Mutations in the $T\Psi C$ Loop of \textit{E. coli} tRNA$^{\text{Lys,3}}$ Have Varied Effects on \textit{In Trans} Complementation of HIV-1 Replication

Wanfeng Yu, Anna McCulley and Casey D Morrow*

Address: Department of Cell Biology, University of Alabama at Birmingham, Birmingham, AL 35294-0024, USA

Email: Wanfeng Yu - wanfengy@uab.edu; Anna McCulley - annam@uab.edu; Casey D Morrow* - caseym@uab.edu

* Corresponding author

Abstract

\textbf{Background:} Human immunodeficiency virus (HIV-1) exclusively selects and utilizes tRNA$^{\text{Lys,3}}$ as the primer for initiation of reverse transcription. Several elements within the $T\Psi C$ stem loop of tRNA$^{\text{Lys,3}}$ are postulated to be important for selection and use in reverse transcription. The post-transcriptional modification at nucleotide 58 could play a role during plus-strand synthesis to stop reverse transcriptase from re-copying the tRNA primer. Nucleotides 53 and 54 within the $T\Psi C$ stem loop of the tRNA have been shown to be important to form the complex between tRNA and the HIV-1 viral genome during initiation of reverse transcription.

\textbf{Results:} To further delineate the features of the $T\Psi C$ stem loop of tRNA$^{\text{Lys,3}}$ in reverse transcription, we have developed a complementation system in which \textit{E. coli} tRNA$^{\text{Lys,3}}$ is provided \textit{in trans} to an HIV-1 genome in which the PBS is complementary to this tRNA. Successful selection and use of \textit{E. coli} tRNA$^{\text{Lys,3}}$ results in the production of infectious virus. We have used this single round infectious system to ascertain the effects that different mutants in the $T\Psi C$ stem loop of tRNA$^{\text{Lys,3}}$ have on complementation. Mutants were designed within the $T\Psi C$ loop (nucleotide 58) and within the stem and loop of the $T\Psi C$ loop (nucleotides 53 and 54). Analysis of the expression of \textit{E. coli} tRNA$^{\text{Lys,3}}$ mutants revealed differences in the capacity for aminoacylation, which is an indication of intracellular stability of the tRNA. Alteration of nucleotide 58 from A to U (A58U), T54G and TG5453CC all resulted in tRNA$^{\text{Lys,3}}$ that was aminoacylated when expressed in cells, while a T54C mutation resulted in a tRNA$^{\text{Lys,3}}$ that was not aminoacylated. Both the A58U and T54G mutated tRNA$^{\text{Lys,3}}$ complemented HIV-1 replication similar to wild type \textit{E. coli} tRNA$^{\text{Lys,3}}$. In contrast, the TG5453CC tRNA$^{\text{Lys,3}}$ mutant did not complement replication.

\textbf{Conclusion:} The results demonstrate that post-transcriptional modification of nucleotide 58 in tRNA$^{\text{Lys,3}}$ is not essential for HIV-1 reverse transcription. In contrast, nucleotides 53 and 54 of tRNA$^{\text{Lys,3}}$ are important for aminoacylation and selection and use of the tRNA$^{\text{Lys,3}}$ in reverse transcription.

\textbf{Background}

The major steps in reverse transcription of retroviral genome have been known for some time [1]. The initiation of reverse transcription occurs at the 5' end of the viral genome at a site designated as the primer-binding site (PBS) [1]. The PBS is an 18-nucleotide region that is complementary to the 3' terminal 18-nucleotides of the tRNA primer used for initiation [1-3]. The reverse tran-
scriptase extends the bound tRNA primer from the PBS resulting in the synthesis of minus strand DNA [4]. The reverse transcriptase then translocates to the 3' end of the viral RNA genome and proceeds to generate a complete minus-strand DNA copy of the viral RNA genome. The RNaseH activity of the viral encoded reverse transcriptase degrades the RNA copy of the viral RNA genome. Incomplete processing of the RNA by the RNaseH activity generates RNA primers for plus-strand DNA synthesis [4]. During plus-strand synthesis, the reverse transcriptase copies the tRNA primer that is attached to the minus-strand DNA to generate a plus-strand copy of the PBS. Complementation between the plus- and minus-strand PBS facilitates the completion of the viral genome, designated as the provirus.

The vast majority of the studies that have analyzed the mechanistic events of reverse transcription have utilized in vitro systems comprised of tRNA, reverse transcriptase, nuclear capsid and synthetic viral RNA/DNA templates. Previous studies have found that the tRNA\textsubscript{Lys,3} and the HIV-1 genome form a complex RNA structure for initiation of reverse transcription. As a consequence of this tRNA:RNA genome interaction, the tRNA\textsubscript{Lys,3} structure is disrupted and new intramolecular bonds are formed. One important new RNA:RNA interaction is between nucleotides 53 and 54 and the first two nucleotides of tRNALys,3 [5,6].

While in vitro studies have been informative in understanding the aspects of reverse transcription, they do not completely recapitulate all of the events in replication of the viral RNA genome. Our laboratory has approached this problem by generating HIV-1 proviruses that require the addition of exogenous tRNA for infectivity. In previous studies, we utilized an HIV-1 proviral genome in which the PBS had been mutated to be complementary to yeast tRNA\textsubscript{Phe}[7-10]. We found that the replication of this genome could be complemented if yeast tRNA\textsubscript{Phe} was supplied in trans. In vitro systems with synthetic tRNA/viral templates have been used to characterize many of the features of reverse transcription [11]. An important question that has been addressed using these systems is the role of modified tRNA bases that might play a role in stopping the reverse transcriptase during the plus-strand DNA synthesis to prevent complete copying of the tRNA primer. Since the completion of the proviral genome is facilitated by complementarity between the minus- and plus-strand DNA copies of the PBS, additional sequences in the plus-strand copy of the PBS as a result of copying of the tRNA primer would compromise the completion of the proviral genome. Previous studies have suggested that the methylated adenosine residue at position 58 (A58) of the tRNA could be a stop signal for the reverse transcriptase [12-14]. Support for this result comes from studies by Renda et al. who found that tRNA\textsubscript{Lys,3} engineered to not be methylated at A58 residue conferred a level of resistance to cells expressing this tRNA [13,14]. Additional studies though have suggested that the methylated A58 residue is not the sole stop determinant in plus-strand DNA synthesis [12].

In a recent study, we have engineered a complementation system which utilizes an HIV-1 proviral genome in which the PBS has been altered to be complementary to the 3' terminal 18-nucleotides of \textit{E. coli} tRNA\textsubscript{Lys,3} [15]. This tRNA maintains many of the unique transcriptional modifications found in mammalian tRNA\textsubscript{Lys,3}, and when expressed in mammalian cells, has been shown to be amionoaclated indicating that it is fully functional [15]. Thus, this system provides an excellent opportunity to directly address the role of the modified nucleotides and tRNA structure in HIV-1 reverse transcription. In the current study, we have found that \textit{E. coli} tRNA\textsubscript{Lys,3} with mutations at nucleotides 58, 54 and 53 in the 3' YψC loop region have varied effects in the production of infectious HIV-1. The results of our studies demonstrate that features in the 3' YψC loop of tRNA\textsubscript{Lys,3} are important for the selection and use of the tRNA as a primer for HIV-1 reverse transcription.

**Materials and methods**

### Tissue culture

293T cells, J53-3BL cells, and HeLa H1 cells were maintained in Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 10% fetal bovine serum (FBS) and 1% antibiotic (Gibco/BRL, Gaithersburg, MD). All cells were cultured in 37°C incubator supplied with 5% CO₂.

### Plasmid construction

The LS9 plasmid containing \textit{E. coli} tRNA\textsubscript{Lys,3} gene and LS9 plasmid containing mammalian tRNA\textsubscript{Lys,3} gene were constructed as previously described [15]. The \textit{E. coli} tRNA\textsubscript{Lys,3} gene in LS9 is located downstream of the human U6snRNA promoter. The A58 in \textit{E. coli} tRNA\textsubscript{Lys,3} was mutated to T using the QuickChange Side-Directed Mutagenesis (Stratagene, La Jolla, CA) with Ec A58U primers:

(\texttt{Ec A58U forward}) 5'GGTGTGCGAGAACCTCAGGACAGC3', (\texttt{Ec A58U reverse}) 5'GTGCGAGTGTTACCTGCTGGACAGAC3'. T54 in \textit{E. coli} tRNA\textsubscript{Lys,3} was substituted to G with Ec T54G primers:

(\texttt{Ec T54G forward}) 5'CAATTGGTCGAGGATCAAGTCCTGCGACCC3', (\texttt{Ec T54G reverse}) 5'GGGTCTCGACGGACATGACCGACAGGACACCC3'.

T54 in \textit{E. coli} tRNA\textsubscript{Lys,3} was also substituted to C with Ec T54C primers:

(\texttt{Ec T54C forward}) 5'CAATTGGTCGAGGATCAAGTCCTGCGAC3', (\texttt{Ec T54C reverse}) 5'GGGTCTCGACGGACATGACCGACAGGACACCC3'. T54553 together were substituted to CC with EcTG/CC primers:

(\texttt{Ec TG/CC forward})
5’GACTTITAATCAATTTGTCAGCAGGCTCAAGTGTTGCA CGACC3′, (Ec TG/CC reverse) 5’GGGTCTGAGGGCTGATGCGGCCCTGCAATTT ATAAGTC3′. All mutations were verified by DNA sequencing.

The PBS of the proviral HIV-1 genome (NL4-3) was substituted for a PBS complementary to the 3′ terminal 18-nucleotides of E. coli tRNA<sup>358</sup> by mutagenesis as described in [15]. A PBS shuttle vector with the substituted PBS was digested using the restriction enzymes BssHII and HpaI in order to release the fragment containing the E. coli tRNA<sup>358</sup> PBS region (an 868-bp fragment). The isolated fragment was ligated into the pNL4-3, which was also digested using the enzymes BssHII and HpaI. Resulting HIV-1 proviral mutant was labeled NL4-EcoLys3. Final mutants were verified by DNA sequencing.

**DNA transfections**

Co-transfections were performed according to the protocol for the Eugene 6 Transfection Reagent (Roche Molecular Biochemicals, Indianapolis, IN). Briefly, 293T cells were seeded at a concentration of 2 × 10<sup>5</sup> cells per well in 6-well plates. 500 ng of NL4-EcoLys3 and 100 ng, 250 ng, 500 ng or 1000 ng LS9 plasmids encoding E. coli tRNA<sup>358</sup> (wild type or mutations) and 3 μl Eugene reagents were added to 100 μl of DMEM. These mixtures were incubated at room temperature for approximately 45 minutes then added drop-wise to 6-well plates. The cells were supplied with fresh media 24 hours post transfection. Supernatants were collected approximately 48 hours post transfection, centrifuged at 3,000 g, and used in JC53-BL assay to determine luciferase activity, which has been determined to correlate to units of infectious virus that is being tested. Supernatants were also assayed for HIV-1 p24 antigen (Beckman Coulter, Miami, FL).

**Analysis of virus infectivity**

Serially diluted supernatants collected from co-transfections were used to infect JC53-BL cells to determine viral infectivity. JC53-BL cells were seeded 24 hours pre-infection. Infected cells were incubated for 2 hours in 37°C incubator supplemented with 5% CO<sub>2</sub>. After 2 hours, DMEM with 10% FBS was added to each well and the cells were incubated for additional 48 hours. To determine luciferase activity, cells were lysed using M-PER Mammalian Protein Extraction Reagent (Pierce, Rockford, IL) and approximated 20 μL of each lysed sample was transferred to a microplate. Reporter Lysis Buffer (Promega, Madison, WI) was added to each sample in the microplate and the light intensity was measured using a Tropix TR717 Microplate Luminometer (Applied Biosystems, Foster City, CA). Uninfected cells in wells represented background luciferase activity which was subtracted from all other samples. Relative Light Units (RLu) per mL were calculated by dividing the luciferase values by their corresponding dilutions. The total amount of virus was determined by the p24 ELISA. The amount of infectious virus was determined as RLU per nanogram p24.

**Analysis of E. coli tRNA<sup>358</sup> expressed in mammalian cells**

E. coli tRNA<sup>358</sup> plasmids were transfected into 293T cells, and total RNA was extracted 48 hours post transfection under acidic conditions to maintain the amino acid-tRNA bond [15]. One-half of the RNA sample was treated with high pH (pH 9) to serve as a de-aminoacylated control. The samples were separated in an acidic polyacrylamide gel as previously described [15]. Northern blot was carried out with NorthernMax-Gly kit (Ambion) using previously described conditions for isolation of total RNA [15]. The probe for E. coli tRNA<sup>358</sup> 5’GGGTCTGAGGGCTGATGCGGCCCTGCAATTT AAAAAAGTC3′ was phosphorylated using the ready to-go kit (Amersham) with [γ-<sup>32</sup>P]-ATP. Hybridization was carried out under standard conditions. The blots were exposed to X-ray film which was developed using an SRX-101A developer (Konica, Wayne, NJ).

**PCR and DNA sequence analysis of the PBS regions from integrated proviruses**

High molecular weight DNA (HMW) was collected using Wizard Genomic DNA Purification Kit (Promega, Madison, WI). The region encompassing the PBS was amplified with PCR using the following primers: (forward) 5‘CTCCTTCTAGCCTGCCTAGTC3′ and (reverse) 5‘GACTTTTAATCAATTGGTCGCAGCCTCAAGTCCTGCA CGACC3′. Following PCR, the products were run on 1% agarose gel and gel extracted with QIAquick Gel Extraction Kit (Qiagen, Valencia, CA) and used for DNA sequencing.

**Results**

**Construction of E. coli tRNA<sup>358</sup> mutants and HIV-1 proviral genomes**

In a recent study, we have described a complementation system which relies on the addition of E. coli tRNA<sup>358</sup> in trans to complement an HIV-1 proviral genome in which the PBS was altered to be complementary to the 3′ terminal 18-nucleotides of this tRNA [15]. E. coli tRNA<sup>358</sup> shares many features with mammalian tRNA<sup>358</sup>. In a previous study, we demonstrated that transfection of the proviral genome with a PBS to E. coli tRNA<sup>358</sup> into mammalian cells required complementation by E. coli tRNA<sup>358</sup> cDNA in order to produce infectious virus [15]. Titration of increasing amounts of E. coli tRNA<sup>358</sup> plasmid resulted in corresponding increase in levels of infectious virus.

In the current study, we have constructed mutants in the E. coli tRNA<sup>358</sup> gene in which the A58 was mutated to T.
To characterize the effect of this mutation we ascertained whether the tRNA could undergo aminoacylation following transfection of tRNA plasmid into mammalian cells. The capacity of the tRNA to undergo aminoacylation is indicative of proper identity elements, three-dimensional folding of the tRNA, and, most probably, inclusion into the translational machinery (Figure 1B). Following transfection of the wild type E. coli tRNA\textsuperscript{\text{Lys,3}} and E. coli tRNA\textsuperscript{\text{Lys,3}} A58U into mammalian cells, we analyzed these cells for the presence of aminoacylated E. coli tRNA\textsuperscript{\text{Lys,3}}. The wild type and mutants were expressed in mammalian cells following transfection and aminoacylation (Figure 1B). Thus, the lack of post-transcriptional A58 modification did not influence the capacity of this tRNA to undergo expression, transport from the nucleus to the cytoplasm, aminoacylation, and presumably inclusion into the translational machinery.

Next, we wanted to determine whether the mutant E. coli tRNA\textsuperscript{\text{Lys,3}} A58U could complement the HIV-1 proviral genome in which the PBS was altered to be complementary to the 3' terminal 18-nucleotides of this tRNA (Figure 1C). For these studies, we utilized increasing amounts of wild type and mutant E. coli tRNA\textsuperscript{\text{Lys,3}} in the transfection. The amount of virus released from the transfections was determined using the JC53-BL assay. Complementation of the proviral genome was demonstrated following transfection of the wild type E. coli tRNA\textsuperscript{\text{Lys,3}}. The levels of infectious virus increased reaching a plateau at approximately 500 ng of plasmid encoding the tRNAs. Importantly, we also found a similar level of complementation following transfection of E. coli tRNA\textsuperscript{\text{Lys,3}} A58U, confirmed by production of infectious virus. In this case, we found no clear difference between the wild type and mutant tRNA for the capacity to complement the HIV-1 proviral genome. Analysis of the PBS from the integrated proviruses after infection revealed that all were complementary to E. coli tRNA\textsuperscript{\text{Lys,3}} (data not shown). We found no evidence for additional tRNA sequences (after nucleotide A58) as would be expected if the reverse transcriptase had not stopped during copy of the tRNA during plus-strand synthesis. Collectively, the results of these studies show that the A58U mutation in tRNA\textsuperscript{\text{Lys,3}} does not impact on the capacity of this tRNA to be selected and used in HIV-1 replication.

Characterization of mutations within the TΨC stem loop of E. coli tRNA\textsuperscript{\text{Lys,3}}

Nucleotides within the TΨC stem loop were modeled to interact with HIV genome in forming the initiation complex between tRNA and viral RNA (Figure 3) [5,6]. A second interaction has also been found involving the tRNA and nucleotides within the stem of the TΨC stem loop that has also been termed the primer activation signal (PAS) [16]. To begin to address the importance of the TΨC stem loop, we made additional mutations in E. coli tRNA\textsuperscript{\text{Lys,3}}. The first mutation targeted nucleotide 54 which is a T (Figure 2A). Two mutations were made in which the thymidine at nucleotide 54 was changed to a G or C (T54G or T54C, respectively). We also made a third mutation in which the TG at nucleotides 54 and 53 were altered to Cs (TG5453CC) (Figure 2B). Each of these mutations were postulated to affect elements of the initiation complex between the tRNA and HIV viral genome.

We first characterized the effects of these mutation on E. coli tRNA\textsuperscript{\text{Lys,3}}. We determined whether the mutant tRNAs would be expressed and aminoacylated. The capacity of the tRNA to be aminoacylated correlates with the stability of this tRNA within the cell. Analysis of the aminoacylation status reveals that both T54G and TG5453CC E. coli tRNA\textsuperscript{\text{Lys,3}} mutants were predominately aminoacylated within the cell, indicating their intracellular stability. In contrast, the T54C mutation in E. coli tRNA\textsuperscript{\text{Lys,3}} was poorly aminoacylated and consequently expressed at a lower level than the other mutants (data not shown).

We next analyzed the capacity of these E. coli tRNA\textsuperscript{\text{Lys,3}} mutants to complement the replication of the HIV viral genome with the PBS complementary to E. coli tRNA\textsuperscript{\text{Lys,3}} (Figure 2C). Fixed amount of pro viral plasmid and increasing amounts of plasmids encoding wild type or mutant E. coli tRNA\textsuperscript{\text{Lys,3}} were co-transfected into mammalian cells. The activity of the resultant virus was determined using the JC53-BL assay. The mutant T54G E. coli tRNA\textsuperscript{\text{Lys,3}} readily complemented the infectivity of the mutant provirus at levels similar to that of the wild type tRNA. It was possible that the T54G E. coli tRNA\textsuperscript{\text{Lys,3}} was slightly more efficient in complementation as evidenced by increased levels at lower amounts of plasmid co-transfected with the proviral genome. However, the peak levels of complementation were similar between T54G E. coli tRNA\textsuperscript{\text{Lys,3}} and the wild type E. coli tRNA\textsuperscript{\text{Lys,3}}. Not surprisingly, the mutant T54C E. coli tRNA\textsuperscript{\text{Lys,3}} did not complement the infectivity (data not shown). Most probably this was due to poor levels of expression as a result of inability to become aminoacylated. Surprisingly, the TG5453CC E. coli tRNA\textsuperscript{\text{Lys,3}} did not complement the mutant proviral genome, even though this tRNA was expressed and aminoacylated in cells. This lack of complementation was observed throughout the entire range of the plasmids used in the titration. Thus, the results of these studies demonstrate that mutations within the TΨC loop can impact the stability of the tRNA (T54G) as well as the capacity of this tRNA to be selected and used in HIV-1 reverse transcription. The results support a role for nucleotides 54 and 53 within the TΨC stem loop in the use of tRNA\textsuperscript{\text{Lys,3}} in reverse transcription.
Complementation of HIV-1 infectivity with *E. coli* tRNA\(^{\text{Lys,3}}\) A58U mutant.

Panel A. Diagram of *E. coli* tRNA\(^{\text{Lys,3}}\) A58U mutant. The base change A58U is indicated. Boldface nucleotides indicate the 3' 18-nucleotides complementary to the PBS of HIV-1 (*E. coli* tRNA\(^{\text{Lys,3}}\)).

Panel B. Expression and aminoacylation of *E. coli* tRNA\(^{\text{Lys,3}}\) and *E. coli* tRNA\(^{\text{Lys,3}}\) A58U in 293T cells following transfection. Total RNA was isolated under acidic conditions to stabilize the amino acid tRNA bond. Approximately one-half was treated with high pH as to break the amino acid-tRNA bond (deAA). Samples were run on an acid polyacrylamide gel and blotted into nitrocellulose. All samples were analyzed with a probe specific for tRNA\(^{\text{Lys,3}}\) [15]. The migration of aminoacylated tRNA (Lane 2) and deacylated controls (Lane 1) are denoted as AA and deAA, respectively. NT is RNA from non-transfected 293T cells.

Panel C. Infectivity of NL4-EcoLys3 complemented by *E. coli* tRNA\(^{\text{Lys,3}}\) or *E. coli* tRNA\(^{\text{Lys,3}}\) A58U. 0.5 μg NL4-EcoLys3 was co-transfected with 0.1, 0.25, 0.5, 1.0 μg plasmids encoding *E. coli* tRNA\(^{\text{Lys,3}}\) or mutant into 293T cells. Virus was collected 48 hours post transfection. The amounts of infectious virus produced from transfection were determined using the JCS3-BL bioassay which measures luciferase activity [19]. The infectivity is determined by the amount of luciferase is divided by the amount of virus as determined by p24 ELISA, to give RLu per nanogram. Values are the average (+SD) from three assays.

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Figure 1

**Complementation of HIV-1 infectivity with *E. coli* tRNA\(^{\text{Lys,3}}\) A58U mutant.** Panel A. Diagram of *E. coli* tRNA\(^{\text{Lys,3}}\) A58U mutant. The base change A58U is indicated. Boldface nucleotides indicate the 3' 18-nucleotides complementary to the PBS of HIV-1 (*E. coli* tRNA\(^{\text{Lys,3}}\)).

Panel B. Expression and aminoacylation of *E. coli* tRNA\(^{\text{Lys,3}}\) and *E. coli* tRNA\(^{\text{Lys,3}}\) A58U in 293T cells following transfection. Total RNA was isolated under acidic conditions to stabilize the amino acid tRNA bond. Approximately one-half was treated with high pH as to break the amino acid-tRNA bond (deAA). Samples were run on an acid polyacrylamide gel and blotted into nitrocellulose. All samples were analyzed with a probe specific for tRNA\(^{\text{Lys,3}}\) [15]. The migration of aminoacylated tRNA (Lane 2) and deacylated controls (Lane 1) are denoted as AA and deAA, respectively. NT is RNA from non-transfected 293T cells.

Panel C. Infectivity of NL4-EcoLys3 complemented by *E. coli* tRNA\(^{\text{Lys,3}}\) or *E. coli* tRNA\(^{\text{Lys,3}}\) A58U. 0.5 μg NL4-EcoLys3 was co-transfected with 0.1, 0.25, 0.5, 1.0 μg plasmids encoding *E. coli* tRNA\(^{\text{Lys,3}}\) or mutant into 293T cells. Virus was collected 48 hours post transfection. The amounts of infectious virus produced from transfection were determined using the JCS3-BL bioassay which measures luciferase activity [19]. The infectivity is determined by the amount of luciferase is divided by the amount of virus as determined by p24 ELISA, to give RLu per nanogram. Values are the average (+SD) from three assays.
Complementation of NL4-EcoLys3 with E. coli tRNALys3 mutants. Panel A. Diagram of E. coli tRNALys3 T54G and T54C. The mutated nucleotide is indicated by a circle.

Panel B. Nucleotide diagram E. coli tRNALys3 TG54GG. The mutated nucleotides are indicated by circles.

Panel C. Infectivity of NL4-EcoLys3 complemented by E. coli tRNALys3, E. coli tRNALys3 T54G, and E. coli tRNALys3 TG5453CC mutants. 293T cells were co-transfected with 0.5 μg of proviral plasmid and with tRNA plasmids that were titrated in at the indicated quantities. Infectivity, for complementation of plasmid NL4-EcoLys3 with E. coli tRNALys3 T54G is specified by open circle; NL4-EcoLys3 with wild type E. coli tRNALys3 is specified by closed circle; NL4-EcoLys3 with E. coli tRNALys3 TG5453CC is specified by open square. The infectivity is determined by the amount of luciferase determined by JC53-BL assay divided by amount of virus (p24 ELISA) to obtain RLu per nanogram. Values are the average (+SD) from three assays.
Potential RNA secondary structure of the complex formed by NL4-EcoLys3 viral RNA and E. coli tRNALys,3

The tRNA:RNA structure was adapted from that described in previous studies [5,6]. The mutations at nucleotides 54 and 53 of E. coli tRNA^Lys,3 are boxed. The T54G mutation would be predicted to destabilize the interaction (G:G), while the TG5453CC mutation should stabilize by favorable G:C interaction.

Figure 3
Potential RNA secondary structure of the complex formed by NL4-EcoLys3 viral RNA and E. coli tRNA^Lys,3.
Discussion

Previous studies have examined the role that post-transcriptional modification of the tRNA$^{\text{Lys,3}}$ have on HIV-1 reverse transcription [12-14,17]. In one study, an in vitro system was established which recapitulates minus-strand strong stop synthesis and the plus-strand DNA synthesis. The results from these studies establish that the modified nucleotide A58 of the natural tRNA$^{\text{Lys,3}}$ was only partially effective as a stop signal [12]. The reverse transcriptase in some instances could transcribe as far as the hypermodified adenosine at A37 in the anticodon loop [17]. Based on the results of these studies, the authors concluded that the modified nucleoside at A58, which is present in all tRNA$^{\text{Lys,3}}$ molecules, appears to be important for both the efficacy and fidelity of plus-strand DNA transfer. Renda et al., extended this work and constructed cell lines or derived MuLV based vectors, to express the A58U tRNA$^{\text{Lys,3}}$ [13,14]. Analysis of the replication of HIV-1 in these cells revealed that it was slower than that observed for replication of the virus in cells which did not express the mutated tRNA$^{\text{Lys,3}}$, although the virus did eventually grow in these cells. Analysis of the resultant virus revealed that it had not undergone alteration in the PBS region. However, the inhibition of HIV-1 replication varied in individual cell clones, with some cell clones showing no inhibition. In addition, the levels of the mutated tRNA$^{\text{Lys,3}}$ were not determined in the individual cell lines, making it difficult to evaluate how the levels of mutant tRNA$^{\text{Lys,3}}$ effect viral replication and cellular metabolism. The results of these experiments suggested that mutations in A58 would have been expected to affect the capacity to produce infectious HIV-1. To further explore the potential of the A58 mutation to inhibit HIV-1 replication, we decided to determine whether the mutant tRNA would complement HIV-1 replication when provided in trans. For our studies, we engineered the HIV-1 proviral genome so that the PBS would be complementary to the 3‘ terminal nucleotides of E. coli tRNA$^{\text{Lys,3}}$. In a recent study, we have shown that E. coli tRNA$^{\text{Lys,3}}$, when provided in trans to this HIV-1 proviral genome, results in production of infectious virus [15]. If the A58 post-transcriptional modification was important for selection and use as the primer, we anticipated that co-transfection with the mutant proviral genome would not result in production of infectious virus. However, we demonstrated that the A58U mutant complemented the infectivity of the HIV-1 proviral genome at levels similar to that observed with the wild type E. coli tRNA$^{\text{Lys,3}}$. At present, we cannot resolve the differences from our study with those of Renda et al [13,14]. The results of our study are consistent with the possibility that additional features of the tRNA$^{\text{Lys,3}}$ are more important than the modified bases for the termination of plus-strand synthesis [12]. Since our in vitro complementation system has all of the appropriate viral and host cell proteins available for the process of reverse transcription, which is not the case entirely for the in vitro system, it is possible that other protein and RNA elements can compensate for the lack of modified bases. Indeed, the in vivo complementation system recapitulates both the selection process as well as the events in reverse transcription. It is also likely that the three dimensional structure of the tRNA impacts on plus-strand DNA stop [12]. Previous studies have found a complex refolding of the tRNA that occurs during initiation of the minus-strand strong stop DNA synthesis [5,6]. As a consequence, new intramolecular bonds are established in the tRNA. The new tRNA structure could be a major determinant in the effective plus-strand strong stop. The viral nucleocapsid could facilitate the maintenance of the new tRNA structure. The intracellular in trans complementation system used in the study then recapitulates the appropriate nucleocapsid-tRNA interactions, which could explain the production of infectious virus using the tRNA$^{\text{Lys,3}}$ mutants. Just how the modified bases affect the new three-dimensional RNA structure is unknown and will require additional studies.

To further explore the role of nucleotides in the tRNA$^{\text{Lys,3}}$ TΨC stem loop during reverse transcription, we made additional mutations at nucleotides 54 and 53. The T54G mutation in E. coli tRNA$^{\text{Lys,3}}$ did not affect the capacity of this tRNA to be aminoacylated or to be used in HIV reverse transcription. In contrast, mutation T54C resulted in lack of aminoacylation and consequently this E. coli tRNA$^{\text{Lys,3}}$ mutant did not complement HIV replication. Analysis of the expression level for T54C E. coli tRNA$^{\text{Lys,3}}$ revealed a decrease, upon comparison to the T54G E. coli tRNA$^{\text{Lys,3}}$, and lack of aminoacylation by the lysyl-tRNA synthetase. The lower levels of expression found for tRNAs that are unable to undergo aminoacylation is consistent with previous studies that have shown unaminoacylated tRNA instability within cells [18]. An interesting result was obtained with the double mutation TG5453CC. In this case, the TG5453CC E. coli tRNA$^{\text{Lys,3}}$ was aminoacylated and generally was produced at levels similar to that of the wild type tRNA. However, this tRNA did not complement the HIV-1 proviral genome at all tRNA plasmid concentrations tested. At present, we believe that the TG5453CC mutation in tRNA$^{\text{Lys,3}}$ precludes or retards the tRNA from forming an initiation complex with the HIV-1 RNA genome. Previous studies have established that nucleotides with the TΨC stem loop within the E. coli tRNA$^{\text{Lys,3}}$ could be involved in two potential steps during initiation of HIV-1 reverse transcription. In the first, this region is postulated to form an intramolecular bond with the 5‘ end of tRNA; this RNA:RNA interaction in tRNA$^{\text{Lys,3}}$ is postulated to help form the structure for the initiation of HIV reverse transcription [5,6] (Figure 3). However, the T54G and TG54CC mutations could be predicted to have different effects on the intramolecular tRNA$^{\text{Lys,3}}$ interactions formed during initiation. The T54G mutation would
be predicted to destabilize the interaction to an unfavorable base pair between nucleotide 1 and 54 (G:G). In contrast, the TG5453CC mutation was expected to promote this interaction through G:C base pairs. Thus, it was surprising that the T54G mutation in E. coli tRNAlys,3 still allowed complementation, while the TG5453CC mutated tRNAlys,3 did not complement. It is possible that sufficient base pair interactions still existed to form the initiation complex, but other tRNA:HIV-1 genome interactions were compromised as a result of the TG5453CC mutation (Figure 3). A second function for this sequence has been found to interact with the HIV-1 genome within the U5 region (PAS) [16]. The TG5453CC mutation would have been predicted to only partially disrupt this interaction, but it could have been sufficient to inhibit initiation. Additional studies will be required to further delineate the critical intra and intermolecular interaction between the TΨC region of tRNAlys,3 and the viral genome for initiation of reverse transcription.

Conclusion
In the current study, we have investigated the contribution that post-transcriptional modification of tRNAlys,3 at nucleotide A58 and nucleotides within the TΨC stem loop (54 and 53) have on the capacity of this tRNAlys,3 to nucleotide A58 and nucleotides within the TΨC stem loop did not complement. It is possible that sufficient base pair interactions still existed to form the initiation complex, but other tRNA:HIV-1 genome interactions were compromised as a result of the TG5453CC mutation (Figure 3). A second function for this sequence has been found to interact with the HIV-1 genome within the U5 region (PAS) [16]. The TG5453CC mutation would have been predicted to only partially disrupt this interaction, but it could have been sufficient to inhibit initiation. Additional studies will be required to further delineate the critical intra and intermolecular interaction between the TΨC region of tRNAlys,3 and the viral genome for initiation of reverse transcription.

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