoknot resists unfolding by a moving translating ribosome. Possible effects of such tension were directly observed in the cryo-electron microscopic (Cryo-EM) images.
of eukaryotic ribosomes stalled in the process of –1 frameshifting in complex with eEF2, tRNA and a frameshifting mRNA pseudoknot (18). The opposing actions of translocation catalyzed by eEF2 and resistance to unfolding by the mRNA strand generate strain that deforms the P-site tRNA which may weaken the codon–anticodon interaction and promote the shift by one nucleotide into the 5′ direction.

The notion that mechanical stability of mRNA structural element is crucial for –1 PRF, has triggered mechanical unfolding experiments of individual mRNA pseudoknot, mutants as well as some of the constituent hairpins using optical tweezers hairpins (19–21). Some of these experiments suggest that frameshift efficiencies correlate with unfolding forces rather than the free-energy difference between the folded and unfolded state (21). This indicates that –1 PRF is kinetically controlled, as has been proposed previously (22).

In HIV-1, translational frameshifting leads to synthesis of the Gag–Pol fusion protein which gives rise to the viral protease, reverse transcriptase (RT) and integrase. This HIV-1 RNA frameshifting signal is a potential target for antiviral therapy (23–26). The exact structure of this RNA frameshifting signal has been the subject of debate. Jacks and collaborators initially proposed that it is a stem–loop structure downstream of the slippery region and the lower stem are much less so, and may relate with unfolding forces rather than the free-energy difference between the folded and unfolded state (21). This indicates that –1 PRF is kinetically controlled, as has been proposed previously (22).

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Questions, however, remain about the structure of the HIV-1 frameshift signal when ribosomes are present and bound at or downstream of the slippery sequence. Here, using Escherichia coli ribosomes, we address this question and probe possible interactions of the HIV-1 frameshift signal with the ribosome by using toeprinting and chemical probing assays. The ability of this eukaryotic mRNA frameshifting signals to promote –1 PRF in the prokaryotic translational machinery has been previously demonstrated (33,34).

**MATERIALS AND METHODS**

**mRNA translocation**

The translocation of mRNA was assayed by toeprinting as described (35,36). mRNA (1 μM) was annealed to primer (2 μM) in 50 mM K-Hepes (pH 7.0) and 100 mM KCl by heating to 90°C for 1 min and placing at room temperature until the temperature reached 45°C. To form the complexes, tight-couple ribosomes (2–5 μM) (37) from *E. coli* MRE600 were added to 0.6 μM of mRNA in 60 mM NH₄Cl, 10 mM Tris–Acetate (pH 7.4) and 20 mM MgCl₂ and incubated at 37°C for 10 min. A first mRNA (4 μM) was added to fill the P site by incubation at 37°C for 10 min and aliquot (0.6 pmol mRNA) was removed to ice for later extension. A second tRNA (4 μM) was added to fill the A site by incubation at 37°C for 10 min and aliquot (0.6 pmol mRNA) was removed to ice for later extension. EF-G was added in buffer (50 mM Tris–HCl (pH 7.6), 20 mM MgCl₂, 100 mM NH₄Cl and 1 mM DTT, 1.5 mM GTP) such that the final concentrations of GTP and EF-G were 300 and 1 μM, respectively. Reactions were incubated at 37°C for 10 min, and aliquots (0.6 pmol mRNA) were removed from each reaction lacking (−G) or containing (+G) EF-G. Each of the aliquots was then extended in parallel with (5′-CTTTATCTTCAAGAGAAAACC-3′) primer, and the product were resolved by 8% denaturing PAGE.

**Stepwise translocation**

Stepwise translocation experiments were done as previously described (15). Tight-couple 70S ribosomes (1 μM final concentration) from *E. coli* MRE600 (39) were incubated with mSP-HIV-1 mRNA (1 μM) in 30 μl binding buffer (10 mM Tris–HCl (pH 7.4), 60 mM NH₄Cl, 10 mM Mg(OAc)₂, 1 μM β-ME) for 10 min at 37°C, followed by addition of tRNAPho (1 μM) and a further 10 min incubation to fill the P site. Aliquots (5 μl) of this reaction were then added to separate tubes containing either GTP (600 μM) (F), GTP + tRNALeu (1 μM) (L), GTP + tRNALeu, EF-G (1 μM) (L’), GTP + tRNALeu,
EF-G + tRNA\textsubscript{Gly} (1 μM) (G), GTP + tRNA\textsubscript{Leu}, EF-G + tRNA\textsubscript{Glu} + tRNA\textsubscript{lys} (1 μM) (K), GTP + tRNA\textsubscript{Leu} + tRNA\textsubscript{Glu} + tRNA\textsubscript{lys} (K'), in binding buffer (final volume, 10 μl). These tubes were incubated for 10 min at 37°C and then placed on ice for the primer extension in toeprinting analysis (4 μl).

Footprints
To toeprint, binding of mRNAs was performed by incubating 70S ribosome (in the range of 50–750 pmol according to the mRNA tested) with mRNA (10–50 pmol) in 50 μl reaction buffer A (20 mM MgCl\textsubscript{2}, 150 mM NH\textsubscript{4}Cl, 80 mM potassium cacodylate, pH 7.2) at 37°C for 10 min. A first tRNA (320 pmol) was added to fill the P site by incubation at 37°C for 10 min. A second tRNA (320 pmol) was added to fill the A site by incubation at 37°C for 10 min. mRNA–tRNA–70S complexes were then purified by ultrafiltration (MICROCON YM-100 100 000 Da, Fisher scientific LABOSI). The ternary complex was diluted in 250 μl of buffer A and distributed into 50 μl aliquots (2 pmol mRNA). Chemical probing (38) was performed by addition of 2, 4 or 8 μl dimethyl sulfate (DMS; 1:10 dilution in 95% ethanol), 2, 4 or 8 μl kethoxal (KE; 19 mg/ml in H\textsubscript{2}O) on 50 μl aliquot, followed by incubation at 37°C for 10 min. All modification reactions were stopped by addition of 150 μl 95% ethanol and 5 μl 3 M sodium acetate followed by rapid mixing. KE-modified samples were adjusted to 25 mM potassium borate (pH 7.0). The pellets were resuspended in 200 μl of 0.3 M sodium acetate, 2.5 mM EDTA and 0.5% SDS (with addition of 25 mM potassium borate for KE samples), extracted three times with phenol, twice with chloroform and resuspended in 10 μl H\textsubscript{2}O for (DMS samples) or in 10 μl 25 mM potassium borate (for KE samples). Primer extension reactions were performed as described (38).

Purified tRNA\textsubscript{lys}, tRNA\textsubscript{Met} and tRNA\textsubscript{phe} were purchased from Sigma, tRNA\textsubscript{Gly} and tRNA\textsubscript{Leu} were gracefully donated by Henry Grosjean.

Messenger RNAs were prepared by in vitro transcription. Plasmid pGENE32 is pUC118 containing a region of phase T4 gene 32 (40) from nucleotide position –54 to +84 (where +1 is the translational start) downstream of an engineered T7 promoter sequence. The introduction of slippery sequence in pGENE32 was performed by site-directed mutagenesis kit (Stratagene). Transcripts with stem–loop were obtained by in vitro transcription of synthetic genes flanked upstream by T7 RNA polymerase promoter region and downstream by a BamHI restriction site. The synthetic genes were constructed by shotgun ligation of 10 DNA fragments (24–30 mers) covering both strands and ligated in the KpnI and BamHI sites of pGENE32. All transcripts have been purified by denaturing PAGE.

His-tagged EF-G was purified from pET24b-fusA in E. coli BL21(DE3) as described (41).

Mechanical unfolding using optical tweezers
RNA was synthesized from a template obtained by polymerase chain reaction (PCR) from bases 3821 to 628 of the pBR322 DNA plasmid, where the frameshifting RNA signal from HIV-1 was cloned into the EcoRI and HindIII restriction sites and a T7 promoter was appended to the template in the course of the PCR reaction (42). The DNA components of the handles were prepared by PCR from pBR322. Handle A (pBR322 bases 3821 to 3) was biotinylated, and one of the primers used to amplify handle B (pBR322 bases 30 to 628) was purchased with a 5’ digoxigenin group.

RNA and DNA handles were resuspended in 10 mM sodium phosphate buffer (pH 6.4), and incubated at a ratio of ~1:1:1 at 90°C for 1 min and transferred to room temperature to cool down gradually, and subsequently diluted to a final concentration (of RNA) of ~1 μM.

Five microliters of anti-digoxigenin-coated polystyrene beads (0.3 nm; diam. 0.49 μm) were mixed with 1 μl of the DNA–RNA hybrid (~1 μM) in binding buffer (10 mM Tris buffer [pH 7.0], 250 mM NaCl, 10 mM MgCl\textsubscript{2}, 0.4% w/v BSA), and incubated at 4°C for overnight on a rotator.

Sample cells were preassembled prior to use. Two thin strip spacers (thickness ~200 μm) were positioned ~5 mm apart on the center of a pre-cleaned microscope slide, and epoxy applied at the outer edges of the spacers. A streptavidin functionalized 24 × 40 mm, no. 1.5 coverglass (Xenopore) was then put on the top of the slide. Before introducing the bead and RNA mixture, the sample cell was surface-coated with acetylated BSA by incubation with binding buffer for 30 min at room temperature and washed with 1 ml of the binding buffer, to prevent any sticking of beads to the surface. The bead and RNA mixture was then introduced into the sample cell, and incubated for 30 min at room temperature, and finally washed with 1 ml of binding buffer to remove any unbound beads and RNA.

Molecules were stretched in 10 mM Tris buffer (pH 7.0), 250 mM NaCl, 10 mM MgCl\textsubscript{2}, or alternatively in 10 mM Tris-acetate (pH 7.4), 60 mM NH\textsubscript{4}Cl, 6 mM β-mercapto, 20 mM MgCl\textsubscript{2}. Unfolding/refolding parameters were statistically indistinguishable for both conditions. The spring constant of the optical tweezers was 0.1–0.2 pN nm\textsuperscript{−1}.

The extension of the unfolded single stranded HIV-1 hairpin, \(x_{SS}F\) was computed as \(x_{SS}(F) = \Delta x + L_{HF}\) for convenience, with \(F = (F_1 + F_2)/2\) the unfolding force and \(\Delta x\) the increase in extension (Figure S2). The increase in contour length (expressed in number of nucleotides) was subsequently computed using the worm-like chain model for polymer elasticity assuming a stretching modulus of 1000 pN, a persistence length of 1 nm (43,44) and inter-phosphate distance of 0.59 nm. Alternatively, one may choose to do the computation explicitly, where it is found that \(x_{SS}(F_2) = \Delta x - (x_{HF}(F_2) - x_{HF}(F_1)) + L_{HF}\). Then however, one needs to fit the section of the force versus extension curve up to the unfolding event with a worm-like chain model in order to compute the extension of the handles, \(x_{HF}(F_2)\) at \(F_2\). When performing this more laborious analysis on a subset of our data, we only find a 1–2 nucleotide difference in contour length compared to the analysis that utilizes the applied approximation \(F = (F_1 + F_2)/2\). Standard free-energies at zero force were computed for each trajectory according to \(\Delta G = F\Delta x - f_{0}^{SS} F dx\) using the worm-like chain model.
with persistence length of 1 nm and stretching modulus of 1000 pN (43).

RESULTS

We used a toeprinting assay to monitor the position and/or structure of the HIV-1 hairpin during the movement of the ribosome along mRNA. This primer-extension inhibition assay has been shown to be a powerful tool for mapping the position of mRNA within 30S and 70S ribosomal complexes that contain tRNA (45). When RT encounters the ribosome (a so-called hard-stop), it terminates cDNA synthesis thereby generating a highly specific toeprint. Alternatively, RT may also stall at mRNA structural elements further downstream from the ribosome that prove too hard for RT to unwind, resulting in what is generally called an extended toeprint.

Translocation on the HIV-1 slippery sequence

We first demonstrated that under simple experimental conditions (where tRNA was non-enzymatically delivered at the ribosomal A site), toeprinting allows localization of the ribosome on the mRNA containing the wild-type HIV-1 slippery sequence (mSP-HIV-1) but not the downstream hairpin. The sequences and secondary structures of the mRNA constructs used are derived from T4 gene 32 mRNA in which we introduced the HIV-1 frameshifting sequences and secondary structures of the HIV-1 slippery sequence (mSP-HIV-1) but not the downstream hairpin. The sequences and secondary structures of the HIV-1 slippery sequence (mSP-HIV-1) but not the downstream hairpin.

Figure 1. Secondary structure of the mRNAs used in this study. The tow domains of the HIV-1 hairpin are labeled: lower stem (FSL) and upper stem (FSU). The slippery sequence is underlined and the Shine-Dalgarno sequence is boxed. The first nucleotide of codon 1 (position +1) is labeled. Toeprints that are observed when each codon is positioned in the ribosomal P-site are shown on the top. Nucleotides that were mutated are indicated in a box for the mSP-SL-CCC-HIV-1 RNA. The limit of accessibility (position +11) of a RNA double helix at ribosomal surface (15) is highlighted by a triangle. The spacer sequence that is expected to be single-stranded in the mRNA tunnel is indicated by a dashed line.

Extended toeprints with the downstream frameshifting HIV-1 hairpin

We then applied this assay to a gene 32 message (mSP-SL-HIV-1), which contains the mRNA frameshifting signal (Figure 2A). In order to avoid any unnecessary ambiguity in interpreting the toeprints, we substituted the phenylalanine codon (u+1u+2u+3) for a methionine codon (a+1a+2g+3) (mSP-SL-HIV-1; Figure 1) to avoid complications due to the presence of the slippery site. This allows tRNA binding in only a single unique reading frame.

When tRNA^{Met} was bound at the P site, a doublet of toeprints at positions +16 and +17 appears when tRNA^{Leu} subsequently binds at the A site (Figure S1A, lane F). This is thought to be due to a conformational change in the ribosomal complex following A-site binding so that a single position of the tRNA in the A site results in a doublet of bands (at positions +16 and +17) (46,47). It is important to note that the signal at +15 did increase, indicating that binding of the tRNA^{Leu} at the A site assisted in the positioning of the mRNA with tRNA^{Phe} preferentially bound to the u+1u+2u+3 codon at the A site assisted in the positioning of the mRNA with tRNA^{Phe} preferentially bound to the u+1u+2u+3 codon. A third toeprint at +17 appears when tRNA^{Leu} subsequently binds at the A site (Figure S1A, lane F) triggered translocation leading to a toeprint at +19.

Extended toeprints with the downstream frameshifting HIV-1 hairpin

When tRNA^{Met} was bound at the P site, a doublet of toeprints at positions +16/17 was detected (Figures S2 and 2A). We note the existence of a stop at position +17 in the control lane (in the absence of ribosomes). However, the signal in presence of ribosome is substantially lower. The ribosome (in the absence of ribosomes). However, the signal in presence of ribosome is substantially lower. The ribosome
produced by ribosomal complexes with mRNAs mSP-Tyr-SL-HIV-1. Complexes with mRNAs mSP-SL-CCC-HIV-1. (C) Toeprints produced by ribosomal complexes with mRNAs mSP-SL-CCC-HIV-1. (C) Toeprints produced by ribosomal complexes with mRNAs mSP-Tyr-SL-HIV-1.

can extend beyond the usual 15 nucleotides buried in the ribosome-mRNA track (48,50). Unlike hard stops, extended toeprints are generally dependent upon change of temperature, RT concentration, or the source of RT. Therefore, we varied the temperature and also tested different RTs to look for extended toeprints in the case of mSP-SL-HIV-1. Two additional reverse transcription stops were identified within the 3' region of the stem–loop (48,50). Unlike hard stops, extended toeprints can extend beyond the usual 15 nucleotides buried in the ribosome-mRNA track (48,50).

Figure 2, Toeprinting analysis of successive EF-G catalyzed translocation steps. (A) mSP-SL-HIV-1, lanes are labeled according to the tRNA species bound to the ribosomal P-site at each step. M, tRNA^Met was bound to the mRNA–ribosome complex followed by stepwise addition of GTP + tRNA^Leu (M'), GTP + tRNA^Leu, EF-G (L), GTP + tRNA^Leu, EF-G + tRNA^Gly (G), GTP + tRNA^Leu, EF-G + tRNA^Gly + tRNA^Lys (K). (B) Toeprints produced by ribosomal complexes with mRNAs mSP-SL-CCC-HIV-1. (C) Toeprints produced by ribosomal complexes with mRNAs mSP-Tyr-SL-HIV-1.

Translocation with the downstream frameshifting HIV-1 stem–loop

We subsequently monitored the position of mSP-SL-HIV-1 within the ribosome allowing a single or multiple rounds of EF-G catalyzed translocation. The signal at positions +43 and +47 disappeared to give a new toeprint at +39 (Figure 2, lane L, and 3B), indicating that the lower stem readily gave way upon just a single translocation step. An identical result was observed in the mSP-SL-HIV-1 RNA which has a wild-type slippery sequence (first codon is u +1t +2u +3) indicating that codon substitution to a methionine codon did not affect the observed toeprints except for the +47 signal that is ambiguous (Figure S1B). Furthermore, upon a single translocation, the +16/+17 toeprint signals disappeared without, however, any occurrence of a new toeprint at position +19 (lane L). The disappearance of the mSP-SL-HIV-1 +16/+17 toeprints upon addition of EF-G is intriguing. We also observed the same phenomenon with the construct containing the wild-type slippery sequence (Figure S1B), and interestingly also with an mRNA containing a pseudoknot (BWYV) bound to ribosome (51). This phenomenon is characteristic to mRNAs containing downstream structural elements, however its cause still need to be elucidated. Ribosomes are known to change conformation upon EF-G binding (52–55). One may speculate that this either blocks RT access directly or stabilizes the mRNA structure.

Any subsequent addition of tRNA^Gly (lane G) and tRNA^Lys (lane K) in presence of EF-G did not further affect the position of the extended toeprint, which remained at +39, indicating that further movement of the mRNA through the ribosome was impaired.

We next tested an mRNA with an additional codon between the AUG and the start of the lower stem (mSP-Tyr-SL-HIV-1 RNA) extending the spacer, which should at least allow for one round of translocation to be visualized before the ribosome stalls. The toeprints at the new positions A+16/U+17 were unambiguously identified (Figure S3), which places the boundary region contacting the ribosome upstream from the bulge region. These bases are in the upper stem for the mSP-SL-HIV-1 RNA. In Figure 2C, where experimental conditions were tuned as to specifically detect the extended toeprints, the toeprint signals in the region +17 prove somewhat weaker than in Figure S3, but are always present. As expected, with the longer spacer, a band at +19 appeared upon addition of EF-G (Figure 2C, lane L). Interestingly, the extended toeprints did not change and appeared at the exact same positions at A+46 and U+50 (corresponding to the A+43 and U+47 in mSP-SL-HIV-1 RNA) (Figure 2C). Subsequent addition of EF-G and tRNA^Tyr and tRNA^Gly produced the same changes in the toeprint pattern as for the mSP-SL-HIV-1 RNA (a shift of 4-nt from +46 to +42). We note that in this case the +19 toeprint remains the strongest toeprint in the upper region of the gel (by comparison with the +17 toeprint) (Figure 2C, lanes Y and G) indicating that translocation, as before, seems to be impaired with the downstream upper stem.

Chemical probing of mRNA with the downstream frameshifting HIV-1 stem–loop

Toeprinting assays fail to provide information on more detailed structural changes that the mRNA might
undergo upon binding to the ribosome. Therefore, a footprint assay was performed to obtain structural information upon the interaction of mSP-SL-HIV-1 mRNA with 70S ribosome–tRNA complexes. In these complexes the ribosome is positioned with the slippery sequence at the decoding site. Results of the chemical probing with KE, 1-cyclohexyl-2-morpholino-carbo-diimide-b-metho-p-toluene sulfonate (CMCT) and DMS are shown in Figure 4. For free mSP-SL-HIV-1 mRNA the chemical modification patterns and levels of reactivities are identical to those previously published (9). Nucleotides in the apical ACAA teraloop as well as A+43 and A+44 from the bulge are found to be reactive to DMS (Figure 4), while nucleotides G+41 and G+42 were reactive to KE. In the lower stem most of the guanine, adenine and uracil bases are accessible to the chemical probes indicating poor stability (9).

Subsequently, mSP-SL-HIV-1 mRNA was probed in complex with the ribosome and tRNA. For each chemical probe tested, bands that were present in a control lane of unmodified mRNA incubated with ribosome were not taken into account. In the complex, the characteristic strong protections at the guanine nucleotides in the Shine–Dalgarno sequence (G–11, G–10) are clearly seen. Base A+1, of the methionine codon is protected from chemical modifications demonstrating, as previously described (56), Watson–Crick pairing with the anticodon of tRNA\textsuperscript{Met}. Interestingly, nucleotide A+6 experienced an increase in reactivity similar to what has previously been seen for a mRNA containing a hairpin (selenocysteine incorporation sequence SECIS) in complex with 30S subunit and tRNA\textsuperscript{Met} (57). In the spacer and hairpin regions, most of the changes in chemical reactivities were detected in the lower stem that is supposed to be close to the ribosomal surface. Bases G+7, G+8 and G+12 were protected from modification by KE (Figure 4). The reactivity of G+9 toward KE slightly increased in a way similar as A+10 toward DMS.

**Mechanical unfolding of the HIV-1 RNA hairpin**

Since toeprinting experiments suggested reduced structural stability of the lower stem, we investigated the
mechanical stability of the HIV-1 hairpins. Individual hairpins, sandwiched between two differentially end-labeled hybrid DNA–RNA handles were unfolded using optical tweezers (42). Molecules were tethered between a streptavidin-coated glass cover slip and anti-digoxygenin coated polystyrene beads (diameter 1 μm), and stretched by moving the piezo-actuated microscope stage while holding the bead with optical tweezers. Force versus extension curves were computed taking into account this experimental geometry (58). Stage velocities were 67 nm/s, so that the system remained at or close to thermodynamic equilibrium as confirmed by the overlap of the stretching and relaxation force versus extension curves (Figure 5B). In addition, repeated unfolding and folding can be observed within single force versus extension curves (Figure 5B), a further indication that the loading rate is sufficiently low as to assure thermodynamic equilibrium. The increase in contour length (expressed in number of nucleotides) upon unfolding was subsequently computed using the worm-like chain model for polymer elasticity assuming a stretching modulus of 1000 pN, a persistence length of 1 nm (43,59) and a inter-phosphate distance of 0.59 nm (see ‘Materials and Methods’ section). Results are summarized in Figure 5C, yielding a mean increase of 25.3 ± 3.4 nm (mean ± SD) nucleotides, consistent with the contour length of the upper stem of the HIV-1 hairpin. The mean unfolding force is 12.8 ± 1.0 pN (mean ± SD), whereas the standard free-energy change at zero force was found as ΔG = 10 ± 2 kcal/mol (mean ± SD). Refolding statistics are summarized in the Figures S4 and S5 and yield a decrease of contour length of 26.0 ± 4.7 nm (mean ± SD) at an average refolding force of 13.1 ± 1.1 pN (mean ± SD), and ΔG = 11 ± 3 kcal/mol (mean ± SD), essentially unchanged from the unfolding statistics, as one would expect when at thermal equilibrium.

**DISCUSSION**

Extended toeprints (albeit weak ones) at +43 and +47 in the pre-translocation state indicate that a fraction of RTs was incapable of unwinding part of the lower stem and

![Figure 4](https://academic.oup.com/nar/article-abstract/37/22/7654/2409901)
bulge region when encountering the HIV-1 hairpin bound at the ribosome. Since such toeprints do not occur with free hairpins, contacts with the ribosome (34) may be responsible for these signals. Extending the spacer sequence by an extra codon yields an identical toeprint; not surprising as it is hard to imagine that a longer spacer would stand in the way of forming ribosomal contacts. This will be discussed further below. Furthermore, we showed that these toeprints are dependent on the structure of the bulged region of the frameshifting signal. The subsequent replacement of the GGA-bulge with a CCC-bulge, known to reduce –1 PRF efficiency (8,32), practically erases the +43 and +47 toeprints, indicating such a mutation affects either contacts with the ribosomes or stability of the hairpin itself. We tend to favor the former possibility as mechanical unfolding of free hairpins indicates the lower stem is fairly weak to begin with. Certainly it remains possible that contact with the ribosome can stabilize part of the lower stem. If such stabilization was to occur, it does not prevent further translocation as a single cycle of EF-G catalyzed translocation moves the toeprint to +39 providing direct evidence for melting of the lower stem. Interestingly, the extended toeprint at +39 remains even under conditions where further translocation is allowed. This suggests that the upper stem is capable of inhibiting translocation of a sizeable fraction of ribosomes. Our mechanical unfolding study supports the notion of a weak lower stem since no transition in the force versus extension curves indicative of unfolding of solely the lower stem have been observed. Although on occasion we suspect such a transition (by visual inspection of the force versus extension data, typically at forces ~6pN and lower), the increase in extension in those cases should be ~4–5 nm (by theory), too small to reliably distinguish from thermal fluctuations at those force levels. We note

Figure 5. Mechanical unfolding of the HIV-1 hairpin. (A) HIV-1 hairpin sequence and structure. (B) Typical force versus extension curves upon stretching and relaxation of a single molecule using optical tweezers. The relaxation curve has been offset by 50 nm for clarity, since both curves would overlap otherwise. (C) Histogram of contour lengths of unfolded structures. (D) Histogram of unfolding forces.
that were these transitions to occur in the deeper part of the force versus extension curve \((F > 6 \text{ pN})\), we should be able to detect a shift of the curve toward the right consistent with unfolding of the lower stem. However, the increases in contour lengths (Figure 5C) observed are consistent with unfolding of solely the upper stem. Not on any occasions have we observed increases in contour lengths consistent with simultaneous unfolding of the lower and upper stems. Since unfolding is hierarchical, the lower stem has to unfold before the upper stem does. We conclude that the lower stem therefore has much reduced mechanical stability compared to the upper stem. Quantitative assessment of the stability of the lower stem is currently not possible in the existing experimental geometry, and requires higher resolution experimental designs (60), if at all possible. The question naturally arises how these findings accord with the observed +43 and +47 toeprint and the dependence of frameshifting upon the lower stem and bulge regions. It seems unlikely that tensions as low as a mere 6 pN and below, can trigger a frameshift (61). Therefore, it is appealing to consider that unfolding of the hairpin in the presence of ribosomes will differ significantly from that when ribosomes are absent. When adhering to a tension-dependent mechanism of \(-1\) frameshifting this suggest that contacts with the ribosome’s exterior surface may indeed stabilize the part of the lower stem of the hairpin. Single-molecule mechanical unfolding experiments in the presence of ribosomes are considered, but beyond the scope of this article.

Our chemical probing experiments provide further insight into the structure of the HIV-1 frameshift signal when bound to the ribosome. The chemical reactivity pattern indicates that almost the entire structure of the hairpin is maintained. Nucleotides from the ACAA tetraloop and the GGA bulge region remained reactive to chemical probes whereas nucleotides in the upper stem were unreactive. As expected, most of the changes in chemical reactivity of the bases are concentrated in the lower stem. The changes observed in the spacer at positions +7, +8 are consistent with this segment of the mRNA being engaged in the 30S mRNA tunnel (13). Further downstream in the spacer sequence, nucleotides \(G_{+12}, A_{+16}\) and \(G_{+19}\) that are expected to be located in the vicinity or at the ribosomal helicase center indeed experienced a change in reactivity. In this segment of the RNA, the DMS reactivity of nucleotide \(A_{+11}\) remained mostly unaffected by the presence of the ribosome. Unfortunately, our chemical probing experiments could not provide useful information on the 3’ strand of the lower stem. In summary, the data demonstrate that the 5’ region of the lower stem experiences structural changes when contacted by the ribosome.

A number of other observations, although of no direct consequence for the interpretation of the extended toeprints, are of interest. First is the observation of the occurrence of an apparent 4-nt translocation step during transfer of tRNA\(^{\text{Phe}}\) from A site to P site. The disappearance and appearance of respectively the +43 and +39 toeprints indicate that the addition of EF-G triggered a movement of the mSP-HIV-1 RNA inside the ribosome of four nucleotides. However, we note that this 4-nt shift is detected on the 3’ strand of the mRNA hairpin that is located outside the ribosome and therefore may not necessarily present a 1:1 reflection of what happened at the coding site. On the other hand, single translocation cycles with only the slippery sequence (mSP-HIV-1) also show a 4-nt step. In this experiment we analyzed the toeprint signals that result from A site, P site tRNA binding and subsequent EF-G catalyzed translocation. The mSP-HIV-1 toeprint at +15 corresponds to tRNA\(^{\text{Phe}}\) that recognizes the overlapping codon, one nucleotide upstream (\(u_{+1}u_{+2}u_{+3}\)). Binding of tRNA\(^{\text{Phe}}\) to the \(u_{+1}u_{+2}u_{+3}\) codon results in an optimal spacing of seven bases between the SD sequence and the start codon (62). The toeprint at +16 corresponds to tRNA\(^{\text{Phe}}\) paired to codon 1 (\(u_{+1}u_{+2}u_{+3}\)). Interestingly, overlapping codons \(u_{+2}u_{+3}u_{+4}\) and \(u_{+3}u_{+4}u_{+5}\) are not pairing with tRNA\(^{\text{Phe}}\) because no toeprints were observed at +17 and +18. This is most likely due to unfavorable spacing between the P-site codon and the Shine-Dalgarno sequence (47,63). After P-site filling and the first round of translocation, the toeprint at +19 is the expected signal for a post-translocation complex with tRNA\(^{\text{Leu}}\) in the P site bound at the \(u_{+4}u_{+5}u_{+6}\) codon. However, the major toeprint signal is at position +20 suggesting that in the post-translocation complex the P site-bound tRNA\(^{\text{Leu}}\) is interacting with the \(u_{+5}u_{+6}u_{+7}\) codon. This corresponds to a 4-nt translocation event. Subsequent translocation cycles seem to force the toeprints back into register. It remains unclear what gives cause to this behavior. We note that a 4-nt translocation has been observed for punctuated mRNA with an extra, unpaired nucleotide between codons (64). Interestingly, Leger and collaborators proposed that slippage and repairing of P-site-bound tRNA\(^{\text{Phe}}\) in the \(-1\) frame would leave an unpaired nucleotide between the Phe and Leu codons (32). But this is unlikely at equilibrium in our assay because A-site binding of tRNA\(^{\text{Leu}}\) mostly repositioned tRNA\(^{\text{Phe}}\) into the canonical frame removing any unpaired nucleotide between Phe and Leu codons.

The P-site tRNA binding triggered a major RT stop at position +17 while only a weak signal at position +16 (the ‘classical toeprint’). Previous toeprinting assays under identical conditions but with different phage T4 mRNAs showed a variety of toeprints signals confined to the +14 to +17 range (45). Thus toeprinting assays are very sensitive to the type of mRNA tested and we note that the toeprint signals with the mSP-HIV-1 and mSP-SL-HIV-1 mRNAs respectively at +16 and +17 fall into this range. Could the observed difference between the two mRNAs be attributed to the hairpin? With unstructured mRNA, we expect a pulling force on the spacer mRNA out of the ribosome purely based on entropic arguments. Such a force presumably exposes base +16 giving rise to the classical toeprint. In the case of the HIV-1 hairpin, physical interactions of the ribosome with the hairpin may provide some sort of strain relieve allowing +16 to slightly relax back into the ribosome. On the other hand, how could such a mechanism be consistent with +16/+17 toeprint obtained when an additional codon was inserted in the spacer, between the AUG codon and the hairpin? If the
hairpin provides strain relieve would one not expect the mRNA to recede further back into ribosome, not only protect +16 but also perhaps +17. However, lacking any quantitative information about tension in the spacer, such extra movement may be too small to also protect +17. These are intriguing possibilities and raise questions that require the design of new experiments, well beyond the scope of this work, for answering.

Site-directed mutagenesis and amino-acid sequencing localized the site of frameshifting to the UUA codon of the HIV-1 slippery sequence (65). The presence of the GC-rich upper stem strongly enhances frameshifting efficiencies. This level of mRNA slippage is enhanced further by the bulge and lower stem. We show here that when the slippery sequence is bound at the decoding site, the ribosome directly influences the spacer sequence in the lower stem that very likely enters the mRNA track. In addition, extended toeprints localized in the upper segment of the lower stem support a physical interaction between this region and the ribosome. These results are in agreement with the importance of the lower stem and bulge regions for the –1 frameshift (8,32,66). It is conceivable that this interaction enhances –1 frameshifting at the particularly slippery UUUUUA sequence (67).

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

ACKNOWLEDGEMENTS

The authors wish to thank Kevin Wilson for the kind gift of the pET24b-fusA vector for His-tagged EF-G.

FUNDING

Young Investigator Award from the Human Frontier Science Program (HFSP) to D.F. and K.V. Funding for open access charge: CNRS

Conflict of interest statement. None declared.

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