HIV-1 Antisense Transcription Is Preferentially Activated in Primary Monocyte-Derived Cells

Sylvain Laverdure,a,b Antoine Gross,a,b Charlotte Arpin-André,a,b Isabelle Clerc,c Bruno Beaumelle,a,b Benoit Barbeau,d and Jean-Michel Mesnarda,b

Université Montpellier 1, Centre d'études d'agents Pathogènes et Biotechnologies pour la Santé (CPBS),* and CNRS, UMR236, CPBS,* Montpellier, France; Institut de Génétique Moléculaire de Montpellier (IGMM), CNRS, UMR5535, Montpellier, France; and Université du Québec à Montréal, Département des sciences biologiques and Centre de recherche BioMed, Montréal (Québec), Canada

In this study, an antisense luciferase-expressing human immunodeficiency virus type 1 (HIV-1) molecular clone was used to infect primary cells. We found that antisense transcription activity from the 3' long terminal repeat (LTR) was significantly more abundant in monocyte-derived cells than in activated T lymphocytes. Moreover, by analyzing antisense transcription in infected monocyte-derived dendritic cells (MDDCs), we observed that the majority of the infected MDDCs with significant antisense transcription activity did not produce Gag. We also confirmed that the negative-strand-encoded antisense protein (ASP) was expressed in monocyte-derived cells.

Human T-cell leukemia virus type 1 (HTLV-1) is the first retrovirus from which the production of antisense transcripts from the 3' long terminal repeat (LTR) has been clearly demonstrated (11, 17, 19, 22). Antisense transcription in human immunodeficiency virus type 1 (HIV-1) has also been characterized (10, 16, 20). Interestingly, such antisense transcripts can potentially encode proteins (9, 14, 15, 18). We have indeed demonstrated that the antisense protein (ASP) can be expressed in HIV-1-infected cells (12). We focused here on the regulation of HIV-1 antisense transcription in primary cells.

In this study, we used pNL4.3LucE−, a proviral DNA in which the Firefly luciferase gene had been inserted in nef (13). We removed its reporter gene to reinsert the luciferase gene in the same position but in the inverse orientation as already described (16). This vector, pNL4.3AsLucE− (Fig. 1A), permits quantification of antisense transcription from the 3' LTR, since the reporter gene was placed under the control of the antisense transcription promoter (16). As these proviral DNAs are env deficient, vesicular stomatitis virus envelope (VSVg)-pseudotyped HIV-1 particles were produced. Their decreased requirement for Nef, together with their high infectious titer and their broad host range, made these viruses useful for our experiments, which involved different target cells for infection (1). Moreover, the restriction of monocyte-derived cells to HIV-1 is also observed in this case (2). To produce pseudotyped viruses, HEK-293T cells were cotransfected with proviral DNA (30 μg) and a VSVg expression vector (18 μg). Cells were grown for 2 days before viruses were collected. Virus titers were determined by infecting Jurkat cells with 2-fold serial dilutions of virus stocks. At 2 days postinfection (p.i.), the percentage of infected cells was determined by flow cytometry following intracellular staining of Gag with K57 antibodies.

Peripheral blood mononuclear cells (PBMC) were isolated fromuffy coats, and CD14+ cells were isolated with anti-human CD14-coated magnetic beads (Miltenyi Biotec). Differentiation of CD14+ monocytes into macrophages or dendritic cells was performed as already described (6, 7). CD4+ T cells were isolated from CD14- peripheral blood lymphocytes with a CD4+ T-cell enrichment kit (Stem Cell Technologies). Cell populations were infected at a multiplicity of infection (MOI) of 2 for both virus strains, and Firefly luciferase activity was measured at different days p.i. Infection of activated primary CD4+ T lymphocytes showed a strong activation of sense transcription from the 5' LTR, while antisense transcription from the 3' LTR remained low (Fig. 1B and E); levels of antisense activity were about 1,000-fold lower than that of sense transcription. Infection of monocyte-derived macrophages (MDMs) again showed higher sense than antisense transcription levels. However, antisense transcription activity was significantly higher than in T cells (Fig. 1E), increasing steadily and reaching 1,000-fold induction at 21 days p.i. (Fig. 1C). Regulation of antisense transcription was quite different in infected monocyte-derived dendritic cells (MDDCs). During the 2 first days of infection, sense and antisense transcription levels were low, and both increased at 6 days p.i. and remained high at up to 21 days p.i. (Fig. 1D and E). Interestingly, levels of antisense transcription in MDDCs were only 25- to 100-fold lower than that of sense transcription.

We next studied the effects of the Tat transactivator on antisense transcription activity. Controversial data have been published so far (16, 20, 21). Most results were obtained from cell lines transfected with a Tat expression plasmid and reporter vectors containing only one LTR. We addressed these results using pNL4.3Luc(Reni)/AsLuc(Fire)E−, corresponding to the NL4.3 molecular clone expressing an antisense Firefly luciferase gene under the control of the 3' LTR and the Renilla luciferase gene after insertion into the env gene under the control of the Tat-dependent sense transcription (Fig. 2A). In addition, we introduced a mutation into the tat gene (pNL4.3Luc(Reni)/AsLuc(Fire)E−ΔTat), blocking its expression. Tat mutation reduced Renilla luciferase activity, while Firefly luciferase ex-

Received 4 July 2012 Accepted 25 September 2012
Published ahead of print 3 October 2012
Address correspondence to Jean-Michel Mesnard, jean-michel.mesnard@cpbs.cnrs.fr.
Copyright © 2012, American Society for Microbiology. All Rights Reserved.
doi:10.1128/JVI.01723-12
FIG 1 HIV-1 antisense transcription during viral infection in primary cells. (A) Structure of the luciferase reporter vectors pNL4.3LucE<sup>-</sup> and pNL4.3AsLucE<sup>-</sup>. The dotted line indicates the untranslated region of env due to the introduction of a stop codon. The dark gray and diagonally striped boxes correspond, respectively, to the Firefly luciferase reporter gene under the control of the 5′ LTR (sense arrow) and the 3′ LTR (antisense arrow) for sense or antisense transcription analyses. (B to D) Analysis of transcriptional sense and antisense activities in primary cells prepared from three different blood donors. Freshly isolated CD4<sup>+</sup> T cells (B) were cultured in plates coated with anti-human CD3 and anti-human CD28 antibodies (3, 8), while monocytes were differentiated into macrophages (C) or dendritic cells (D). On day 5, cells were infected with NL4.3LucE<sup>-</sup> or NL4.3AsLucE<sup>-</sup> virions pseudotyped with VSVg (MOI = 2). At different days p.i., cell extracts were prepared and normalized for protein content, and Firefly luciferase activity was subsequently measured. Luciferase values are expressed as fold increases relative to values measured in uninfected cells for each day p.i. Values represent means ± standard errors of the means (SEM) (n = 3). Luciferase activities are presented on a logarithmic scale. (E) Comparison of levels of antisense transcription activity in primary cells infected with NL4.3AsLucE<sup>-</sup> virions. Reported values are the average luminescence levels determined from the three experiments represented in panels B, C, and D. Results are represented as the luciferase activity values normalized for protein content. Values represent means ± SEM. dpi, days postinfection.

FIG 2 Study of effect of Tat on antisense transcription. (A) Generation of a proviral construct expressing the antisense Firefly luciferase gene under the control of the 3′ LTR (antisense arrow) and the sense Renilla luciferase gene under the control of the 5′ LTR (sense arrow). pNL4.3Luc(Reni)/AsLuc(Fire)E<sup>-</sup> was derived from pNL4.3AsLucE<sup>-</sup> as described for Fig. 1A. The Nhel-KpnI fragment of the pNL4.3AsLucE<sup>-</sup> env gene was replaced by the Nhel-KpnI Renilla luciferase gene from pRL-TK (Promega) (23). The dark gray box and the star correspond to the Renilla luciferase gene cloned into the env gene and the stop codon introduced in the tat gene, respectively. (B) Effect of Tat on sense and antisense transcription. HEK 293T cells were cotransfected with 1 μg of the dual-luciferase-expressing proviral DNA, 1 μg of pCMV-Tat or empty pCMV-Neo, and 1 μg of pcDNA3.1-lacZ. Cells were grown for 2 days and then lysed in passive lysis buffer (Promega). Cell extracts were normalized for protein content, and Firefly and Renilla luciferase activities were measured using the Genofax A and C reporter assay systems (Yelen Corp., France), respectively. Luciferase activities were normalized to β-galactosidase activity. Luciferase activities are presented on a logarithmic scale. Results are presented as fold induction compared to the control (CTL) data (white columns), which correspond to levels measured in transfected cells without reporter vector. Values represent means ± standard deviations (SD) (n = 3).
pression was not altered by the absence of Tat (Fig. 2B). When pNL4.3Luc(Reni)/AsLuc(Fire)E ΔTat was cotransfected with a Tat expression vector, sense transcription, unlike antisense transcription, was stimulated.

Higher antisense transcription activity was preferentially observed in HIV-1-infected MDDCs at 14 days p.i. However, the question remained whether both transcriptional activities could coexist within the same cell. Therefore, we analyzed the antisense transcription in single infected MDDCs. To identify cells capable of activating antisense transcription from the 3′ LTR, we inserted egfp under the control of the antisense transcription promoter (see Fig. 3A). The novel virus (NL4.3AsegfpE) was pseudotyped with VSVg to infect MDDCs. As a control, we produced virions from a sense enhanced green fluorescent protein (EGFP)-expressing NL4.3 clone (pNL4.3egfpE) in which egfp had been inserted in nef (Fig. 3B). At 14 days p.i., transcriptional activity of each infected MDDC was assessed by analyzing the resulting EGFP expression. Expression of gag was also analyzed by intracellular staining of Gag. Few single cells infected with NL4.3AsegfpE virus were able to express both Gag and EGFP, while that was not the case for the NL4.3segfpE virus-infected cells (Fig. 3). A total of 70.5% ± 3.1% of the EGFP-positive cells infected with NL4.3AsegfpE virus did not produce Gag, while 86.5% ± 1.5% of the EGFP-positive cells infected with NL4.3segfpE virus produced Gag (Fig. 3C). Interestingly, 90.1% ± 1.2% of Gag-expressing cells infected with NL4.3AsegfpE virus did not express antisense egfp (Fig. 3D). When the same approach was used in the presence of zidovudine (20 μM), Gag could not be detected, confirming that the Gag signal was not derived from the infecting viruses (data not shown).

ASP expression was next studied in HIV-1-infected primary cells. Cells were infected with pseudotyped NL4.3ASP-Flag virions, expressing an ASP tagged with the Flag epitope at its C-terminal end (12). While no staining was detected in activated T lymphocytes, we were able to detect ASP expression in MDMs (Fig. 4A) and MDDCs (Fig. 4B and C). No staining was detected in MDMs (Fig. 4D) or MDDCs (Fig. 4E) infected with NL4.3ASPMut12-Flag virions containing a stop codon at amino acid 12 of ASP (12).

We have provided strong evidence for the existence of antisense transcription in infected primary cells and for the production of ASP in monocyte-derived cells. Indeed, HIV-1 antisense transcription is more active in monocyte-derived cells than in activated T cells and unaltered by Tat expression. Moreover, at 14

![Image](https://via.placeholder.com/150)
days p.i., about two-thirds of HIV-1-infected MDDCs showing efficient antisense transcription activity did not express the structural Gag protein. Interestingly, our previous results had already suggested this inverse correlation between Gag and negative-stranded-encoded protein expression in HTLV-1-infected cells (4, 5, 11).

ACKNOWLEDGMENTS

This work was supported by institutional grants from the Centre National de la Recherche Scientifique (CNRS) and the Université Montpellier 1 (UM 1), a grant to J.-M.M. from the CNRS for an international project of scientific cooperation, a grant to J.-M.M. from the Agence Nationale de Recherches sur le Sida et les hépatites virales (ANRS), and a grant to B.B. from The Canadian Foundation for AIDS Research (CanFAR). B.B. holds a Canada Research Chair in Human Retrovirology (Tier 2). S.L. was supported by fellowships from the Ministère de l’Enseignement Supérieur et de la Recherche, and I.C. was supported by ANRS and Sidaction.

Confocal microscopy was performed using Montpellier RIO Imaging (MRI) facilities.

REFERENCES

1. Aiken C. 1997. Pseudotyping human immunodeficiency virus type 1 (HIV-1) by the glycoprotein of vesicular stomatitis virus targets HIV-1

---

**FIG 4** ASP expression in infected monocyte-derived cells. MDM (A and D) or MDDC (B, C, and E) cells were infected with VSVg-pseudotyped virus NL4.3ASP-Flag (A to C) or NL4.3ASPMut12-Flag (D and E) virions. At 14 days p.i., ASP expression was visualized by fluorescence microscopy after immunostaining with a primary anti-Flag antibody and a secondary antibody coupled to Alexa Fluor 568. The morphology of the cells was assessed by Normaski differential interference contrast (NDIC).
entry to an endocytic pathway and suppresses both the requirement for Nef and the sensitivity to cyclosporin A. J. Virol. 71:5871–5877.


