Nucleocapsid mutations turn HIV-1 into a DNA-containing virus

Laurent Houzet1,2,3, Zakia Morichaud1,2,3, Ludovic Didierlaurent1,2,3, Delphine Muriaux4, Jean-Luc Darlix4 and Marylène Mougel1,2,3,*

1Université Montpellier 1, Centre d’études d’agents Pathogènes et Biotechnologies pour la Santé (CPBS), 2CNRS, UMR 5236, CPBS, 4 Bd Henri IV, CS69033, 34965 Montpellier, 3Université Montpellier 2, CPBS, 34095 Montpellier and 4LaboRetro, Unité de virologie humaine INSERM U758, IFR128, ENS, 46 allée d’Italie, Lyon, France

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ABSTRACT

Retroviruses replicate by converting their positive sense genomic RNA into double-stranded DNA that is subsequently integrated into the host genome. This conversion is catalyzed by reverse transcriptase (RT) early after virus entry into the target cell and is chaperoned by the nucleocapsid protein (NC). In HIV-1, NC is composed of small basic domains flanking two highly conserved CCHC zinc fingers that specifically interact with the genomic RNA and RT. Through specific interactions with the genomic RNA and RT, and possibly with cellular factors, the NC zinc fingers were found to play critical roles in HIV-1 assembly and budding, and later in proviral DNA synthesis and integration. Therefore, intact NC zinc fingers are needed throughout the virus replication cycle. Here, we report for the first time that deleting either one or the two NC zinc fingers leads to an unexpected premature viral DNA synthesis in virus producer cells and the production of non-infectious particles with a high level of viral DNA. In addition to providing the first example of reverse transcription during the late steps of HIV-1 replication, these findings emphasize the fact that the NC zinc fingers are a major target for new drugs against HIV-1.

INTRODUCTION

Conversion of the positive sense genomic RNA of retroviruses into DNA is a complex multistep process that is initiated from a cellular tRNA annealed to the 5’ end of the genomic RNA and culminates in the synthesis of a double-stranded DNA copy of the genomic RNA flanked by two long terminal repeats (LTR). Reverse transcription is catalyzed by the viral reverse transcriptase (RT) present in a large nucleoprotein complex where viral nucleocapsid protein (NC) molecules coat the genomic RNA (1,2). NC is encoded by Gag and is a small basic protein with nucleic acid-binding and chaperone properties, found in all retroviruses (3). In lentiviruses, such as HIV-1, NC is formed of small basic domains flanking two highly conserved CCHC zinc fingers (4) (Figure 1).

Originally, the intact NC zinc fingers were shown to be required for genomic RNA packaging (5–8) and later to be involved in virus assembly and budding (3,9–11). The selective packaging of the genomic RNA is thought to be directed by specific interactions between NC, notably the zinc fingers, and the packaging Psi signal in the genomic 5’ UTR (12). Recently, we observed that disruption of either one or the two zinc fingers impaired the intracellular Gag trafficking to the budding site, and simultaneously reduced the levels of Gag processing and of virion production (13) probably due to a lost interaction with ALIX (14).

NC also plays key roles in the reverse transcription process that needs its nucleic acid-binding and chaperone activities and its interaction with RT (3,15). Although the exact mechanism by which NC facilitates nucleic acid rearrangements is not completely understood, in vitro assays mimicking the different steps of the reverse transcription reaction clearly showed that NC chaperones primer tRNA annealing to the initiation site (PBS), and the two obligatory strand transfers that are required to generate the complete proviral DNA flanked by two LTR (3,16). However, the precise role of the NC zinc fingers in viral DNA synthesis has been difficult to evaluate in newly infected cells. Current findings show that mutating conserved residues in the first or the second CCHC zinc

*To whom correspondence should be addressed. Tel: +33 4 67 60 0232; Fax: +33 4 67 60 4420; Email: mmougel@univ-montp1.fr

The authors wish it to be known that, in their opinion, the second and the third authors be regarded as joint Second Authors
finger leads to some reverse-transcription defects with reduced DNA synthesis and stability in infected cells, and a drastic reduction of viral DNA integration (8,17,18).

Interestingly, the reverse-transcription reaction appears to be tightly controlled during the late steps of HIV-1 replication since the full-length viral DNA (FL DNA) synthesis is completed only after virions infect target cells (19). How this is regulated is yet poorly understood.

Since the NC zinc fingers appear to be required throughout the viral replication cycle, we investigated their role in the temporal control of reverse transcription. We undertook a detailed quantitative analysis of the viral nucleic acid production throughout the replication cycle by qPCR and qRT–PCR. By measuring the effects of NC zinc finger deletions on the conversion of both the genomic and spliced RNA species into DNA, we discovered that viral particles released from the cells expressing HIV-1 NC mutants, contained a high level of DNA. This report shows for the first time that RT can occur in HIV-1-producing cells, converting HIV-1 into a DNA-like virus similar to hepatitis B virus and spumaviruses. Thus, viral DNA synthesis should be regulated to ensure HIV-1 replication in target cells and appears to be a new crucial role for NC.

**MATERIALS AND METHODS**

**Plasmids, cell culture**

The HIV-1 pNL4-3 molecular clone was used to generate constructs with deletion of ZF1 (pNL4-3ΔZF1), ZF2 (pNL4-3ΔZF 2) or both ZF1 and ZF 2 (pNL4-3Δ ZF 1 ZF 2). These mutant constructs have been described elsewhere (8,13). The human HeLa LTRHIV-1-Luc (20) (kind gift of S. Emiliani), 293T, HeLa cell lines and the stably CD4/CXCR4-coexpressing-293 cells (42CD4) (21) were grown in Dulbecco’s modified Eagle’s medium (DMEM) supplemented with glutamine (2 mM), penicillin, streptomycin and 10% (v/v) heat-inactivated fetal calf serum.

**Transfection, virus preparation and infection**

Transfections of 293T cells were performed with 3 × 10⁶ cells divided 1 day before in 100 mm dishes by calcium phosphate precipitation with 8 μg of HIV-1 plasmid DNA for 48 h. When RT inhibitors were used, 293T cells were pretreated 3 h before transfection with 50 μM AZT or 50 μM Nevirapine and transfection was pursued for 48 h in presence of the drug. In all cases, 6 h after transfection cells were trypsinized, extensively washed with fresh medium and divided into a new plate to eliminate plasmid in excess. The amount of HIV-1 particles in the supernatant was determined using a HIV-1 CA p24 core antigen enzyme-linked immunosorbent assay (ELISA) Kit (Beckman Coulter) or in vitro standard RT enzyme assay (22). Virions were purified from filtered culture supernatants by centrifugation through a 20% sucrose cushion at 30 000 r.p.m. in an SW32 rotor for 1 h 30 at 4°C.

Infections of the 42CD4 cells were performed in presence of polybrene (2 μg/ml) during 24 h as previously described (23).
For luciferase reporter assay, virions produced from transfected 293T cells were quantified by measuring p24 antigen in culture supernatants (ELISA), purified and treated with DNase. Virion DNA was extracted and used to transfect HeLa cells stably expressing the HIV-1 promoter/enhancer LTR luciferase construct. HeLa LTRHIV-1-Luc cells were lyzed 72 h posttransfection. Cell protein extracts were standardized by means of the Bradford assay and luciferase activity was monitored using a luminometer.

For the natural endogenous reverse transcription assays (NERT), the virion-containing media of 293T cells were removed 44 h posttransfection, and cell culture was incubated for 4 h in different conditions: standard medium, medium without FBS (minus dNTP), supplemented or not with 100 μM dNTP and with or without 50 μM Nevirapine (AIDS Research and Reference Reagent Program, Division of AIDS, NIAID, NIH). To evaluate the effect of time on NERT, the virions produced during the 4 h of culture in standard medium, were collected and incubated for an additional 4 h in absence of cells.

DNA and RNA extraction and analysis

Nucleic acids extraction from virions was performed as previously described (23). Briefly, purified virions were incubated with 8 U of DNase (RQ1, Promega) at 37°C for 45 min before extraction of the nucleic acids by phenol/chloroform and ethanol-precipitated. DNase treatment reduced contaminations by the transfecting-plasmid DNA (pNL4-3) under the level of the FL DNA. Thus, proportions of pNL4-3 in wt and mutant DNA samples were <20% and 1% of total DNA detected, respectively. Cellular DNA was extracted with DNAzol (MRC) according to the manufacturer’s instructions. To avoid any contamination with viral cDNA associated with the particles, cells were trypsinized and extensively washed with PBS before DNA extraction. Cellular RNA was extracted with TriReagent according to manufacturer’s instructions. Nucleic acids were quantitated by measuring optical absorption at 260 nm.

To check that intracellular DNA samples were not contaminated by DNA-containing virions, untransfected cells were incubated for 24 h with supernatant containing HIV-1 ΔZF2 virions ( issued from 48 h production). After DNA extraction, contaminating multispliced cDNA levels were determined by qPCR.

In vitro reverse transcription for RT–PCR experiments was performed as described (24) with oligo(dT) primer and 1/20 aliquots of virion RNA samples. qPCR was achieved with 2.5% of the RT-d(T) reaction, or with 50 ng of intravirion and cellular nucleic acid samples. Systematically, cellular GAPDH gene level was determined for standardization. qPCR assays were performed with SYBR Green Kit (Roche) with the RotorGene (Labgene) systems. A standard curve was generated from 50 to 500 000 copies of pNL4.3 plasmid. Each assay included a control without RT indicating a DNA contamination level <0.1% of the HIV genomic RNA. Sequences of primers described in Figure 2 and in (23), and detailed PCR conditions will be provided on request.

Nucleic acid levels in virion or cell were normalized with respect to p24 (determined by ELISA) and GAPDH gene, respectively.

The HIV-1 singly spliced RNAs and corresponding cDNAs were detected in virions (100 ng p24) by standard PCR (25). In addition to the DNase treatment of the purified virions, an additional DNase treatment was performed on nucleic acid extracts to allow specific detection of FL RNA by RT–PCR and as a control for the nature of the viral cDNAs.

RESULTS

Deletions of the NC zinc fingers decreased the packaging efficiency of the FL RNA, but not of the viral spliced RNA

To study the control exerted by NC on DNA synthesis by RT, we used HIV-1 mutants with a deletion of either the first (ΔZF1), the second (ΔZF2) or both (ΔZF1ZF2) NC zinc fingers and transsected the corresponding DNA in human 293T cells (Figure 1). As previously reported, the NC mutant particles were noninfectious and were produced in reduced amounts containing less genomic RNA (FL RNA) (Figure 1) (7,13). Reduction of genomic RNA packaging was monitored by qRT–PCR (Figure 2A) on purified particles and was found to be from 10% to 1.5% of the wild-type level (Figure 1). In addition, we examined the packaging of the spliced viral RNAs that were shown to be specifically incorporated into HIV-1 particles (25). The NC zinc-finger deletions did not impair their packaging (Figures 1 and S1A) and the mutant particles contained roughly similar levels of genomic and spliced RNAs (Figure 1).

NC zinc-finger deletions resulted in the accumulation of viral DNA in virions

Infection performed with these mutant particles showed that the levels of newly made viral DNA were decreased in target cells (Figure 3, left panel) in agreement with previous results (8,17,18). Indeed, the canonical view of reverse transcription is that it mostly takes place in newly infected cells. However, some reverse transcription can take place prior to cell entry since low levels of viral DNA have been found in HIV-1 virions (23,26–30), and termed NERT for natural endogenous RT. The chaperoning role of NC protein in reverse transcription suggested to us that it could regulate the timing of viral DNA synthesis. Thus, we performed an in-depth analysis of the viral DNA content in HIV-1 NC mutant particles which have been extensively treated with DNase to remove any contaminant plasmid DNA due to cell transfection, and purified. We used qPCR for the quantitative monitoring of ss-cDNA, Gag and FL DNA. The levels of viral DNA resulting from reverse transcription of the spliced RNAs have also been monitored (Figure 2A and B).

Surprisingly, all NC mutant particles contained a high level of DNA compared with wild-type virions, notably the ΔZF2 virions, leading to DNA-to-RNA ratio increase of up to three orders of magnitude (Figure 3, right panel). Previously, we found that spliced HIV-1 RNAs were...
reverse-transcribed as efficiently as the genomic RNA (23). Thus, we also monitored the level of spliced DNAs which represent an ideal marker for viral DNA quantitation due to the lack of pNL4.3 plasmid DNA contaminations (Figure 2A). There was also a drastic enhancement of spliced cDNAs in these NC mutant virions, with about a 1000-fold increase in the /C1 ZF2 particles (Figure 3, right panel and Figure S1B). Similar results were obtained with HIV-1 particles obtained by transfecting HeLa cells (Figure S2).

A careful examination of the newly made virion DNA shows that a fraction of it corresponded to the full-length proviral DNA such as in /C1 ZF2 virions. Interestingly, DNA purified from /C1 ZF2 particles was active once transfected into the reporter cell-line HeLa LTRHIV-1-Luc since it activated the Tat-mediated luciferase expression (Figure 4).

Taken together these results suggest that deletion of the zinc fingers did not impair chaperoning of the reverse transcription by NC, in agreement with in vitro data (31). Interestingly, the /ZF2 virions contained even higher amount of viral DNA products (Figure 3, right panel) than that generated by the wild-type HIV early after cell infection (Figure 3, left panel). Thus, while the NC zinc fingers control at least in part genomic RNA packaging, these conserved NC motifs could also regulate the timing of viral DNA synthesis during the HIV-1 replication.

Viral DNA in zinc finger mutant virions did not result from NERT activity

Thus, we asked whether the virion DNA was produced in virus particles by a NERT activity somehow stimulated by the NC zinc-finger deletions. Stimulation of an otherwise marginal cDNA synthesis in virions is usually obtained by adding dNTP to the extracellular milieu together with virion permeabilization agent (32,33). Because the /ZF1, /ZF2 and /ZF1ZF2 particles exhibit an abnormal morphology with a round immature core structure (13)
Figure 1, we hypothesized that an abnormal core structure may confer NERT stimulation through a natural permeability to the dNTP that are possibly present in the fetal bovine serum (FBS) of the cell-culture media.

Since the completion of NERT takes place by 4 h (27), the DNA level in NC mutant particles should increase over time without prior virion permeabilization. However, the DNA level in particles remained the same with or without a 4 h incubation at 37°C (Figure 5A), indicating that viral DNA synthesis was already complete before incubation, probably at the time of particles release from the producer cells. In agreement with this conclusion, addition or deprivation of dNTP or treatment with the HIV-1 RT inhibitor Nevirapine during the 4 h period of virus production did not influence levels of the different viral DNA species found in ΔZF2 particles (Figure 5B).

Viral DNA was synthesized in HIV-producer cells

To examine whether reverse transcription did take place in HIV-1 producer cells, we pretreated cells with Nevirapine 3 h prior to DNA transfection, and maintained drug treatment until the viral particles were collected. Under these conditions, virion release and RNA levels in ΔZF2 virions were not affected (Figure S3), but ss-cDNA, Gag and FL DNAs and spliced cDNAs were all found in very low quantities in HIV-1 ΔZF2 virions (Figures 6, S1C and S4A). Similar results were obtained with AZT, except as expected for the initial ss-cDNA product that is known to be poorly responsive to the AZT chain terminator (23,34) (Figures S3 and S4B). Taken together, these findings indicate that viral DNA synthesis can take place in cells producing HIV-1 zinc finger mutant particles.

This prompted us to investigate whether newly made ss-cDNA, Gag and FL DNAs were present in HIV-1 producer cells. However, despite extensive DNase treatment of these cells, there was residual plasmid DNA present in the transfected cells. To circumvent this difficulty we analyzed the levels of spliced cDNAs corresponding to the reverse transcription of the spliced viral RNAs. As shown in Figure 7A, spliced viral cDNAs were abundant in cells transfected with the ΔZF2 mutant DNA, but not in those transfected with the wild-type HIV-1 DNA. In support of this, addition of the RT inhibitor Nevirapine or AZT prevented spliced cDNA synthesis in these producer cells (Figures 7B and S5).

Upon HIV-1 infection, the double-stranded linear FL DNA is synthesized and the ends can be joined by host ligases generating 2-LTR-circles (35,36). Therefore, detection of circular DNA with an LTR–LTR junction in the ΔZF2 transfected-cell (Figure 7C) indicates that some of the reverse transcripts made in producer cells are double-stranded full-length molecules. Our data clearly showed that NC zinc-finger mutation did not interfere with bona fide reverse transcription process but caused viral DNA synthesis to take place in HIV-1-producing cells, and to generate DNA-containing particles (Figure 8).
DISCUSSION

Soon after infection, viral DNA synthesis is thought to take place within reverse transcription complexes (RTC) containing mature RT, integrase (IN), Vpr, the genomic RNA and NC protein molecules (3, 37). The findings reported here favor the notion that the conserved NC zinc fingers exert, either directly or indirectly, a control over the timing of viral DNA synthesis during the late stages of HIV-1 replication. The start of viral DNA synthesis probably follows viral core formation and processing, and necessitates mature RT protein (38–40).

In producer cells, there is an accumulation of Gag and Gag-Pol precursors, containing NC and RT domains, respectively. Nevertheless, mature viral proteins and enzymes are also most probably present as indicated by the detection of mature capsid protein (p24) in HIV-1-producer cells [see for examples (13, 14, 41, 42)]. In addition, both Gag and Gag-Pol precursors can also contribute to the reverse transcription reaction. The RT domain of Pol, which has some enzymatic activity (43), can select the replication tRNALys3 primer (44, 45). The Gag–NC was found to chaperone primer tRNALys3 annealing to the genomic primer-binding site (PBS) (46) that is required for the initiation of cDNA synthesis by RT. Such a cellular environment should allow the start of viral reverse transcription and thus the presence of viral DNA in newly formed virions (47) (Figure 3, right panel).

Gag–NC is also implicated in the protease-mediated processing of Gag via its binding to the viral RNA (42, 48, 49). Although Gag processing was partially impaired when the NC zinc fingers were deleted, mature viral proteins were still easily detected in cells expressing the NC zinc fingers mutants (13, 42) and could then endorse the late RT activity observed in cells producing HIV-1 with deleted NC zinc fingers.

Interestingly, the NC zinc fingers are important determinants that are also involved in the control of HIV-1 assembly, trafficking and budding (3, 11, 50). During these processes, the NC domain mediates important protein–protein interactions with other viral proteins such as Gag-Pol, Vpr and Vif (51), and with cellular proteins, actin (52), ALIX (14), topoisomerase I (53), Staufen (54) and APOBEC3G (3, 10). These multiple Gag–NC functions likely contribute to the temporal fine-tuning of the replication steps including the timing of RT (Figure 8).

Then, how could we tentatively explain that viral DNA synthesis can take place in cells producing HIV-1 particles harboring a deletion in either one of the two zinc fingers?

The kinetics of the concerted HIV-1 assembly and budding processes might be slowed down by zinc finger deletions, allowing more time for viral DNA synthesis before particle release. Indeed, mutating or deleting the NC zinc fingers reduced the production of the NC mutant viral particles (Figure 1) with a drastic change in Gag

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**Figure 5.** No increase of NERT activity over time in ΔZF2 virions. (A) A stock of ΔZF2 virions produced during 4 h (t time) was incubated for 4 h at 37°C (t + 4 h) and viral cDNA levels were determined by qPCR (n = 3 ± SD). (B) The intravirion DNA yield was neither dependent on dNTP concentration nor Nevirapine (50 μM). DNA was analyzed in ΔZF2 virions produced by transfected 293T cells grown during 4 h in the different culture conditions (n = 3 ± SD).

**Figure 6.** Viral DNA synthesis in HIV-transfected cells. Analysis of viral DNA content of ΔZF2 virions released from cells treated or not with RT inhibitor Nevirapine (50 μM) (n = 3 ± SD).
localization (13). More precisely, they caused an accumulation of Gag in the cytoplasm or at the plasma membrane, but not at the level of late endosomes as for wild-type Gag (13), probably due to the fact that Gag–NC interactions with genomic RNA and with the cellular budding factor ALIX were impaired (14). Thus, we favor the notion that late RT is taking place in core structures formed of nonprocessed and processed Gag and Gag-Pol molecules. According to electron microscopy analysis of NC mutant cores (8,13) (Figure 1), it is likely that such intracellular viral core complexes are poorly condensed thus facilitating the reverse transcription reaction.

Figure 7. Large amounts of viral DNA in HIV-1 ΔZF2-producing cells. (A) Spliced viral cDNAs were measured by qPCR in 293T cells transfected or not with ΔZF2 or wild-type plasmids (n = 3 ± SD). (B) Effects of Nevirapine (50 μM) and AZT (50 μM) treatments on spliced viral cDNA levels were measured by qPCR in 293T cells producing ΔZF2 virions (n = 3 ± SD). (C) Circular DNA forms were detected by PCR with primers flanking the 2LTR junction in both the infected cells (42CD4) and transfected cells (293T). All DNA samples were normalized to GAPDH gene copy number. A representative experiment is shown.

Figure 8. Cartoon depicting conceptual differences between early and late RT activation during HIV-1 replication with intact NC or with a deletion of an NC zinc finger.
An alternative mechanism by which zinc finger mutations may stimulate late RT is the alleviation of cofactor(s) activity that negatively regulates RT. Several studies reported a Vif/APOBEC related regulation of the RT (51.55–57). These two proteins interact directly with Gag–NC and genomic RNA (58–60). Interestingly, Vif shares RNA chaperone activity with NC and Gag–NC proteins (51). Thus, in the context of the 293T cells in which APOBEC is poorly expressed, Vif may compete with NC-associated functions in assembly complexes and RTC. Indeed, in vitro Vif inhibits the NC-mediated tRNALys3 annealing and the initiation of RT (51). Then, Vif inhibitory effect could be relieved by the deletion of the NC zinc fingers which prevents Gag–NC/ Vif interactions, and consequently, late RT could be stimulated by Gag–NC. It will thus be of interest to investigate viral DNA synthesis in cells expressing HIV-1 NC mutants and APOBEC3G.

The occurrence of late RT inside producer cells is also a property of the hepadnaviruses (e.g. hepatitis B virus) and foamy viruses which lack NC zinc fingers and release viral DNA-containing particles (61). Even though the life cycle of the simian foamy virus is poorly understood (62), the fact that HIV-1 NC zinc finger mutants could engage a similar DNA replication strategy (Figure 8) constitutes a fundamental issue and could bring an alternative explanation for the presence of viral DNA in HIV-1 particles isolated from the peripheral blood and semen of HIV-1-infected patients (29,47).

Last, these new findings on the role of HIV-1 further highlight the fact that the conserved zinc finger motifs should be viewed as a major target for new drugs inhibiting both the late and early steps of HIV-1 replication (3,63).

**SUPPLEMENTARY DATA**

Supplementary Data are available at NAR Online.

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